Shaped by their environment: variation in blue whale morphology across three productive coastal ecosystems

Data analysis code

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1/26/2023

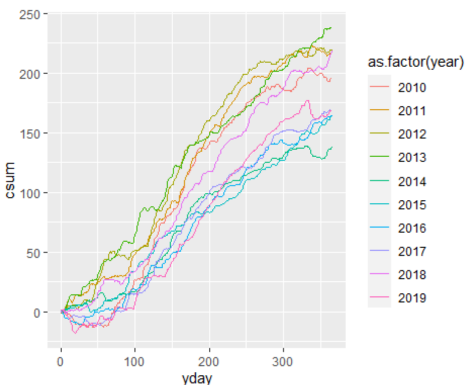
```
library(dplyr)
library(ggplot2)
library(tidyverse)
library(zoo)
library(lubridate)
library(ggplot2)
library(gridExtra)
library(ggpubr)
library(maps)
library(mapproj)
library(maptools)
library(ggstar)
# upwelling data
cuti daily <- read.csv("CUTI daily.csv",header = TRUE)</pre>
STB upwelling <- read.csv("STB upwelling 2009-2019.csv")
GC_wind <- read.csv("GC_ASCATwind_2009-2019.csv")
# productivity data
Chla.df <- read.csv("VIIRS_Chla.csv")</pre>
# morphometric data
BAI.df <- read.csv("./photogrammetry/sapphire_bai.csv")</pre>
TL.df <- read.csv("./photogrammetry/sapphire TL.csv")</pre>
UASeffort.df <- read.csv("./photogrammetry/UAS_samping_effort_corrected.csv")</pre>
```

Upwelling seasonality comparison

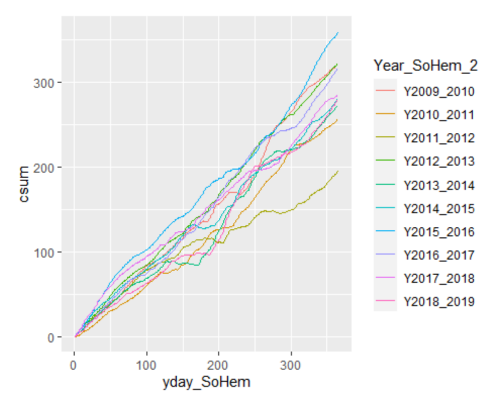
```
#### California ####
#########################
cuti_daily$date <- as.Date(with(cuti_daily, paste(year, month, day,sep="-")),
"%Y-%m-%d")
cuti_daily$yday <- yday(cuti_daily$date)

## LOOP TO CALCULATE CUMULATIVE UPWELLING FOR ALL YEARS</pre>
```

```
Years <- unique(cuti daily$year)
Years <- Years[23:32]
# apply rolling mean to whole CUTI dataset, by year - for plotting
CA_upwelling_ByYear.list <- list()</pre>
for (i in 1:length(Years)) {
  Year.i <- Years[[i]]</pre>
  ## calculate running mean of cumulative upwelling indext for the given year
  cuti_i <- filter(cuti_daily, year==Year.i)</pre>
  cuti_i <- dplyr::select(cuti_i,c("year","date","yday", "X37N"))</pre>
  cuti_i$csum <- cumsum(cuti_i$X37N) # cumulative sum</pre>
  CA_upwelling_ByYear.list[[i]] <- cuti_i</pre>
}
cuti_daily_2 <- do.call("rbind", CA_upwelling_ByYear.list)</pre>
# Quick plot
ggplot() +
  geom_line(data = cuti_daily_2, aes(x = yday, y = csum, color =
as.factor(year)))
```



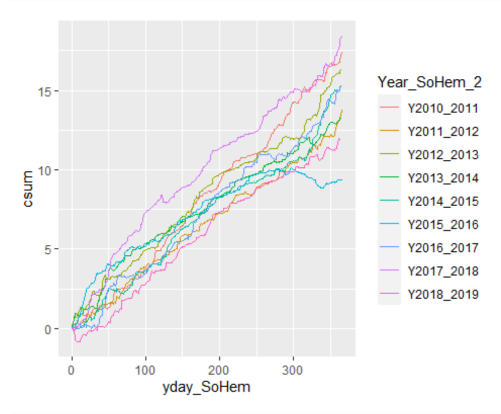
```
#### New Zealand ####
STB_upwelling$Date <- as.Date(STB_upwelling$Date)</pre>
## LOOP TO CALCULATE CUMULATIVE UPWELLING FOR ALL YEARS
Years <- unique(STB_upwelling$Year_SoHem_2)</pre>
Years <- Years[2:11] # get rid of partial (split) years at beginning and end
of time series
# apply rolling mean to whole upwelling dataset, by year (SoHem) - for
plotting
STB upwelling ByYear.list <- list()</pre>
for (i in 1:length(Years)) {
  Year_SoHem.i <- Years[[i]]</pre>
  ## calculate running mean of cumulative upwelling indext for the given year
  STBindex i <- filter(STB upwelling, Year SoHem 2==Year SoHem.i)
  STBindex_i$csum <- cumsum(STBindex_i$UpwellingIndex.MUR) # cumulative sum</pre>
  STB_upwelling_ByYear.list[[i]] <- STBindex_i</pre>
}
STB upwelling 2 <- do.call("rbind", STB upwelling ByYear.list)
# Quick plot
ggplot() +
  geom_line(data = STB_upwelling_2, aes(x = yday_SoHem, y = csum, color =
Year_SoHem_2))
```



```
#### Chile ####
GC wind$Date <- as.Date(GC wind$Date)</pre>
# fill NAs
GC_wind$Windstress <- na.locf(GC_wind$Windstress)</pre>
GC_wind$windstress_zonal <- na.locf(GC_wind$windstress_zonal)</pre>
GC_wind$windstress_meridional <- na.locf(GC_wind$windstress_meridional)
## LOOP TO CALCULATE CUMULATIVE WIND STRESS FOR ALL YEARS
Years <- unique(GC_wind$Year_SoHem_2)</pre>
Years <- Years[2:10] # get rid of partial (split) years at beginning and end
of time series
# apply rolling mean to whole wind dataset, by year (SoHem) - for plotting
GC_zonal_ByYear.list <- list()</pre>
for (i in 1:length(Years)) {
  Year_SoHem.i <- Years[[i]]</pre>
  ## calculate running mean of cumulative upwelling indext for the given year
  CGzonal_i <- filter(GC_wind, Year_SoHem_2==Year_SoHem.i)</pre>
  CGzonal i$csum <- cumsum(CGzonal i$windstress zonal) # cumulative sum
  GC_zonal_ByYear.list[[i]] <- CGzonal_i</pre>
```

```
GC_zonal_2 <- do.call("rbind", GC_zonal_ByYear.list)

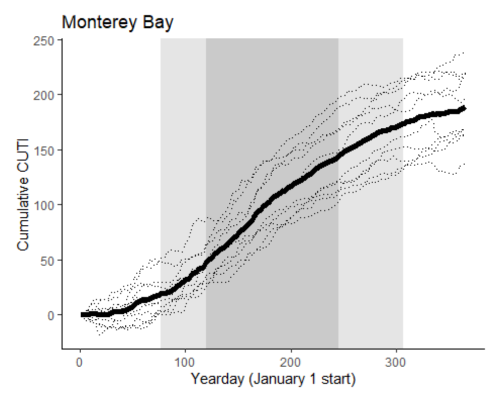
# Quick plot
ggplot() +
   geom_line(data = GC_zonal_2, aes(x = yday_SoHem, y = csum, color = Year_SoHem_2))</pre>
```



```
ca total <- max(ca clim$csumclim,na.rm = TRUE)
ca pct10 <- which.min(abs(ca clim$csumclim-(0.1*ca total)))</pre>
ca_pct25 <- which.min(abs(ca_clim$csumclim-(0.25*ca_total)))</pre>
ca pct50 <- which.min(abs(ca clim$csumclim-(0.5*ca total)))</pre>
ca_pct75 <- which.min(abs(ca_clim$csumclim-(0.75*ca_total)))</pre>
ca pct90 <- which.min(abs(ca clim$csumclim-(0.9*ca total)))</pre>
# pull dates for 50% accumulation window
ca pct25
## [1] 120
cuti daily 2$date[cuti daily 2$yday==ca pct25]
## [1] "2010-04-30" "2011-04-30" "2012-04-29" "2013-04-30" "2014-04-30"
## [6] "2015-04-30" "2016-04-29" "2017-04-30" "2018-04-30" "2019-04-30"
ca pct75
## [1] 245
cuti_daily_2$date[cuti_daily_2$yday==ca_pct75]
## [1] "2010-09-02" "2011-09-02" "2012-09-01" "2013-09-02" "2014-09-02"
## [6] "2015-09-02" "2016-09-01" "2017-09-02" "2018-09-02" "2019-09-02"
ca_pct75 - ca_pct25
## [1] 125
# yday 119-239; 29 April - 27 August; 120 days
## New Zealand
# calculate climatological mean cumulative upwelling curve
nz_clim <- data.frame(matrix(ncol = 2, nrow = 366))</pre>
colnames(nz_clim) <- c("yday_SoHem","csumclim")</pre>
for (i in 1:366) {
  c <- STB upwelling 2 %>% filter(yday SoHem == i)
  nz_clim$yday_SoHem[i] <- i</pre>
  nz_clim$csumclim[i] <- mean(c$csum,na.rm = TRUE)</pre>
# calculate 50%, 25-75%, and 10-90% yeardays on the climatological upwelling
curve
nz total <- max(nz clim$csumclim,na.rm = TRUE)</pre>
nz_pct10 <- which.min(abs(nz_clim$csumclim-(0.1*nz_total)))</pre>
nz_pct25 <- which.min(abs(nz_clim$csumclim-(0.25*nz_total)))</pre>
nz pct50 <- which.min(abs(nz clim$csumclim-(0.5*nz total)))</pre>
nz pct75 <- which.min(abs(nz clim$csumclim-(0.75*nz total)))</pre>
nz_pct90 <- which.min(abs(nz_clim$csumclim-(0.9*nz_total)))</pre>
```

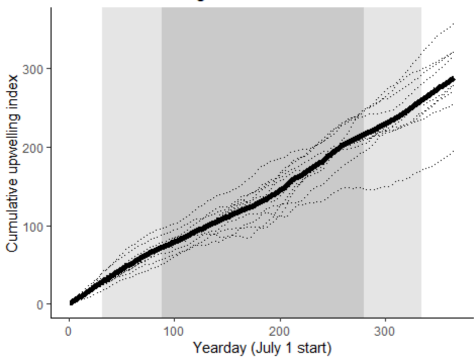
```
# pull dates for 50% accumulation window
nz pct25
## [1] 88
STB_upwelling_2$Date[STB_upwelling_2$yday_SoHem==nz_pct25]
## [1] "2009-09-26" "2010-09-26" "2011-09-26" "2012-09-26" "2013-09-26"
## [6] "2014-09-26" "2015-09-26" "2016-09-26" "2017-09-26" "2018-09-26"
nz_pct75
## [1] 280
STB_upwelling_2$Date[STB_upwelling_2$yday_SoHem==nz_pct75]
## [1] "2010-04-06" "2011-04-06" "2012-04-05" "2013-04-06" "2014-04-06"
## [6] "2015-04-06" "2016-04-05" "2017-04-06" "2018-04-06" "2019-04-06"
nz_pct75 - nz_pct25
## [1] 192
# yday_Sohem 85-275; 23 September - 1 April; 190 days
## Chile
# calculate climatological mean cumulative wind stress curve
gc_clim_zonal <- data.frame(matrix(ncol = 2, nrow = 366))</pre>
colnames(gc clim zonal) <- c("yday SoHem","csumclim")</pre>
for (i in 1:366) {
  c <- GC zonal 2 %>% filter(yday SoHem == i)
  gc_clim_zonal$yday_SoHem[i] <- i</pre>
  gc_clim_zonal$csumclim[i] <- mean(c$csum,na.rm = TRUE)</pre>
}
# calculate 50%, 25-75%, and 10-90% yeardays on the climatological upwelling
gc_zonal_total <- max(gc_clim_zonal$csumclim,na.rm = TRUE)</pre>
gc_zonal_pct10 <- which.min(abs(gc_clim_zonal$csumclim-(0.1*gc_zonal_total)))</pre>
gc_zonal_pct25 <- which.min(abs(gc_clim_zonal$csumclim-</pre>
(0.25*gc zonal total)))
gc_zonal_pct50 <- which.min(abs(gc_clim_zonal$csumclim-(0.5*gc_zonal_total)))</pre>
gc_zonal_pct75 <- which.min(abs(gc_clim_zonal$csumclim-</pre>
(0.75*gc zonal total)))
gc_zonal_pct90 <- which.min(abs(gc_clim_zonal$csumclim-(0.9*gc_zonal_total)))</pre>
# pull dates for 50% accumulation window
gc_zonal_pct25
## [1] 81
```

```
GC zonal 2$Date[GC zonal 2$yday SoHem==gc zonal pct25]
## [1] "2010-09-19" "2011-09-19" "2012-09-19" "2013-09-19" "2014-09-19"
## [6] "2015-09-19" "2016-09-19" "2017-09-19" "2018-09-19"
gc_zonal_pct75
## [1] 286
GC_zonal_2$Date[GC_zonal_2$yday_SoHem==gc_zonal_pct75]
## [1] "2011-04-12" "2012-04-11" "2013-04-12" "2014-04-12" "2015-04-12"
## [6] "2016-04-11" "2017-04-12" "2018-04-12" "2019-04-12"
gc_zonal_pct75 - gc_zonal_pct25
## [1] 205
# yday Sohem 75-266; 13 September - 23 March; 191 days
#### Climatology plots w/ percent accumulation for seasonality ####
## only include yday 1-365 (366 is less common and therefore not robust in
the climatological curve)
ca clim <- ca clim[1:365,]
nz clim <- nz clim[1:365,]</pre>
gc_clim_zonal <- gc_clim_zonal[1:365,]</pre>
## California
annual_CUTI_CA.plot <- ggplot() +</pre>
  annotate("rect", fill = "gray", alpha = 0.4,
          xmin = ca_pct10, xmax = ca_pct90,
          ymin = -Inf, ymax = Inf) +
  annotate("rect", fill = "gray", alpha = 0.7,
          xmin = ca_pct25, xmax = ca_pct75,
          ymin = -Inf, ymax = Inf) +
  geom_line(data = cuti_daily_2, aes(x = yday, y = csum, group = year),
linetype = "dotted") +
  geom line(data = ca clim, aes(x = yday, y = csumclim), size = 2) +
  theme classic() +
  theme(legend.position = "none") +
  xlab("Yearday (January 1 start)") +
  ylab("Cumulative CUTI") +
  ggtitle("Monterey Bay")
annual_CUTI_CA.plot
```

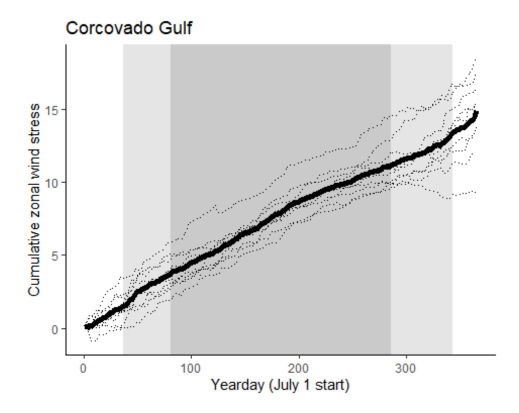


```
## New Zealand
annual_upwelling_STB.plot <- ggplot() +</pre>
  annotate("rect", fill = "gray", alpha = 0.4,
           xmin = nz_pct10, xmax = nz_pct90,
           ymin = -Inf, ymax = Inf) +
  annotate("rect", fill = "gray", alpha = 0.7,
           xmin = nz_pct25, xmax = nz_pct75,
           ymin = -Inf, ymax = Inf) +
  geom_line(data = STB_upwelling_2, aes(x = yday_SoHem, y = csum, group =
Year_SoHem), linetype = "dotted") +
  geom_line(data = nz_clim, aes(x = yday_SoHem, y = csumclim), size = 2) +
  theme_classic() +
  theme(legend.position = "none") +
  xlab("Yearday (July 1 start)") +
  ylab("Cumulative upwelling index") +
  ggtitle("South Taranaki Bight")
annual_upwelling_STB.plot
```

South Taranaki Bight



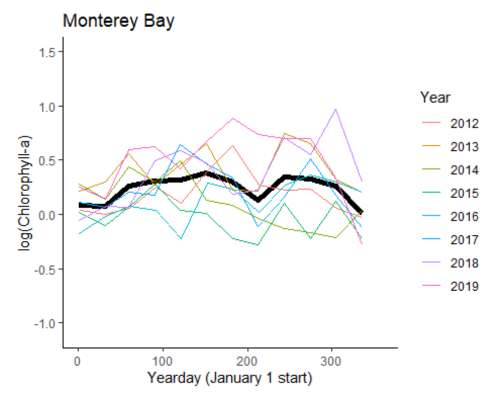
```
## Chile
annual_zonal_GC.plot <- ggplot() +</pre>
  annotate("rect", fill = "gray", alpha = 0.4,
           xmin = gc_zonal_pct10, xmax = gc_zonal_pct90,
           ymin = -Inf, ymax = Inf) +
  annotate("rect", fill = "gray", alpha = 0.7,
           xmin = gc_zonal_pct25, xmax = gc_zonal_pct75,
           ymin = -Inf, ymax = Inf) +
  geom_line(data = GC_zonal_2, aes(x = yday_SoHem, y = csum, group =
Year_SoHem), linetype = "dotted") +
  geom_line(data = gc_clim_zonal, aes(x = yday_SoHem, y = csumclim), size =
2) +
  theme classic() +
  theme(legend.position = "none") +
  xlab("Yearday (July 1 start)") +
  ylab("Cumulative zonal wind stress") +
  ggtitle("Corcovado Gulf")
annual zonal GC.plot
```



Productivity seasonality comparison

```
# only include years that overlap with upwelling data
Chla.df <- Chla.df[Chla.df$Year < 2020, ]</pre>
Chla.df$Chl_STB[Chla.df$Year_SoHem=="Y2020_2021"] <- NA
Chla.df$Chl GC[Chla.df$Year SoHem=="Y2020 2021"] <- NA
Chla.df$logChl_STB[Chla.df$Year_SoHem=="Y2020_2021"] <- NA</pre>
Chla.df$logChl GC[Chla.df$Year SoHem=="Y2020 2021"] <- NA
# Separate Chl data by year
Chla.df$Year <- as.factor(Chla.df$Year)</pre>
# Aggregate monthly mean values
Chla_annual_CA <- aggregate(logChl_CA ~ Month, Chla.df, mean)</pre>
# Assign yearday to monthly mean, for plotting purposes
Chla_annual_CA$yday <- Chla.df$yday[Chla.df$Year=="2014" &
Chla.df$Month==Chla_annual_CA$Month]
# Aggregate monthly mean values
Chla_annual_STB <- aggregate(logChl_STB ~ Month, Chla.df, mean)</pre>
# Assign yearday to monthly mean, for plotting purposes
Chla_annual_STB$yday_SoHem <- Chla.df$yday_SoHem[Chla.df$Year=="2014" &</pre>
Chla.df$Month==Chla annual STB$Month]
```

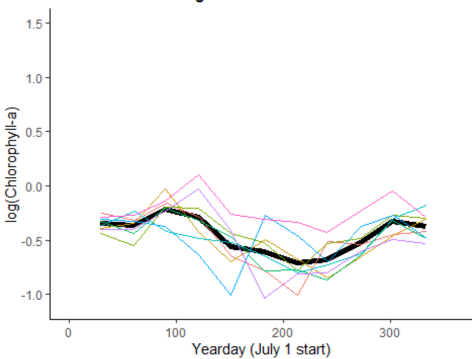
```
# Aggregate monthly mean values
Chla annual GC <- aggregate(logChl GC ~ Month, Chla.df, mean)
# Assign yearday to monthly mean, for plotting purposes
Chla_annual_GC$yday_SoHem <- Chla.df$yday_SoHem[Chla.df$Year=="2014" &
Chla.df$Month==Chla_annual_GC$Month]
Chl CA.plot <- ggplot() +
  geom_line(data = Chla_annual_CA, aes(x=yday, y=logChl_CA), size=2) +
  geom_line(data = Chla.df,aes(x=yday, y=logChl_CA, group = Year,
color=Year)) +
  ylim(-1.1,1.5) + xlim(0,360) +
  theme_classic() +
  #theme(legend.position = "none") +
  xlab("Yearday (January 1 start)") +
  ylab("log(Chlorophyll-a)") +
  ggtitle("Monterey Bay")
Chl CA.plot
```



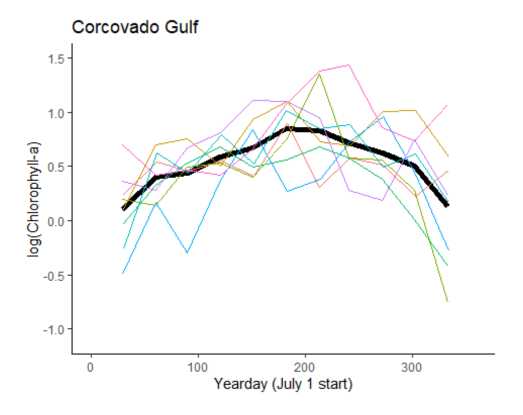
```
Chl_STB.plot <- ggplot() +
   geom_line(data = Chla_annual_STB, aes(x=yday_SoHem, y=logChl_STB), size=2)
+
   geom_line(data = Chla.df,aes(x=yday_SoHem, y=logChl_STB, group = Year,
color=Year)) +
   ylim(-1.1,1.5) + xlim(0,360) +
   theme_classic() +
   theme(legend.position = "none") +</pre>
```

```
xlab("Yearday (July 1 start)") +
ylab("log(Chlorophyll-a)") +
ggtitle("South Taranaki Bight")
Chl_STB.plot
```

South Taranaki Bight



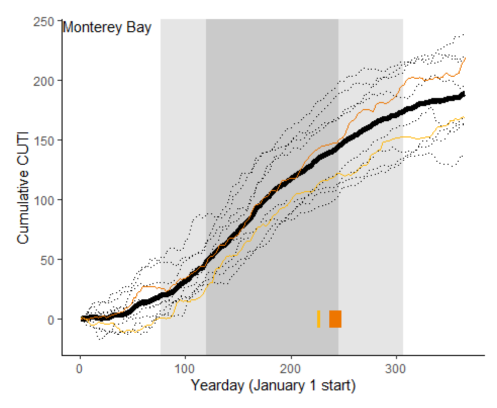
```
Chl_GC.plot <- ggplot() +
    geom_line(data = Chla_annual_GC, aes(x=yday_SoHem, y=logChl_GC), size=2) +
    geom_line(data = Chla.df,aes(x=yday_SoHem, y=logChl_GC, group = Year,
color=Year)) +
    ylim(-1.1,1.5) + xlim(0,360) +
    theme_classic() +
    theme(legend.position = "none") +
    xlab("Yearday (July 1 start)") +
    ylab("log(Chlorophyll-a)") +
    ggtitle("Corcovado Gulf")
Chl_GC.plot</pre>
```



Configure seasonality plots: highlight UAS data years, combine with total length and body condition data by year

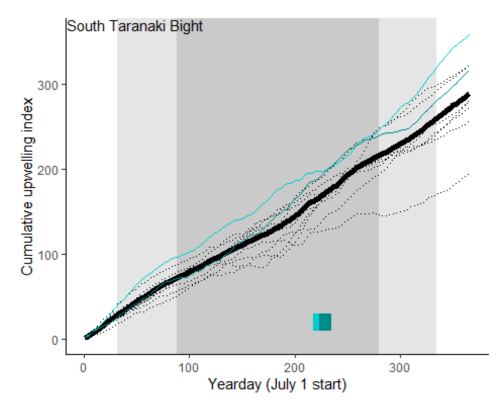
```
# UPWELLING
## California
annual_CUTI_CA.plot <- ggplot() +</pre>
  annotate("rect", fill = "gray", alpha = 0.4,
           xmin = ca pct10, xmax = ca pct90,
           ymin = -Inf, ymax = Inf) +
  annotate("rect", fill = "gray", alpha = 0.7,
           xmin = ca_pct25, xmax = ca_pct75,
           ymin = -Inf, ymax = Inf) +
  geom_line(data = cuti_daily_2, aes(x = yday, y = csum, group = year),
linetype = "dotted") +
  geom_line(data = ca_clim, aes(x = yday, y = csumclim), size = 2) +
  geom_line(data = cuti_daily_2[cuti_daily_2$year=="2017",], aes(x = yday, y
= csum, group = year), color="darkgoldenrod1") +
  geom_line(data = cuti_daily_2[cuti_daily_2$year=="2018",], aes(x = yday, y
= csum, group = year), color="darkorange2") +
  geom_line(aes(x=c(225,228), y=0), color="darkgoldenrod1", size=6) +
  geom\_line(aes(x=c(237,248), y=0), color="darkorange2", size=6) +
  theme classic() +
```

```
theme(legend.position = "none") +
    xlab("Yearday (January 1 start)") +
    ylab("Cumulative CUTI") +
    annotate("text", x= -Inf, y = Inf, label = "Monterey Bay", vjust=1,
hjust=-0.01)
annual_CUTI_CA.plot
```



```
## New Zealand
annual_upwelling_STB.plot <- ggplot() +</pre>
  annotate("rect", fill = "gray", alpha = 0.4,
           xmin = nz pct10, xmax = nz pct90,
           ymin = -Inf, ymax = Inf) +
  annotate("rect", fill = "gray", alpha = 0.7,
           xmin = nz_pct25, xmax = nz_pct75,
           ymin = -Inf, ymax = Inf) +
  geom_line(data = STB_upwelling_2, aes(x = yday_SoHem, y = csum, group =
Year SoHem), linetype = "dotted") +
  geom_line(data = nz_clim, aes(x = yday_SoHem, y = csumclim), size = 2) +
  geom_line(data = STB_upwelling_2[STB_upwelling_2$Year_SoHem=="2016",],
aes(x = yday_SoHem, y = csum, group = Year_SoHem), color="cyan3") +
  geom_line(data = STB_upwelling_2[STB_upwelling_2$Year_SoHem=="2017",],
aes(x = yday_SoHem, y = csum, group = Year_SoHem), color="cyan4") +
  geom_line(aes(x=c(217,223), y=20), color="cyan3", size=6) +
  geom_line(aes(x=c(223,235), y=20), color="cyan4", size=6) +
```

```
theme_classic() +
  theme(legend.position = "none") +
  xlab("Yearday (July 1 start)") +
  ylab("Cumulative upwelling index") +
  annotate("text", x= -Inf, y = Inf, label = "South Taranaki Bight",
  vjust=1, hjust=-0.01)
annual_upwelling_STB.plot
```

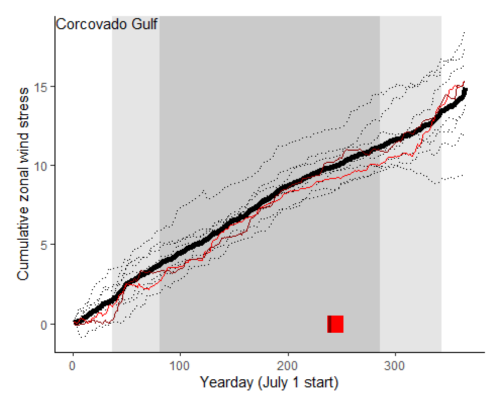


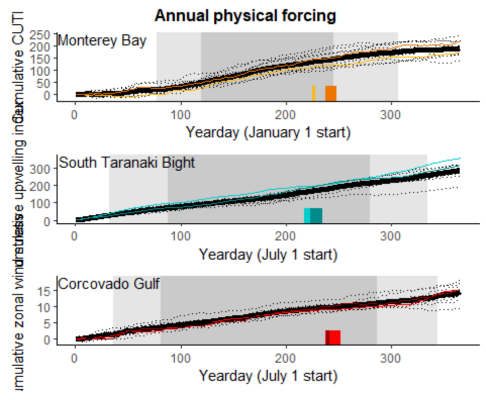
```
## Chile
annual_zonal_GC.plot <- ggplot() +</pre>
  annotate("rect", fill = "gray", alpha = 0.4,
           xmin = gc_zonal_pct10, xmax = gc_zonal_pct90,
           ymin = -Inf, ymax = Inf) +
  annotate("rect", fill = "gray", alpha = 0.7,
           xmin = gc_zonal_pct25, xmax = gc_zonal_pct75,
           ymin = -Inf, ymax = Inf) +
  geom_line(data = GC_zonal_2, aes(x = yday_SoHem, y = csum, group =
Year_SoHem), linetype = "dotted") +
  geom_line(data = gc_clim_zonal, aes(x = yday_SoHem, y = csumclim), size =
2) +
  geom line(data = GC zonal 2[GC zonal 2$Year SoHem=="2015",], aes(x =
yday_SoHem, y = csum, group = Year_SoHem), color="red") +
  geom_line(data = GC_zonal_2[GC_zonal_2$Year_SoHem=="2017",], aes(x =
yday_SoHem, y = csum, group = Year_SoHem), color="dark red") +
```

```
geom_line(aes(x=c(237, 252), y=0), color="red", size=6) +
geom_line(aes(x=c(238, 241), y=0), color="dark red", size=6) +

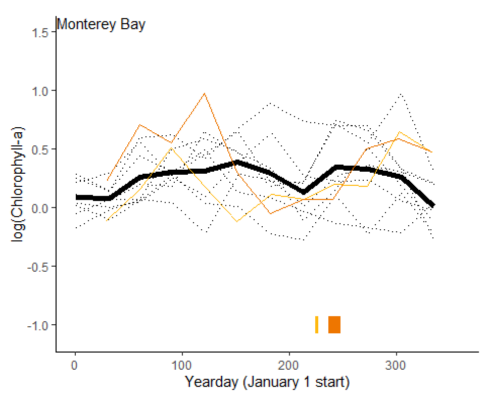
theme_classic() +
theme(legend.position = "none") +
xlab("Yearday (July 1 start)") +
ylab("Cumulative zonal wind stress") +
annotate("text", x= -Inf, y = Inf, label = "Corcovado Gulf", vjust=1,
hjust=-0.01)

annual_zonal_GC.plot
```

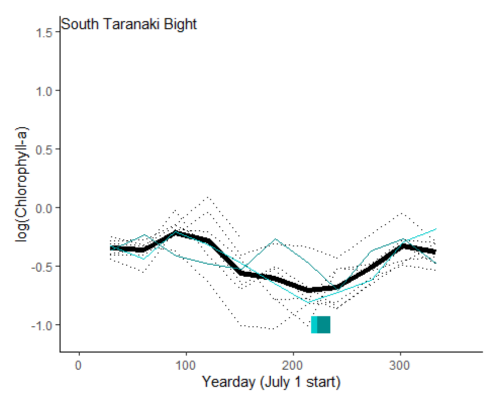




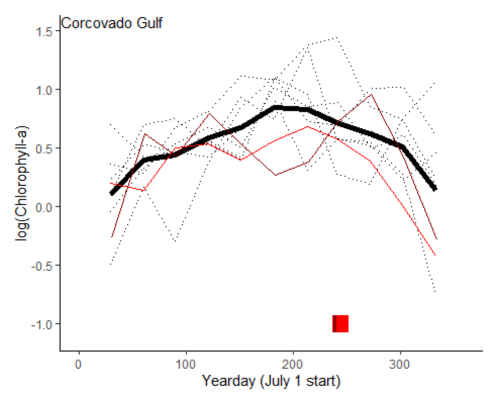
```
# PRODUCTIVITY
Chl CA.plot <- ggplot() +
  geom_line(data = Chla_annual_CA, aes(x=yday, y=logChl_CA), size=2) +
  geom_line(data = Chla.df,aes(x=yday, y=logChl_CA, group = Year), linetype =
"dotted") +
  geom line(data = Chla.df[Chla.df$Year=="2017",], aes(x = yday SoHem, y =
logChl CA, group = Year), color="darkgoldenrod1") +
  geom line(data = Chla.df[Chla.df$Year=="2018",], aes(x = yday SoHem, y =
logChl_CA, group = Year), color="darkorange2") +
  geom_line(aes(x=c(225,228), y=-1), color="darkgoldenrod1", size=6) +
  geom_line(aes(x=c(237,248), y=-1), color="darkorange2", size=6) +
  ylim(-1.1,1.5) + xlim(0,360) +
  theme_classic() +
  theme(legend.position = "none") +
  xlab("Yearday (January 1 start)") +
  ylab("log(Chlorophyll-a)") +
  annotate("text", x= -Inf, y = Inf, label = "Monterey Bay", vjust=1,
hjust=-0.01)
Chl_CA.plot
```

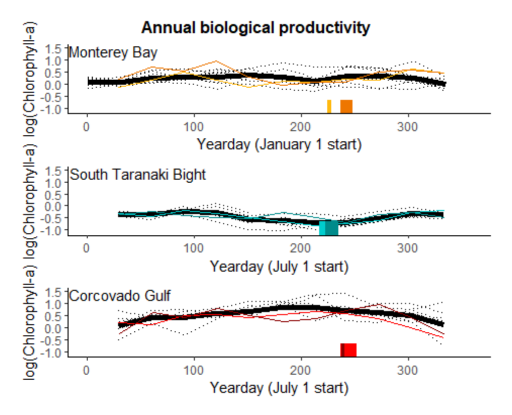


```
Chl_STB.plot <- ggplot() +</pre>
  geom_line(data = Chla_annual_STB, aes(x=yday_SoHem, y=logChl_STB), size=2)
  geom line(data = Chla.df,aes(x=yday_SoHem, y=logChl_STB, group =
Year_SoHem), linetype = "dotted") +
  geom_line(data = Chla.df[Chla.df$Year_SoHem=="Y2015_2016",], aes(x =
yday_SoHem, y = logChl_STB, group = Year_SoHem), color="cyan3") +
  geom line(data = Chla.df[Chla.df$Year SoHem=="Y2016 2017",], aes(x =
yday_SoHem, y = logChl_STB, group = Year_SoHem), color="cyan4") +
  geom_line(aes(x=c(217,223), y=-1), color="cyan3", size=6) +
  geom_line(aes(x=c(223,235), y=-1), color="cyan4", size=6) +
  ylim(-1.1,1.5) + xlim(0,360) +
  theme_classic() +
  theme(legend.position = "none") +
  xlab("Yearday (July 1 start)") +
  ylab("log(Chlorophyll-a)") +
  annotate("text", x= -Inf, y = Inf, label = "South Taranaki Bight",
vjust=1, hjust=-0.01)
Chl STB.plot
```



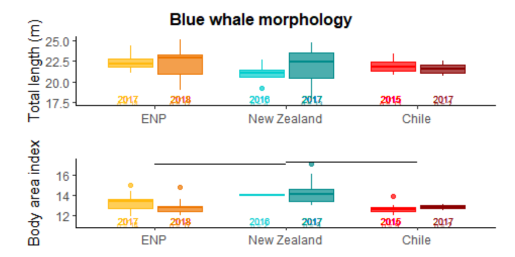
```
Chl_GC.plot <- ggplot() +</pre>
  geom_line(data = Chla_annual_GC, aes(x=yday_SoHem, y=logChl_GC), size=2) +
  geom_line(data = Chla.df,aes(x=yday_SoHem, y=logChl_GC, group =
Year_SoHem), linetype = "dotted") +
  geom_line(data = Chla.df[Chla.df$Year_SoHem=="Y2014_2015",], aes(x =
yday_SoHem, y = logChl_GC, group = Year_SoHem), color="red") +
  geom_line(data = Chla.df[Chla.df$Year_SoHem=="Y2016_2017",], aes(x =
yday SoHem, y = logChl GC, group = Year SoHem), color="dark red") +
  geom_line(aes(x=c(237, 252), y=-1), color="red", size=6) +
  geom_line(aes(x=c(238, 241), y=-1), color="dark red", size=6) +
  ylim(-1.1,1.5) + xlim(0,360) +
  theme_classic() +
  theme(legend.position = "none") +
  xlab("Yearday (July 1 start)") +
  ylab("log(Chlorophyll-a)") +
  annotate("text", x= -Inf, y = Inf, label = "Corcovado Gulf", vjust=1,
hjust=-0.01
Chl_GC.plot
```





```
# BODY CONDITION
BAI.df$Location <- factor(BAI.df$Location, levels = c("CA", "NZ", "Chile"))
TL.df$Location <- factor(TL.df$Location, levels = c("CA", "NZ", "Chile"))
TL.df$Location_Year <- paste0(TL.df$Location, "_", TL.df$Year)</pre>
TL samplesizes.df <- aggregate(AID ~ Location + Year, TL.df, length)
TL_samplesizes.df$Location_Year <- paste0(TL_samplesizes.df$Location, "_",
TL_samplesizes.df$Year)
TL samplesizes.df$Location <- factor(TL samplesizes.df$Location, levels =
c("CA", "NZ", "Chile"))
BAI.df$Location_Year <- paste0(BAI.df$Location, "_", BAI.df$Year)
BAI_samplesizes.df <- aggregate(AID ~ Location + Year, BAI.df, length)</pre>
BAI_samplesizes.df$Location_Year <- paste0(BAI_samplesizes.df$Location, " ",
BAI samplesizes.df$Year)
BAI_samplesizes.df$Location <- factor(BAI_samplesizes.df$Location, levels =
c("CA", "NZ", "Chile"))
fillcols b <- c("darkgoldenrod1", "darkorange2", "red", "dark red", "cyan3",
"cyan4")
TL.plot <- ggplot(data = TL.df, aes(y=TL.mean, x=Location)) +
  geom boxplot(aes(color = Location Year, fill = Location Year), alpha = 0.7)
  scale_color_manual(values = fillcols_b) + scale_fill_manual(values =
```

```
fillcols b) +
  geom text(aes(x = Location, y = 18, label = Year, color=Location Year),
position=position_dodge(width=0.8), size=2.5) +
  geom text(data = TL samplesizes.df, aes(x = Location, y = 17.5, label =
paste("n =", AID), color=Location_Year),
            position=position_dodge(width=0.8), size=2) +
ylab("Total length (m)") + xlab(" ") + guides(fill="none", color="none") +
scale_x_discrete(labels=c("ENP", "New Zealand", "Chile")) +
  theme classic()
#TL.plot
BAI.plot <- ggplot(data = BAI.df, aes(y=BAI.mean, x=Location)) +
  geom_boxplot(aes(color = Location_Year, fill = Location_Year), alpha = 0.7)
  scale color manual(values = fillcols b) + scale fill manual(values =
fillcols b) +
  geom_text(aes(x = Location, y = 11.5, label = Year, color=Location_Year),
position=position_dodge(width=0.8), size=2.5) +
  geom_text(data = BAI_samplesizes.df, aes(x = Location, y = 11.1, label =
paste("n =", AID), color=Location_Year),
            position=position dodge(width=0.8), size=2) +
  geom_segment(aes(x="CA", xend="NZ", y=17.1, yend=17.1)) +
  geom_segment(aes(x="NZ", xend="Chile", y=17.35, yend=17.35)) +
  geom_text(aes(x="CA", y=17.1, label="*"), vjust=-0.005, hjust=-6, size=3.5)
  geom_text(aes(x="NZ", y=17.35, label="*"), vjust=-0.005, hjust=-6,
size=3.5) +
  ylab("Body area index") + xlab(" ") + guides(fill="none", color="none") +
  scale x discrete(labels=c("ENP", "New Zealand", "Chile")) +
  theme classic()
#BAI.plot
morph.plot <- ggarrange(TL.plot, BAI.plot, nrow = 3, align = "hv")</pre>
morph.plot <- annotate figure(morph.plot, top = text grob("Blue whale</pre>
morphology",
               color = "Black", face = "bold", size = 12))
morph.plot
```



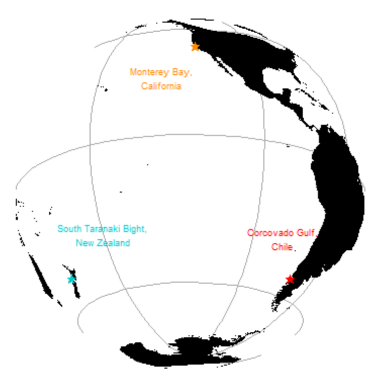
ALL TOGETHER NOW

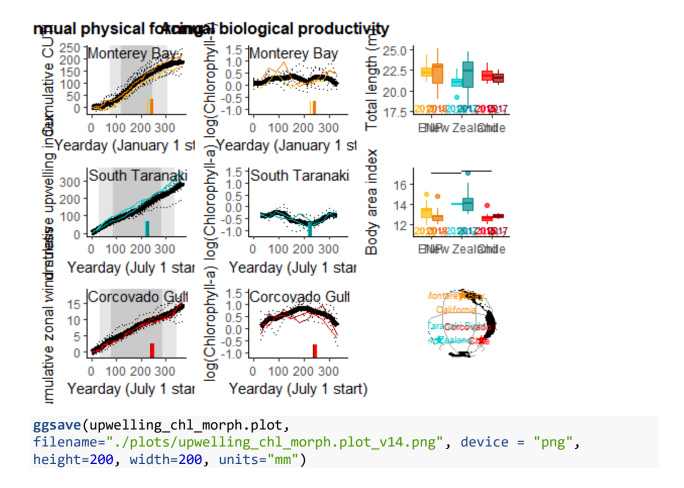
```
upwelling_chl_morph.plot <- ggarrange(upwelling_season_combined.plot,
Chl_annual_combined.plot, morph.plot, nrow = 1, align = "hv")
upwelling_chl_morph.plot
```

nrual physical fokcingat biological productivitywhale morpholog 250 Monterey Bay 25.0 1.5 Monterey Bay Chlorophy 200 Worker 150 - 1 1.0 £ 22.5 20.0 0.0 -0.5Total -1.017.5 ğ 0 100 200 300 ⊟NMeW Zeal@midle _ ভূYearday (January 1 st index mulative zonal windratrlasise upwelling Chlorophyll 1.5 1.0 South Taranaki South Taranaki 16 Body area 0.5 14 -0.512 -1.0) Bo 100 200 300 100 200 300 ⊟NMeW Zeal@midle Yearday (July 1 star Yearday (July 1 star ® Chlorophyll. 1.5 Corcovado Gull Corcovado Gull 0.5 0.0 -0.5-1.0 ğ 100 200 300 0 100 200 300 Yearday (July 1 star Yearday (July 1 start)

```
## Add map with study regions
# In the maps package, world2 is the Pacific centered map with longitude
[0,360]
world2 <- map_data("world2")</pre>
pacific.map <- ggplot() +</pre>
  \#geom\ map(data = world\ map,\ map=world\ map,\ aes(x=long,\ y=lat,\ map\ id=id),
fill="black") +
  geom_map(data = world2, map=world2, aes(x=long, y=lat, map_id=region),
fill="black") +
  coord_map("ortho", orientation = c(-20,235,0)) + # requires mapproj package
  geom_star(aes(x=-123, y=37), color = "darkorange", fill = "darkorange",
size=1.75) +
  geom_star(aes(x=173, y=-40), color = "cyan3", fill = "cyan3", size=1.75) +
  geom star(aes(x=-74, y=-43), color = "red", fill = "red", size=1.75) +
  annotate(geom = "text", x=-135, y=22, label = "Monterey Bay,\nCalifornia",
color = "darkorange", size = 2.5) +
  annotate(geom = "text", x=-160, y=-30, label = "South Taranaki Bight,\nNew
Zealand", color = "cyan3", size = 2.5) +
  annotate(geom = "text", x=-87, y=-31, label = "Corcovado Gulf, \nChile",
color = "red", size = 2.5) +
  xlab("") + ylab("") +
  theme(panel.grid.major = element_line(color = "dark gray"),
```

```
panel.grid.minor = element_line(color = "dark gray"),
    axis.text = element_blank(), axis.ticks = element_blank(),
    panel.background = element_blank(),
    plot.background = element_rect(fill = "transparent", color = NA))
pacific.map
```





MORPHOLOGICAL COMPARISONS

Data wrangling

```
EE.df <- read.csv("./Photogrammetry/sapphire_EE.csv")
Fs.df <- read.csv("./Photogrammetry/sapphire_Fs.csv")
Fw.df <- read.csv("./Photogrammetry/sapphire_Fw.csv")
JL.df <- read.csv("./Photogrammetry/sapphire_JL.csv")
RB.df <- read.csv("./Photogrammetry/sapphire_RB.csv")
Tail.df <- read.csv("./Photogrammetry/sapphire_tail.csv")

morph_df.list <- list(EE.df, Fs.df, Fw.df, JL.df, RB.df, Tail.df)

morph_sample_sizes.df <- data.frame(matrix(ncol = 4, nrow = length(morph_df.list)))
colnames(morph_sample_sizes.df) <- c("measurement", "n_CA", "n_Chile", "n_NZ")

for (i in 1:length(morph_df.list)) {</pre>
```

```
morph i <- morph df.list[[i]]</pre>
  morph_sample_sizes.df[i,1] <- colnames(morph_i[4])</pre>
  morph sample sizes.df[i,2] <-
length(morph i$Location[morph i$Location=="CA"])
  morph_sample_sizes.df[i,3] <-</pre>
length(morph i$Location[morph i$Location=="Chile"])
  morph sample sizes.df[i,4] <-
length(morph_i$Location[morph_i$Location=="NZ"])
  morph_i$Location <- factor(morph_i$Location, levels = c("CA", "NZ",</pre>
"Chile"))
}
morph sample sizes.df
##
         measurement n_CA n_Chile n_NZ
## 1
         EE.std.mean
                           12
                                      10
                                            15
## 2
         Fs.std.mean
                           11
                                      13
                                            16
         Fw.std.mean
## 3
                           11
                                      15
                                            14
## 4
         JL.std.mean
                           12
                                            17
                                      11
## 5
         RB.std.mean
                           12
                                      14
                                            15
## 6 Tail.std.mean
                            6
                                      13
                                              6
# Re-order factor levels
EE.df$Location <- factor(EE.df$Location, levels = c("CA", "NZ", "Chile"))</pre>
Fs.df$Location <- factor(Fs.df$Location, levels = c("CA", "NZ", "Chile"))
Fw.df$Location <- factor(Fw.df$Location, levels = c("CA", "NZ", "Chile"))
JL.df$Location <- factor(JL.df$Location, levels = c("CA", "NZ", "Chile"))
RB.df$Location <- factor(RB.df$Location, levels = c("CA", "NZ", "Chile"))
Tail.df$Location <- factor(Tail.df$Location, levels = c("CA", "NZ", "Chile"))</pre>
```

Configure morphology plots

```
geom_segment(aes(x="CA", xend="Chile", y=0.152, yend=0.152)) +
  geom_text(aes(x="CA", y=0.15, label="*"), vjust=-0.005, hjust=-5, size=3.5)
  geom_text(aes(x="CA", y=0.152, label="*"), vjust=-0.005, hjust=-10,
size=3.5) +
  scale_color_manual(values = fillcols_a) + scale_fill_manual(values =
fillcols a) +
  ylab("Eye-to-eye") + xlab(" ") + guides(fill="none", color="none") +
  scale_x_discrete(labels=c("ENP", "New Zealand", "Chile")) +
 theme classic()
# EE.plot
JL.plot <- ggplot(data = JL.df) +</pre>
  geom_pointrange(aes(x=Location, y=JL.std.mean, ymin=JL.std.lower,
ymax=JL.std.upper, color = Location, fill = Location),
                  position = position jitter(), linetype='dashed', alpha =
0.7, size=0.1) +
  geom_boxplot(aes(y=JL.std.mean, x=Location, color = Location, fill =
Location), alpha = 0.7) +
  geom_segment(aes(x="CA", xend="NZ", y=0.2455, yend=0.2455)) +
  geom\_segment(aes(x="CA", xend="Chile", y=0.248, yend=0.248)) +
  geom_text(aes(x="CA", y=0.2455, label="*"), vjust=-0.005, hjust=-5,
size=3.5) +
  geom text(aes(x="CA", y=0.248, label="*"), vjust=-0.005, hjust=-10,
size=3.5) +
  scale color manual(values = fillcols a) + scale fill manual(values =
fillcols a) +
  ylab("Jaw length") + xlab(" ") + guides(fill="none", color="none") +
  scale_x_discrete(labels=c("ENP", "New Zealand", "Chile")) +
  theme classic()
# JL.plot
RB.plot <- ggplot(data = RB.df) +
  geom pointrange(aes(x=Location, y=RB.std.mean, ymin=RB.std.lower,
ymax=RB.std.upper, color = Location, fill = Location),
                  position = position_jitter(), linetype='dashed', alpha =
0.7, size=0.1) +
  geom boxplot(aes(y=RB.std.mean, x=Location, color = Location, fill =
Location), alpha = 0.7 +
  geom_segment(aes(x="CA", xend="NZ", y=0.227, yend=0.227)) +
  geom_segment(aes(x="CA", xend="Chile", y=0.23, yend=0.23)) +
  geom_text(aes(x="CA", y=0.227, label="*"), vjust=-0.005, hjust=-5,
size=3.5) +
  geom_text(aes(x="CA", y=0.23, label="*"), vjust=-0.005, hjust=-10,
size=3.5) +
```

```
scale_color_manual(values = fillcols_a) + scale_fill_manual(values =
fillcols a) +
  ylab("Rostrum-to-blowhole") + xlab(" ") + guides(fill="none", color="none")
  scale x discrete(labels=c("ENP", "New Zealand", "Chile")) +
  theme classic()
# RB.plot
Fs.plot <- ggplot(data = Fs.df) +
  geom pointrange(aes(x=Location, y=Fs.std.mean, ymin=Fs.std.lower,
ymax=Fs.std.upper, color = Location, fill = Location),
                  position = position_jitter(), linetype='dashed', alpha =
0.7, size=0.1) +
  geom_boxplot(aes(y=Fs.std.mean, x=Location, color = Location, fill =
Location), alpha = 0.7 +
  geom_segment(aes(x="CA", xend="NZ", y=0.271, yend=0.271)) +
  geom segment(aes(x="NZ", xend="Chile", y=0.275, yend=0.275)) +
  geom_text(aes(x="CA", y=0.271, label="*"), vjust=-0.005, hjust=-5,
size=3.5) +
  geom_text(aes(x="NZ", y=0.275, label="*"), vjust=-0.005, hjust=-5,
size=3.5) +
  scale_color_manual(values = fillcols_a) + scale_fill_manual(values =
fillcols a) +
  ylab("Fluke span") + xlab(" ") + guides(fill="none", color="none") +
  scale_x_discrete(labels=c("ENP", "New Zealand", "Chile")) +
  theme classic()
# Fs.plot
Fw.plot <- ggplot(data = Fw.df) +
  geom_pointrange(aes(x=Location, y=Fw.std.mean, ymin=Fw.std.lower,
ymax=Fw.std.upper, color = Location, fill = Location),
                  position = position jitter(), linetype='dashed', alpha =
0.7, size=0.1) +
  geom boxplot(aes(y=Fw.std.mean, x=Location, color = Location, fill =
Location), alpha = 0.7 +
  geom_segment(aes(x="CA", xend="NZ", y=0.074, yend=0.074)) +
  geom_text(aes(x="CA", y=0.074, label="*"), vjust=-0.005, hjust=-5,
size=3.5) +
  scale_color_manual(values = fillcols_a) + scale_fill_manual(values =
fillcols a) +
  ylab("Fluke width") + xlab(" ") + guides(fill="none", color="none") +
  scale_x_discrete(labels=c("ENP", "New Zealand", "Chile")) +
 theme_classic()
# Fw.plot
```

```
Tail.plot <- ggplot(data = Tail.df) +</pre>
  geom pointrange(aes(x=Location, y=Tail.std.mean, ymin=Tail.std.lower,
ymax=Tail.std.upper, color = Location, fill = Location),
                    position = position_jitter(), linetype='dashed', alpha =
0.7, size=0.1) +
  geom_boxplot(aes(y=Tail.std.mean, x=Location, color = Location, fill =
Location), alpha = 0.7 +
  geom_segment(aes(x="CA", xend="Chile", y=0.245, yend=0.245)) +
  geom_text(aes(x="CA", y=0.245, label="*"), vjust=-0.005, hjust=-10,
size=3.5) +
  scale_color_manual(values = fillcols_a) + scale_fill_manual(values =
fillcols a) +
  ylab("Tail length") + xlab(" ") + guides(fill="none", color="none") +
  scale_x_discrete(labels=c("ENP", "New Zealand", "Chile")) +
  theme_classic()
# Tail.plot
morph.multiplot_nopecs <- grid.arrange(EE.plot, JL.plot, RB.plot, Fs.plot,</pre>
Fw.plot, Tail.plot,
                                    nrow=2, ncol=3)
                                             0.23
0.21
0.19
0.17
0.15
   0.15
                          0.24
0.14
0.13
0.12
                       Jaw length
                          0.20
   0.11
        ⊟NA€W Zeal@mide
                               ⊟NAPw Zeal@mide
                                                     ⊟NMaPw Zeal@mide
                          0.07
   0.26
                                                0.23
                       Fluke width
Fluke span
                                              rail length
   0.24
   0.22
                                                0.19
                          0.04
   0.20
                          0.03
```

ggsave(morph.multiplot_nopecs, filename="./plots/morph.multiplot_v4.png",
device = "png", height=200, width=180, units="mm")

⊟NAPw Zeal@mide

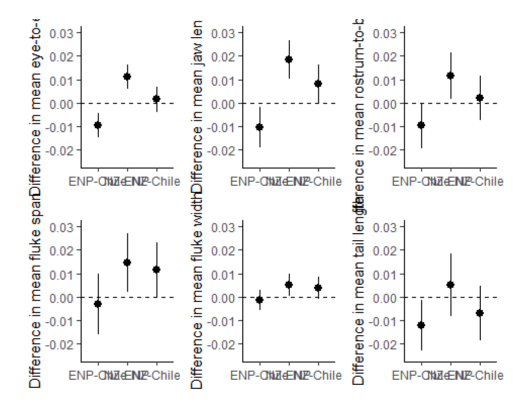
⊟NA€W Zeal@mide

⊟NMeW Zeal@midle

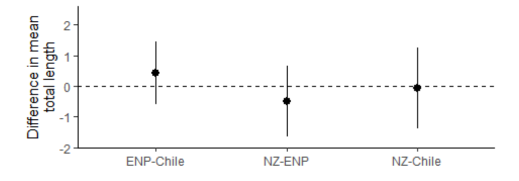
Plot MCMC ANOVA results

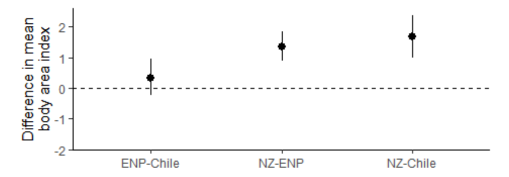
```
TL_anova.df <- read.csv("./Photogrammetry/MCMC ANOVA
outputs/TL summary results.csv")
BAI anova.df <- read.csv("./Photogrammetry/MCMC ANOVA
outputs/BAI summary results.csv")
EE anova.df <- read.csv("./Photogrammetry/MCMC ANOVA
outputs/EE summary results.csv")
Fs_anova.df <- read.csv("./Photogrammetry/MCMC ANOVA
outputs/Fs summary results.csv")
Fw_anova.df <- read.csv("./Photogrammetry/MCMC ANOVA</pre>
outputs/Fw_summary_results.csv")
JL_anova.df <- read.csv("./Photogrammetry/MCMC ANOVA</pre>
outputs/JL summary results.csv")
RB_anova.df <- read.csv("./Photogrammetry/MCMC ANOVA
outputs/RB summary results.csv")
Tail_anova.df <- read.csv("./Photogrammetry/MCMC ANOVA</pre>
outputs/Tail_summary_results.csv")
TL anova.plot <- ggplot(data = TL anova.df[!is.na(TL anova.df$label),]) +
  geom_pointrange(aes(x = label, y = means, ymin = HPD_lwr, ymax = HPD_upr))
  geom abline(intercept = 0, slope = 0, lty = 2) +
  ylim(-1.8, 2.4) +
  xlab("") + ylab("Difference in mean\ntotal length") +
  scale_x_discrete(labels=c("ENP-Chile", "NZ-ENP", "NZ-Chile")) +
  theme_classic()
BAI_anova.plot <- ggplot(data = BAI_anova.df[!is.na(BAI_anova.df$label),]) +
  geom_pointrange(aes(x = label, y = means, ymin = HPD_lwr, ymax = HPD_upr))
  geom abline(intercept = 0, slope = 0, lty = 2) +
  ylim(-1.8, 2.4) +
  xlab("") + ylab("Difference in mean\nbody area index") +
  scale_x_discrete(labels=c("ENP-Chile", "NZ-ENP", "NZ-Chile")) +
  theme classic()
EE_anova.plot <- ggplot(data = EE_anova.df[!is.na(EE_anova.df$label),]) +</pre>
  geom pointrange(aes(x = label, y = means, ymin = HPD lwr, ymax = HPD upr))
  geom_abline(intercept = 0, slope = 0, lty = 2) +
  vlim(-0.025, 0.03) +
  xlab("") + ylab("Difference in mean eye-to-eye") +
  scale_x_discrete(labels=c("ENP-Chile", "NZ-ENP", "NZ-Chile")) +
  theme classic()
JL anova.plot <- ggplot(data = JL anova.df[!is.na(JL anova.df$label),]) +</pre>
  geom_pointrange(aes(x = label, y = means, ymin = HPD_lwr, ymax = HPD_upr))
```

```
geom abline(intercept = 0, slope = 0, lty = 2) +
  ylim(-0.025, 0.03) +
  xlab("") + ylab("Difference in mean jaw length") +
  scale x discrete(labels=c("ENP-Chile", "NZ-ENP", "NZ-Chile")) +
  theme_classic()
RB_anova.plot <- ggplot(data = RB_anova.df[!is.na(RB_anova.df$label),]) +</pre>
  geom_pointrange(aes(x = label, y = means, ymin = HPD_lwr, ymax = HPD_upr))
  geom abline(intercept = 0, slope = 0, lty = 2) +
  ylim(-0.025, 0.03) +
  xlab("") + ylab("Difference in mean rostrum-to-blowhole") +
  scale_x_discrete(labels=c("ENP-Chile", "NZ-ENP", "NZ-Chile")) +
  theme classic()
Fs anova.plot <- ggplot(data = Fs anova.df[!is.na(Fs anova.df$label),]) +
  geom pointrange(aes(x = label, y = means, ymin = HPD lwr, ymax = HPD upr))
  geom_abline(intercept = 0, slope = 0, lty = 2) +
  vlim(-0.025, 0.03) +
  xlab("") + ylab("Difference in mean fluke span") +
  scale_x_discrete(labels=c("ENP-Chile", "NZ-ENP", "NZ-Chile")) +
  theme classic()
Fw anova.plot <- ggplot(data = Fw anova.df[!is.na(Fw anova.df$label),]) +
  geom_pointrange(aes(x = label, y = means, ymin = HPD_lwr, ymax = HPD_upr))
  geom abline(intercept = 0, slope = 0, lty = 2) +
  ylim(-0.025, 0.03) +
  xlab("") + ylab("Difference in mean fluke width")
  scale x discrete(labels=c("ENP-Chile", "NZ-ENP", "NZ-Chile")) +
  theme_classic()
Tail anova.plot <- ggplot(data = Tail anova.df[!is.na(Tail anova.df$label),])</pre>
  geom pointrange(aes(x = label, y = means, ymin = HPD lwr, ymax = HPD upr))
  geom_abline(intercept = 0, slope = 0, lty = 2) +
  ylim(-0.025, 0.03) +
  xlab("") + ylab("Difference in mean tail length") +
  scale_x_discrete(labels=c("ENP-Chile", "NZ-ENP", "NZ-Chile")) +
  theme classic()
morph.multiplot_anova <- grid.arrange(EE_anova.plot, JL_anova.plot,</pre>
RB anova.plot, Fs anova.plot, Fw anova.plot, Tail anova.plot,
                                nrow=2, ncol=3)
```



```
ggsave(morph.multiplot_anova,
filename="./plots/morph.multiplot_anovaresults_v1.png", device = "png",
height=200, width=180, units="mm")
TL_BAI_anova <- grid.arrange(TL_anova.plot, BAI_anova.plot, nrow=2, ncol=1)
```





```
ggsave(TL_BAI_anova, filename="./plots/TL_BAI.multiplot_anovaresults_v1.png",
device = "png", height=110, width=80, units="mm")
```

Allometry

```
EE_abs.df <- read.csv("./Photogrammetry/absolute morph</pre>
measurements/sapphire_EE_abs.csv")
Fs_abs.df <- read.csv("./Photogrammetry/absolute morph
measurements/sapphire Fs abs.csv")
Fw_abs.df <- read.csv("./Photogrammetry/absolute morph</pre>
measurements/sapphire_Fw_abs.csv")
JL_abs.df <- read.csv("./Photogrammetry/absolute morph</pre>
measurements/sapphire_JL_abs.csv")
RB abs.df <- read.csv("./Photogrammetry/absolute morph</pre>
measurements/sapphire RB abs.csv")
Tail_abs.df <- read.csv("./Photogrammetry/absolute morph</pre>
measurements/sapphire Tail abs.csv")
# Re-order factor levels
EE_abs.df$Location <- factor(EE_abs.df$Location, levels = c("CA", "NZ",</pre>
"Chile"))
Fs abs.df$Location <- factor(Fs abs.df$Location, levels = c("CA", "NZ",
"Chile"))
Fw_abs.df$Location <- factor(Fw_abs.df$Location, levels = c("CA", "NZ",
"Chile"))
JL_abs.df$Location <- factor(JL_abs.df$Location, levels = c("CA", "NZ",</pre>
```

```
"Chile"))
RB abs.df$Location <- factor(RB abs.df$Location, levels = c("CA", "NZ",
"Chile"))
Tail abs.df$Location <- factor(Tail abs.df$Location, levels = c("CA", "NZ",
"Chile"))
## Plot on log-log scale
log EE allometry.plot <- ggplot() +</pre>
  geom_point(data = EE_abs.df, aes(x = log(TL.mean), y = log(EE.mean), color
= Location)) +
  geom smooth(data = EE abs.df, aes(x = log(TL.mean), y = log(EE.mean), color
= Location, fill = Location), method = lm, alpha = 0.15, se=FALSE) +
  geom\_pointrange(data = EE\_abs.df, aes(x = log(TL.mean), y = log(EE.mean),
                                         color = Location, ymin =
log(EE.lower), ymax = log(EE.upper)), fatten = 0, alpha = 0.15) +
  geom pointrange(data = EE abs.df, aes(x = log(TL.mean), y = log(EE.mean),
                                         color = Location, xmin =
log(TL.lower), xmax = log(TL.upper)), fatten = 0, alpha = 0.15) +
  scale_color_manual(values = fillcols_a, labels=c("ENP", "New Zealand",
"Chile")) +
  scale fill manual(values = fillcols_a, labels=c("ENP", "New Zealand",
"Chile")) + labs(color="Population", fill="Population") +
  xlab("log(Total length)") + ylab("log(Eye-to-eye)") + xlim(2.7,3.4) +
  theme classic()
log Fw allometry.plot <- ggplot() +</pre>
  geom point(data = Fw abs.df, aes(x = log(TL.mean), y = log(Fw.mean), color
= Location)) +
  geom smooth(data = Fw abs.df, aes(x = log(TL.mean), y = log(Fw.mean), color
= Location, fill = Location), method = lm, alpha = 0.15, se=FALSE) +
  geom pointrange(data = Fw abs.df, aes(x = log(TL.mean), y = log(Fw.mean),
                                        color = Location, ymin =
log(Fw.lower), ymax = log(Fw.upper)), fatten = 0, alpha = 0.15) +
  geom\_pointrange(data = Fw\_abs.df, aes(x = log(TL.mean), y = log(Fw.mean),
                                         color = Location, xmin =
log(TL.lower), xmax = log(TL.upper)), fatten = 0, alpha = 0.15) +
  scale color manual(values = fillcols a, labels=c("ENP", "New Zealand",
"Chile")) +
  scale fill_manual(values = fillcols_a, labels=c("ENP", "New Zealand",
"Chile")) + labs(color="Population", fill="Population") +
  xlab("log(Total length)") + ylab("log(Fluke width)") + xlim(2.7,3.4) +
  theme classic()
log Fs allometry.plot <- ggplot() +</pre>
  geom_point(data = Fs_abs.df, aes(x = log(TL.mean), y = log(Fs.mean), color
= Location)) +
  geom\_smooth(data = Fs\_abs.df, aes(x = log(TL.mean), y = log(Fs.mean), color
= Location, fill = Location), method = lm, alpha = 0.15, se=FALSE) +
```

```
geom pointrange(data = Fs abs.df, aes(x = log(TL.mean), y = log(Fs.mean),
                                        color = Location, ymin =
log(Fs.lower), ymax = log(Fs.upper)), fatten = 0, alpha = 0.15) +
  geom_pointrange(data = Fs_abs.df, aes(x = log(TL.mean), y = log(Fs.mean),
                                        color = Location, xmin =
log(TL.lower), xmax = log(TL.upper)), fatten = 0, alpha = 0.15) +
  scale_color_manual(values = fillcols_a, labels=c("ENP", "New Zealand",
"Chile")) +
  scale fill manual(values = fillcols_a, labels=c("ENP", "New Zealand",
"Chile")) + labs(color="Population", fill="Population") +
  xlab("log(Total length)") + ylab("log(Fluke span)") + xlim(2.7,3.4) +
  theme classic()
log JL allometry.plot <- ggplot() +</pre>
  geom point(data = JL abs.df, aes(x = log(TL.mean), y = log(JL.mean), color
= Location)) +
  geom\_smooth(data = JL\_abs.df, aes(x = log(TL.mean), y = log(JL.mean), color
= Location, fill = Location), method = lm, alpha = 0.15, se=FALSE) +
  geom\_pointrange(data = JL\_abs.df, aes(x = log(TL.mean), y = log(JL.mean),
                                        color = Location, ymin =
log(JL.lower), ymax = log(JL.upper)), fatten = 0, alpha = 0.15) +
  geom pointrange(data = JL abs.df, aes(x = log(TL.mean), y = log(JL.mean),
                                        color = Location, xmin =
log(TL.lower), xmax = log(TL.upper)), fatten = 0, alpha = 0.15) +
  scale_color_manual(values = fillcols_a, labels=c("ENP", "New Zealand",
"Chile")) +
  scale fill manual(values = fillcols a, labels=c("ENP", "New Zealand",
"Chile")) + labs(color="Population", fill="Population") +
  xlab("log(Total length)") + ylab("log(Jaw length)") + xlim(2.7,3.4) +
  theme classic()
log RB allometry.plot <- ggplot() +</pre>
  geom_point(data = RB_abs.df, aes(x = log(TL.mean), y = log(RB.mean), color
= Location)) +
  geom smooth(data = RB abs.df, aes(x = log(TL.mean), y = log(RB.mean), color
= Location, fill = Location), method = lm, alpha = 0.15, se=FALSE) +
  geom\_pointrange(data = RB\_abs.df, aes(x = log(TL.mean), y = log(RB.mean),
                                        color = Location, ymin =
log(RB.lower), ymax = log(RB.upper)), fatten = 0, alpha = 0.15) +
  geom pointrange(data = RB abs.df, aes(x = log(TL.mean), y = log(RB.mean),
                                        color = Location, xmin =
log(TL.lower), xmax = log(TL.upper)), fatten = 0, alpha = 0.15) +
  scale_color_manual(values = fillcols_a, labels=c("ENP", "New Zealand",
"Chile")) +
  scale fill manual(values = fillcols a, labels=c("ENP", "New Zealand",
"Chile")) + labs(color="Population", fill="Population") +
  xlab("log(Total length)") + ylab("log(Rostrum-to-blowhole)") +
xlim(2.7,3.4) +
  theme classic()
```

```
log Tail allometry.plot <- ggplot() +</pre>
  geom point(data = Tail abs.df, aes(x = log(TL.mean), y = log(Tail.mean),
color = Location)) +
  geom_smooth(data = Tail_abs.df, aes(x = log(TL.mean), y = log(Tail.mean),
color = Location, fill = Location), method = lm, alpha = 0.15, se=FALSE) +
  geom_pointrange(data = Tail_abs.df, aes(x = log(TL.mean), y =
log(Tail.mean),
                                          color = Location, ymin =
log(Tail.lower), ymax = log(Tail.upper)), fatten = 0,alpha = 0.15)+
  geom_pointrange(data = Tail_abs.df, aes(x = log(TL.mean), y =
log(Tail.mean),
                                          color = Location, xmin =
log(TL.lower), xmax = log(TL.upper)), fatten = 0, alpha = 0.15) +
  scale color manual(values = fillcols a, labels=c("ENP", "New Zealand",
"Chile")) +
  scale fill manual(values = fillcols a, labels=c("ENP", "New Zealand",
"Chile")) + labs(color="Population", fill="Population") +
  xlab("log(Total length)") + ylab("log(Tail length)") + xlim(2.7,3.4) +
  theme classic()
allometry.multiplot<- ggarrange(log_EE_allometry.plot, log_JL_allometry.plot,
log_RB_allometry.plot, log_Fs_allometry.plot, log_Fw_allometry.plot,
log Tail allometry.plot, nrow=2, ncol=3, common.legend = TRUE,
legend="bottom")
ggsave(allometry.multiplot, filename="./plots/allometry.multiplot_v4.png",
device = "png", height=200, width=180, units="mm")
## Linear models on log-log scale to get slopes
# EE
summary(lm(log(EE.mean) ~ log(TL.mean), data =
EE_abs.df[EE_abs.df$Location=="CA",]))
##
## Call:
## lm(formula = log(EE.mean) ~ log(TL.mean), data =
EE abs.df[EE abs.df$Location ==
       "CA", ])
##
##
## Residuals:
##
        Min
                  1Q
                       Median
                                    3Q
                                            Max
## -0.10018 -0.02869 0.01335 0.03251 0.05925
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
##
                             0.5655 -3.891 0.003003 **
## (Intercept) -2.2005
## log(TL.mean)
                 1.0272
                             0.1815
                                     5.661 0.000209 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```
##
## Residual standard error: 0.04901 on 10 degrees of freedom
## Multiple R-squared: 0.7622, Adjusted R-squared: 0.7384
## F-statistic: 32.05 on 1 and 10 DF, p-value: 0.0002093
summary(lm(log(EE.mean) ~ log(TL.mean), data =
EE_abs.df[EE_abs.df$Location=="NZ",]))
##
## Call:
## lm(formula = log(EE.mean) ~ log(TL.mean), data =
EE_abs.df[EE_abs.df$Location ==
##
       "NZ", ])
##
## Residuals:
        Min
                    10
                          Median
                                        30
                                                 Max
## -0.056850 -0.013036 0.004618 0.012006 0.044951
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
                            0.25085 -7.231 6.63e-06 ***
## (Intercept) -1.81401
                            0.08086 11.518 3.41e-08 ***
## log(TL.mean) 0.93133
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.02614 on 13 degrees of freedom
## Multiple R-squared: 0.9108, Adjusted R-squared: 0.9039
## F-statistic: 132.7 on 1 and 13 DF, p-value: 3.411e-08
summary(lm(log(EE.mean) ~ log(TL.mean), data =
EE abs.df[EE abs.df$Location=="Chile",]))
##
## Call:
## lm(formula = log(EE.mean) ~ log(TL.mean), data =
EE abs.df[EE abs.df$Location ==
##
       "Chile", ])
##
## Residuals:
                       Median
        Min
                  1Q
                                    3Q
                                            Max
## -0.05772 -0.01856 -0.01133 0.01879
                                        0.08439
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
                 -2.1442
                             1.3173
                                    -1.628
## (Intercept)
                                              0.1422
## log(TL.mean)
                 1.0341
                             0.4253
                                      2.431
                                              0.0411 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.0455 on 8 degrees of freedom
```

```
## Multiple R-squared: 0.4249, Adjusted R-squared: 0.353
## F-statistic: 5.911 on 1 and 8 DF, p-value: 0.04112
# JL
summary(lm(log(JL.mean) ~ log(TL.mean), data =
JL abs.df[JL abs.df$Location=="CA",]))
##
## Call:
## lm(formula = log(JL.mean) ~ log(TL.mean), data =
JL_abs.df[JL_abs.df$Location ==
       "CA", ])
##
##
## Residuals:
                          Median
                    1Q
                                        3Q
                                                 Max
## -0.047292 -0.031999 0.001039 0.026694
                                           0.077323
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                 -1.9947
                             0.4656 -4.284
                                             0.0016 **
## log(TL.mean)
                 1.1223
                             0.1494
                                     7.512 2.04e-05 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.04035 on 10 degrees of freedom
## Multiple R-squared: 0.8495, Adjusted R-squared: 0.8344
## F-statistic: 56.43 on 1 and 10 DF, p-value: 2.035e-05
summary(lm(log(JL.mean) ~ log(TL.mean), data =
JL_abs.df[JL_abs.df$Location=="NZ",]))
##
## Call:
## lm(formula = log(JL.mean) ~ log(TL.mean), data =
JL_abs.df[JL_abs.df$Location ==
##
      "NZ", ])
##
## Residuals:
##
        Min
                  1Q
                      Median
                                    3Q
                                            Max
## -0.07820 -0.04969 0.01382 0.02363 0.07333
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
                                    -3.646 0.00239 **
## (Intercept)
                 -1.5485
                             0.4247
                                     7.327 2.5e-06 ***
## log(TL.mean)
                 1.0073
                             0.1375
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.04942 on 15 degrees of freedom
## Multiple R-squared: 0.7816, Adjusted R-squared: 0.767
## F-statistic: 53.68 on 1 and 15 DF, p-value: 2.499e-06
```

```
summary(lm(log(JL.mean) ~ log(TL.mean), data =
JL_abs.df[JL_abs.df$Location=="Chile",]))
##
## Call:
## lm(formula = log(JL.mean) ~ log(TL.mean), data =
JL_abs.df[JL_abs.df$Location ==
       "Chile", ])
##
## Residuals:
                                        30
                                                 Max
##
         Min
                    10
                          Median
## -0.064848 -0.029084 -0.007023 0.015020 0.082056
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                 -3.5648
                             1.1701
                                    -3.047 0.01387 *
                                      4.354
## log(TL.mean)
                  1.6474
                             0.3784
                                             0.00184 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.04468 on 9 degrees of freedom
## Multiple R-squared: 0.6781, Adjusted R-squared: 0.6423
## F-statistic: 18.96 on 1 and 9 DF, p-value: 0.00184
# RB
summary(lm(log(RB.mean) ~ log(TL.mean), data =
RB abs.df[RB abs.df$Location=="CA",]))
##
## Call:
## lm(formula = log(RB.mean) ~ log(TL.mean), data =
RB abs.df[RB abs.df$Location ==
       "CA", ])
##
##
## Residuals:
        Min
                    10
                          Median
                                        30
                                                 Max
## -0.105251 -0.031721 -0.006171 0.031356 0.112717
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
                                    -2.725 0.021361 *
## (Intercept)
                             0.6788
                 -1.8501
                                      4.756 0.000774 ***
## log(TL.mean)
                  1.0359
                             0.2178
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.05883 on 10 degrees of freedom
## Multiple R-squared: 0.6934, Adjusted R-squared: 0.6627
## F-statistic: 22.62 on 1 and 10 DF, p-value: 0.0007735
summary(lm(log(RB.mean) ~ log(TL.mean), data =
RB_abs.df[RB_abs.df$Location=="NZ",]))
```

```
##
## Call:
## lm(formula = log(RB.mean) ~ log(TL.mean), data =
RB abs.df[RB abs.df$Location ==
       "NZ", ])
##
##
## Residuals:
        Min
                 10
                      Median
                                    30
                                            Max
## -0.12220 -0.06053 0.02727 0.04456
                                        0.12395
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                 -2.9437
                             0.7978 -3.690 0.002723 **
                             0.2602
## log(TL.mean)
                 1.4138
                                     5.433 0.000115 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.0742 on 13 degrees of freedom
## Multiple R-squared: 0.6942, Adjusted R-squared: 0.6707
## F-statistic: 29.52 on 1 and 13 DF, p-value: 0.0001145
summary(lm(log(RB.mean) ~ log(TL.mean), data =
RB_abs.df[RB_abs.df$Location=="Chile",]))
##
## Call:
## lm(formula = log(RB.mean) ~ log(TL.mean), data =
RB abs.df[RB abs.df$Location ==
##
      "Chile", ])
##
## Residuals:
##
        Min
                    10
                          Median
                                        3Q
                                                 Max
## -0.111575 -0.036991 0.009874 0.032526 0.093451
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                 -2.1555
                             1.2530
                                      -1.72
                                              0.1110
                             0.4058
                                       2.84
                                              0.0149 *
## log(TL.mean)
                 1.1527
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.06053 on 12 degrees of freedom
## Multiple R-squared: 0.402, Adjusted R-squared: 0.3522
## F-statistic: 8.068 on 1 and 12 DF, p-value: 0.01488
summary(lm(log(Fw.mean) ~ log(TL.mean), data =
Fw_abs.df[Fw_abs.df$Location=="CA",]))
##
## Call:
```

```
## lm(formula = log(Fw.mean) ~ log(TL.mean), data =
Fw abs.df[Fw abs.df$Location ==
       "CA", ])
##
##
## Residuals:
##
         Min
                          Median
                                        30
                                                 Max
                    1Q
## -0.113352 -0.042860
                        0.005397 0.047801
                                            0.098603
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
                                    -4.054 0.00287 **
## (Intercept)
                 -3.3025
                             0.8147
                                      4.176 0.00239 **
## log(TL.mean)
                  1.0923
                             0.2615
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.07046 on 9 degrees of freedom
## Multiple R-squared: 0.6596, Adjusted R-squared: 0.6218
## F-statistic: 17.44 on 1 and 9 DF, p-value: 0.00239
summary(lm(log(Fw.mean) ~ log(TL.mean), data =
Fw abs.df[Fw abs.df$Location=="NZ",]))
##
## Call:
## lm(formula = log(Fw.mean) ~ log(TL.mean), data =
Fw_abs.df[Fw_abs.df$Location ==
       "NZ", ])
##
##
## Residuals:
        Min
                  1Q
                       Median
                                    3Q
                                            Max
## -0.20616 -0.06069 -0.01284 0.06375
                                        0.16121
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
                             1.2423 -1.800
## (Intercept)
                 -2.2358
                                              0.0971 .
                                      1.942
                                              0.0760 .
## log(TL.mean)
                  0.7788
                             0.4011
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.1021 on 12 degrees of freedom
## Multiple R-squared: 0.239, Adjusted R-squared:
## F-statistic: 3.77 on 1 and 12 DF, p-value: 0.07603
summary(lm(log(Fw.mean) ~ log(TL.mean), data =
Fw_abs.df[Fw_abs.df$Location=="Chile",]))
##
## Call:
## lm(formula = log(Fw.mean) ~ log(TL.mean), data =
Fw_abs.df[Fw_abs.df$Location ==
       "Chile", ])
##
```

```
##
## Residuals:
##
        Min
                  10
                       Median
                                    3Q
                                            Max
## -0.10541 -0.05508 0.01553 0.05697
                                        0.13929
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
                                    -2.447
## (Intercept)
                 -3.7605
                             1.5366
                                              0.0294 *
                             0.4977
                                      2.512
                                              0.0260 *
## log(TL.mean)
                  1.2502
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.07424 on 13 degrees of freedom
## Multiple R-squared: 0.3268, Adjusted R-squared: 0.275
## F-statistic: 6.31 on 1 and 13 DF, p-value: 0.02599
# Fs
summary(lm(log(Fs.mean) ~ log(TL.mean), data =
Fs_abs.df[Fs_abs.df$Location=="CA",]))
##
## Call:
## lm(formula = log(Fs.mean) ~ log(TL.mean), data =
Fs abs.df[Fs abs.df$Location ==
##
       "CA", 1)
##
## Residuals:
                       Median
        Min
                  1Q
                                    3Q
                                            Max
## -0.11607 -0.06030 -0.01901 0.06721 0.12543
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                 -0.7240
                             1.0459
                                    -0.692
                                              0.5062
## log(TL.mean)
                  0.7374
                             0.3358
                                      2.196
                                              0.0557 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.09045 on 9 degrees of freedom
## Multiple R-squared: 0.3489, Adjusted R-squared: 0.2766
## F-statistic: 4.823 on 1 and 9 DF, p-value: 0.05569
summary(lm(log(Fs.mean) ~ log(TL.mean), data =
Fs abs.df[Fs abs.df$Location=="NZ",]))
##
## Call:
## lm(formula = log(Fs.mean) ~ log(TL.mean), data =
Fs abs.df[Fs abs.df$Location ==
       "NZ", ])
##
##
## Residuals:
```

```
Min 10
                       Median
                                    30
                                            Max
## -0.11016 -0.01219 0.01087 0.02014
                                        0.10822
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
                                    -1.408 0.181000
## (Intercept)
                 -0.6485
                             0.4606
## log(TL.mean)
                  0.7327
                             0.1492
                                      4.912 0.000229 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.05064 on 14 degrees of freedom
## Multiple R-squared: 0.6328, Adjusted R-squared: 0.6065
## F-statistic: 24.12 on 1 and 14 DF, p-value: 0.0002293
summary(lm(log(Fs.mean) ~ log(TL.mean), data =
Fs_abs.df[Fs_abs.df$Location=="Chile",]))
##
## Call:
## lm(formula = log(Fs.mean) \sim log(TL.mean), data =
Fs_abs.df[Fs_abs.df$Location ==
##
       "Chile", ])
##
## Residuals:
                          Median
         Min
                    10
                                        30
                                                 Max
## -0.095616 -0.025451 -0.002776 0.012743 0.163422
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
##
                 -3.0183
                             1.4788
                                     -2.041
## (Intercept)
                                              0.0660 .
## log(TL.mean)
                  1.4845
                             0.4799
                                      3.094
                                              0.0102 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.06545 on 11 degrees of freedom
## Multiple R-squared: 0.4652, Adjusted R-squared:
## F-statistic: 9.57 on 1 and 11 DF, p-value: 0.01022
# Tail
summary(lm(log(Tail.mean) ~ log(TL.mean), data =
Tail_abs.df[Tail_abs.df$Location=="CA",]))
##
## Call:
## lm(formula = log(Tail.mean) ~ log(TL.mean), data =
Tail abs.df[Tail abs.df$Location ==
##
       "CA", ])
##
## Residuals:
                                                        5
##
                                  3
                                             4
                                                                   6
            1
                       2
## -0.0001675 -0.0084285 0.0262343 -0.0481266 0.0201263 0.0103620
```

```
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
                             0.6378
                                    -2.178
## (Intercept)
                 -1.3888
                                              0.0950 .
                                      4.593
## log(TL.mean)
                  0.9423
                             0.2051
                                              0.0101 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.02995 on 4 degrees of freedom
## Multiple R-squared: 0.8406, Adjusted R-squared: 0.8008
## F-statistic: 21.1 on 1 and 4 DF, p-value: 0.01008
summary(lm(log(Tail.mean) ~ log(TL.mean), data =
Tail abs.df[Tail abs.df$Location=="NZ",]))
##
## Call:
## lm(formula = log(Tail.mean) ~ log(TL.mean), data =
Tail_abs.df[Tail_abs.df$Location ==
##
       "NZ", ])
##
## Residuals:
##
          20
                    21
                              22
                                        23
                                                  24
                        0.034276 0.021668 -0.129720
##
    0.035858 0.035001
                                                      0.002917
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                 -0.5832
                             1.2881
                                     -0.453
                                               0.674
## log(TL.mean)
                  0.6875
                             0.4182
                                      1.644
                                               0.176
##
## Residual standard error: 0.07244 on 4 degrees of freedom
## Multiple R-squared: 0.4032, Adjusted R-squared: 0.254
## F-statistic: 2.703 on 1 and 4 DF, p-value: 0.1755
summary(lm(log(Tail.mean) ~ log(TL.mean), data =
Tail abs.df[Tail abs.df$Location=="Chile",]))
##
## Call:
## lm(formula = log(Tail.mean) ~ log(TL.mean), data =
Tail_abs.df[Tail_abs.df$Location ==
       "Chile", ])
##
##
## Residuals:
##
         Min
                    10
                          Median
                                        3Q
                                                 Max
## -0.065579 -0.027709 -0.008747 0.016876 0.086271
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                 -2.4553
                             1.0146
                                    -2.420 0.03400 *
                                     3.972 0.00219 **
## log(TL.mean) 1.3059
                             0.3288
```

```
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.04828 on 11 degrees of freedom
## Multiple R-squared: 0.5892, Adjusted R-squared: 0.5518
## F-statistic: 15.77 on 1 and 11 DF, p-value: 0.00219
```

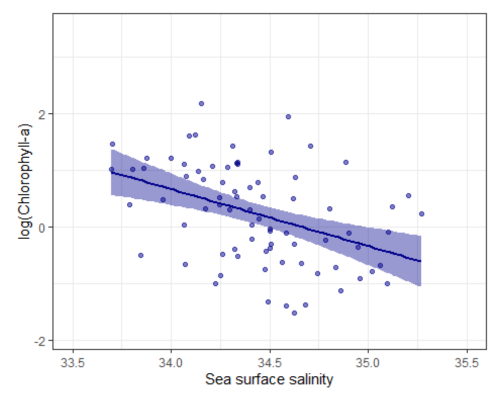
Salinity and Chlorophyll-a in Chile

Influence of freshwater input on productivity

```
GC_oc.df <- read.csv("GC_oc.csv")

## SSS and Chl-a

SSSvChla.p <- ggplot(data = GC_oc.df, aes(y=log(Chla), x=SSS)) +
    geom_point(color = "dark blue", alpha = 0.5) + geom_smooth(method = lm,
color = "dark blue", fill = "dark blue") +
    theme_bw() + xlim(33.5, 35.5) + xlab("Sea surface salinity") +
ylab("log(Chlorophyll-a)")</pre>
SSSvChla.p
```



```
ggsave(SSSvChla.p, filename="./plots/SSSvChla.png", device = "png",
height=80, width=100, units="mm")
```

```
# Pearson's correlation
cor.test(x = GC_oc.df$SSS, y = log(GC_oc.df$Chla), method = "pearson")
##
## Pearson's product-moment correlation
##
## data: GC_oc.df$SSS and log(GC_oc.df$Chla)
## t = -4.0397, df = 76, p-value = 0.0001268
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## -0.5879630 -0.2183404
## sample estimates:
## cor
## -0.4204429
```