SEACAR Oyster Analysis

Last compiled on 29 January, 2025

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Important Notes

The purpose of this script is to group oyster data, create managed area statistics, perform line analysis on density, percent live, and shell height, and create reports in pdf and Word document form for Oyster data.

All scripts and outputs can be found on the SEACAR GitHub repository:

```
https://github.com/FloridaSEACAR/SEACAR Trend Analyses
```

This markdown file is designed to be compiled by Oyster_ReportRender.R (https://github.com/FloridaSE ACAR/SEACAR_Trend_Analyses/blob/main/Oyster/Oyster_ReportRender.R).

This script is based off of code originally written by Stephen Durham.

Compiled and edited by J.E. Panzik and Tyler Hill for SEACAR.

Libraries and Settings

Loads libraries used in the script. Loads the Segoe UI font for use in the figures. The inclusion of scipen option limits how frequently R defaults to scientific notation. Sets default settings for displaying warning and messages in created document, and sets figure dpi.

```
library(lubridate)
library(tidyverse)
library(data.table)
library(glue)
library(glue)
library(kableExtra)
options(scipen=999)
knitr::opts_chunk$set(
   warning=FALSE,
   message=FALSE,
   dpi=200,
   fig.pos = 'H'
)
options(kableExtra.auto_format = FALSE)
options(knitr.kable.NA = '-')
```

File Import

Imports file that is determined in the Oyster ReportRender.R script.

The command fread is used because of its improved speed while handling large data files. Updates column names.

The latest version of Oyster data is available at: https://usf.box.com/s/6dtcrcdkrtc33fqw5kl5fjo6guo0anrw

The file being used for the analysis is: All_OYSTER_Parameters-2024-Dec-08.txt

Generate Universal Reef IDs

Individual IDs are given to reefs based on location within GIS files. This process now occurs separately from the Oyster habitat scripts and the UniversalReefID will be maintained within the database.

Data Setup & Filtering

Documentation on database filtering is provided here: SEACAR Documentation- Analysis Filters and Calculations.docx

Identifies and removes outliers in the data and various programs.

```
#Make sure column formats are correct-I am still getting an "NAs introduced
#by coercion" warning on the LiveDate calculation,
#but I'm not sure what is going on because when I spot-check the output,
#it does not look like it is introducing NAs...
oysterraw[, `:=` (RowID=as.integer(RowID),
                  ProgramID=as.integer(ProgramID),
                  LocationID=as.integer(LocationID),
                  ProgramName=as.character(ProgramName),
                  ProgramLocationID=as.character(ProgramLocationID),
                  QuadIdentifier=as.character(QuadIdentifier),
                  ReefIdentifier=as.character(ReefIdentifier),
                  UniversalReefID=as.factor(UniversalReefID),
                  LiveDate=as.integer(ifelse(!is.na(LiveDate_Qualifier) &
                                               str_detect(LiveDate,
                                                          "...-.."),
                                             paste0(str sub(LiveDate, 1, 4)),
```

```
round(as.numeric(LiveDate)))),
                  LiveDate_Qualifier=as.character(LiveDate_Qualifier),
                  LiveDate_MinEstDate=as.numeric(LiveDate_MinEstDate),
                  LiveDate MaxEstDate=as.numeric(LiveDate MaxEstDate),
                  SampleAge_Stdev=as.numeric(SampleAge_Stdev),
                  #GISUniqueID=as.logical(GISUniqueID),
                  Year=as.integer(Year),
                  Month=as.integer(Month),
                  ManagedAreaName=as.character(ManagedAreaName),
                  SurveyMethod=as.character(SurveyMethod),
                  PercentLiveMethod=as.character(PercentLiveMethod),
                  HabitatClassification=as.character(HabitatClassification),
                  MinimumSizeMeasured_mm=as.character(MinimumSizeMeasured_mm),
                  NumberMeasured n=as.character(NumberMeasured n),
                  QuadSize_m2=as.numeric(QuadSize_m2),
                  MADup=as.integer(MADup),
                  Density_m2=as.numeric(Density_m2),
                  PercentLive_pct=as.numeric(PercentLive_pct),
                  ShellHeight_mm=as.numeric(ShellHeight_mm),
                  Number_of_Oysters_Counted_Total_Count =
                    as.integer(Number_of_Oysters_Counted_Total_Count),
                  Number_of_Oysters_Counted_Live_Count =
                    as.integer(Number_of_Oysters_Counted_Live_Count),
                  Number_of_Oysters_Counted_Dead_Count =
                    as.integer(Number of Oysters Counted Dead Count),
                  ObsIndex=as.integer(ObsIndex))]
#Calculate Density_m2 values for ProgramID==4016 & 4042
oysterraw[ProgramID==4016, Density m2 :=
            Number_of_Oysters_Counted_Live_Count/as.numeric(QuadSize_m2)]
oysterraw[ProgramID==4042 & !is.na(Number_of_Oysters_Counted_Live_Count),
          Density m2 :=
            Number_of_Oysters_Counted_Live_Count/as.numeric(QuadSize_m2)]
#Remove "25" values from total counts column, make all "PercentLiveMethod"
#values the same, and calculate estimated live Density for ProgramID==5074 and
oysterraw <- oysterraw[RowID %in%</pre>
                         setdiff(
                           oysterraw[, RowID],
                           oysterraw[ProgramID ==5074 &
                                       Number_of_Oysters_Counted_Total_Count==25, RowID]), ]
oysterraw[ProgramID==5074, PercentLiveMethod := "Estimated percent"]
oysterraw[ProgramID==5074, SampleDate :=
            unique(oysterraw[ProgramID==5074 &
                               !is.na(Number_of_Oysters_Counted_Total_Count),
                             SampleDate])[1]]
#Some PercentLiveMethod values for ID4042 are NA
oysterraw[ProgramID==4042 | ProgramID==4016,
          PercentLiveMethod := "Point-intercept"]
#make sure quadrat identifiers are unique
oysterraw[, QuadIdentifier_old := QuadIdentifier]
```

```
oysterraw[, QuadIdentifier := paste(UniversalReefID,
                                    LocationID, Year, Month,
                                    QuadIdentifier_old, sep="_")]
oysterraw[, MA_plotlab := paste0(ManagedAreaName, "_", HabitatClassification)]
subtidal <- c(4044, 5007, 5071, 5073)</pre>
oysterraw[, Subtidal := ifelse(ProgramID %in% subtidal, 1, 0)][, Subtidal := as.logical(Subtidal)]
#Create variables for relative year and size class category for data that
#should be included in analyses and counts of live oysters measured
for(i in unique(oysterraw$ManagedAreaName)){
  oysterraw[ManagedAreaName==i & !is.na(LiveDate), `:=`
            (RelYear=(LiveDate-min(LiveDate))+1,
              YearDiff=min(LiveDate)-1,
              #adding 1 to each RelYear to avoid min(RelYear) == 0,
              #because it is used later as an index for plotting years so
              #it needs to start from 1
              SizeClass=fcase(ShellHeight_mm >= 25 &
                                ShellHeight_mm < 75, "25to75mm",
                              ShellHeight_mm >= 75, "o75mm",
                              default=NA))]
  oysterraw[ManagedAreaName==i & !is.na(LiveDate),
            counts := length(ShellHeight_mm), by=c("QuadIdentifier")]
}
#Remove unrealistically high shell heights from ID 5017
oysterraw <- setdiff(oysterraw, oysterraw[ProgramID==5017 & ShellHeight_mm >= 165, ])
#Create data table to save model results
oysterresults <- data.table(indicator=character(),</pre>
                            managed_area=character(),
                            habitat_class=character(),
                            size_class=character(),
                            live_date_qual=character(),
                            n_programs=integer(),
                            programs=list(),
                            filename=character(),
                            effect=character(),
                            component=character(),
                            group=character(),
                            term=character(),
                            estimate=numeric(),
                            std.error=numeric().
                            conf.low=numeric(),
                            conf.high=numeric())
#How many years of data for each managed area/habitat class/indicator combination?
setDT(oysterraw)
oysterraw[!is.na(Density_m2), `:=` (nyrpar="Density_m2",
                                    nyears=length(unique(Year))),
          by=MA_plotlab]
oysterraw[!is.na(PercentLive_pct), `:=` (nyrpar="PercentLive_pct",
```

```
nyears=length(unique(Year))),
          by=MA_plotlab]
oysterraw[!is.na(ShellHeight_mm), `:=` (nyrpar="ShellHeight_mm",
                                         nyears=length(unique(Year))),
          by=MA_plotlab]
MAinclude <- distinct(oysterraw[, .(MA_plotlab, nyrpar, nyears)])</pre>
# View(MAinclude[!is.na(nyrpar) & nyears >= 5, ])
oysterraw[str detect(MA plotlab, "Pine Island Sound"),
           `:=` (MA plotlab=ifelse(str detect(ProgramLocationID,
                                               "Reference")
                                      str_detect(ProgramLocationID,
                                                 "Control"),
                                    "Pine Island Sound Aquatic Preserve_Natural",
                                    "Pine Island Sound Aquatic Preserve_Restored"),
                 HabitatClassification=ifelse(str_detect(ProgramLocationID,
                                                          "Reference") |
                                                 str_detect(ProgramLocationID,
                                                            "Control"),
                                               "Natural", "Restored"))]
```

Managed Area Statistics

Gets summary statistics for each managed area. Uses piping from dplyr package to feed into subsequent steps. Sets of summary statistics are performed for Density, Shell Height, and Percent Living. The following steps are performed:

- 1. Group data that have the same ManagedAreaName, Year, Month, nyrpar, LiveDate_Qualifier, SizeClass, and HabitatClassification.
 - Second summary statistics do not use the Month grouping and are only for ManagedAreaName, Year, and the other oyster parameters.
 - Third summary statistics do not use Year grouping and are only for ManagedAreaName, Month, and the other oyster parameters.
 - Fourth summary statistics are only grouped based on ManagedAreaName and the other oyster parameters
 - Determines the years that the minimum and maximum species richness occurred
- 2. For each group, provide the following information: Parameter Name (ParameterName), Number of Entries (N_Data), Lowest Value (Min), Largest Value (Max), Median, Mean, Standard Deviation, and a list of all Programs included in these measurements.
- 3. Sort the data in ascending (A to Z and 0 to 9) order based on ManagedAreaName then Year then Month
- 4. Write summary stats to a pipe-delimited .txt file in the output directory
 - Oyster Output Files in SEACAR GitHub (https://github.com/FloridaSEACAR/SEACAR_Tren d_Analyses/tree/main/Oyster/output)

##Density

```
oysterraw$SizeClass[oysterraw$SizeClass=="25to75mm"] <- "25-75mm"
oysterraw$SizeClass[oysterraw$SizeClass=="35to75mm"] <- "35-75mm"
oysterraw$SizeClass[oysterraw$SizeClass=="075mm"] <- ">75mm"

# Create summary statistics for each managed area based on Year and Month
# intervals.

MA_YM_Stats <- oysterraw[oysterraw$nyrpar=="Density_m2",] %>%
group_by(AreaID, ManagedAreaName, Year, Month, nyrpar,
```

```
LiveDate_Qualifier, SizeClass, HabitatClassification) %>%
   dplyr::summarize(N Data=length(Density m2[!is.na(Density m2)]),
             Min=min(Density_m2[!is.na(Density_m2)]),
             Max=max(Density_m2[!is.na(Density_m2)]),
             Median=median(Density_m2[!is.na(Density_m2)]),
             Mean=mean(Density_m2[!is.na(Density_m2)]),
             StandardDeviation=sd(Density_m2[!is.na(Density_m2)]),
             Programs=paste(sort(unique(ProgramName), decreasing=FALSE),
                              collapse=', '),
             ProgramIDs=paste(sort(unique(ProgramID), decreasing=FALSE),
                              collapse=', '))
setnames(MA_YM_Stats, c("nyrpar", "LiveDate_Qualifier",
                        "HabitatClassification"),
         c("ParameterName", "ShellType", "HabitatType"))
MA_YM_Stats$ShellType[MA_YM_Stats$ShellType=="Exact"] <- "Live Oyster Shells"
MA_YM_Stats$ShellType[MA_YM_Stats$ShellType=="Estimate"] <- "Dead Oyster Shells"
# Puts the data in order based on ManagedAreaName, Year, then Month
MA_YM_Stats <- as.data.table(MA_YM_Stats[order(MA_YM_Stats$ManagedAreaName,
                                               MA_YM_Stats$Year,
                                               MA_YM_Stats$Month,
                                               MA_YM_Stats$ShellType,
                                               MA_YM_Stats$SizeClass,
                                               MA_YM_Stats$HabitatType), ])
# Writes summary statistics to file
fwrite(MA YM Stats, paste0(out dir,"/Density/Oyster Dens MA MMYY Stats.txt"),
       sep="|")
# Removes variable storing data to improve computer memory
rm(MA_YM_Stats)
# Create summary statistics for each managed area based on Year intervals
MA_Y_Stats <- oysterraw[oysterraw$nyrpar=="Density_m2",] %>%
   group_by(AreaID, ManagedAreaName, Year, nyrpar, LiveDate_Qualifier,
            SizeClass, HabitatClassification) %>%
   dplyr::summarize(N_Data=length(Density_m2[!is.na(Density_m2)]),
             Min=min(Density_m2[!is.na(Density_m2)]),
             Max=max(Density_m2[!is.na(Density_m2)]),
             Median=median(Density_m2[!is.na(Density_m2)]),
             Mean=mean(Density_m2[!is.na(Density_m2)]),
             StandardDeviation=sd(Density_m2[!is.na(Density_m2)]),
             Programs=paste(sort(unique(ProgramName), decreasing=FALSE),
                              collapse=', '),
             ProgramIDs=paste(sort(unique(ProgramID), decreasing=FALSE),
                              collapse=', '))
setnames(MA_Y_Stats, c("nyrpar", "LiveDate_Qualifier",
                        "HabitatClassification"),
         c("ParameterName", "ShellType", "HabitatType"))
MA_Y_Stats$ShellType[MA_Y_Stats$ShellType=="Exact"] <- "Live Oyster Shells"
MA_Y_Stats$ShellType[MA_Y_Stats$ShellType=="Estimate"] <- "Dead Oyster Shells"
# Puts the data in order based on ManagedAreaName then Year
MA_Y_Stats <- as.data.table(MA_Y_Stats[order(MA_Y_Stats$ManagedAreaName,
                                             MA_Y_Stats$Year,
                                             MA_Y_Stats$ShellType,
                                             MA_Y_Stats$SizeClass,
```

```
MA_Y_Stats$HabitatType), ])
# Writes summary statistics to file
fwrite(MA_Y_Stats, paste0(out_dir,"/Density/Oyster_Dens_MA_Yr_Stats.txt"),
# Removes variable storing data to improve computer memory
rm(MA_Y_Stats)
# Create summary statistics for each managed area based on Month intervals.
MA M Stats <- oysterraw[oysterraw$nyrpar=="Density m2",] %>%
   group by (AreaID, ManagedAreaName, Month, nyrpar,
            LiveDate Qualifier, SizeClass,
            HabitatClassification) %>%
   dplyr::summarize(N_Data=length(Density_m2[!is.na(Density_m2)]),
             Min=min(Density_m2[!is.na(Density_m2)]),
             Max=max(Density_m2[!is.na(Density_m2)]),
             Median=median(Density_m2[!is.na(Density_m2)]),
             Mean=mean(Density_m2[!is.na(Density_m2)]),
             StandardDeviation=sd(Density_m2[!is.na(Density_m2)]),
             Programs=paste(sort(unique(ProgramName), decreasing=FALSE),
                              collapse=', '),
             ProgramIDs=paste(sort(unique(ProgramID), decreasing=FALSE),
                              collapse=', '))
setnames(MA_M_Stats, c("nyrpar", "LiveDate_Qualifier",
                        "HabitatClassification"),
         c("ParameterName", "ShellType", "HabitatType"))
MA M Stats$ShellType[MA M Stats$ShellType=="Exact"] <- "Live Oyster Shells"
MA M Stats $ShellType [MA M Stats $ShellType == "Estimate"] <- "Dead Oyster Shells"
# Puts the data in order based on ManagedAreaName then Month
MA_M_Stats <- as.data.table(MA_M_Stats[order(MA_M_Stats$ManagedAreaName,
                                             MA M Stats $ Month,
                                             MA_M_Stats$ShellType,
                                             MA_M_Stats$SizeClass,
                                             MA_M_Stats$HabitatType), ])
# Writes summary statistics to file
fwrite(MA_M_Stats, pasteO(out_dir,"/Density/Oyster_Dens_MA_Mo_Stats.txt"),
       sep="|")
# Removes variable storing data to improve computer memory
rm(MA_M_Stats)
# Create summary overall statistics for each managed area.
MA_Ov_Stats <- oysterraw[oysterraw$nyrpar=="Density_m2",] %>%
   group_by(AreaID, ManagedAreaName, nyrpar,
            LiveDate Qualifier, SizeClass,
            HabitatClassification) %>%
   dplyr::summarize(N Years=length(unique(
     LiveDate[!is.na(LiveDate) & !is.na(Density_m2)])),
             SufficientData=ifelse(N_Years>=5, TRUE, FALSE),
             EarliestLiveDate=min(LiveDate[!is.na(Density_m2)]),
             LatestLiveDate=max(LiveDate[!is.na(Density_m2)]),
             LastSampleDate=max(SampleDate),
             N_Data=length(Density_m2[!is.na(Density_m2)]),
             Min=min(Density_m2[!is.na(Density_m2)]),
             Max=max(Density_m2[!is.na(Density_m2)]),
```

```
Median=median(Density_m2[!is.na(Density_m2)]),
             Mean=mean(Density_m2[!is.na(Density_m2)]),
             StandardDeviation=sd(Density_m2[!is.na(Density_m2)]),
             Programs=paste(sort(unique(ProgramName), decreasing=FALSE),
                              collapse=', '),
             ProgramIDs=paste(sort(unique(ProgramID), decreasing=FALSE),
                              collapse=', '))
setnames(MA Ov Stats, c("nyrpar", "LiveDate Qualifier",
                        "HabitatClassification"),
         c("ParameterName", "ShellType", "HabitatType"))
MA_Ov_Stats$ShellType[MA_Ov_Stats$ShellType=="Exact"] <- "Live Oyster Shells"
MA_Ov_Stats$ShellType[MA_Ov_Stats$ShellType=="Estimate"] <- "Dead Oyster Shells"
# Puts the data in order based on ManagedAreaName
MA_Ov_Stats <- as.data.table(MA_Ov_Stats[order(MA_Ov_Stats$ManagedAreaName,
                                               MA_Ov_Stats$ShellType,
                                               MA_Ov_Stats$SizeClass,
                                               MA_Ov_Stats$HabitatType), ])
# Replaces blank ProgramIDs with NA (missing values)
MA_Ov_Stats$ProgramIDs <- replace(MA_Ov_Stats$ProgramIDs,
                                  MA_Ov_Stats$ProgramIDs=="", NA)
MA_Ov_Stats$Programs <- replace(MA_Ov_Stats$Programs,
                                  MA_Ov_Stats$Programs=="", NA)
# Write overall statistics to file
fwrite(MA Ov Stats, pasteO(out dir,"/Density/Oyster Dens MA Overall Stats.txt"),
       sep="|")
# Removes variable storing data to improve computer memory
rm(MA_Ov_Stats)
##Shell Height
# Create summary statistics for each managed area based on Year and Month
# intervals.
MA_YM_Stats <- oysterraw[oysterraw$nyrpar=="ShellHeight_mm",] %>%
   group_by(AreaID, ManagedAreaName, Year, Month, nyrpar,
            LiveDate_Qualifier, SizeClass, HabitatClassification) %>%
   dplyr::summarize(N_Data=length(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Min=min(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Max=max(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Median=median(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Mean=mean(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             StandardDeviation=sd(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Programs=paste(sort(unique(ProgramName), decreasing=FALSE),
                              collapse=', '),
             ProgramIDs=paste(sort(unique(ProgramID), decreasing=FALSE),
                              collapse=', '))
setnames(MA_YM_Stats, c("nyrpar", "LiveDate_Qualifier",
                        "HabitatClassification"),
         c("ParameterName", "ShellType", "HabitatType"))
MA_YM_Stats$ShellType[MA_YM_Stats$ShellType=="Exact"] <- "Live Oyster Shells"
MA_YM_Stats$ShellType[MA_YM_Stats$ShellType=="Estimate"] <- "Dead Oyster Shells"
# Puts the data in order based on ManagedAreaName, Year, then Month
MA_YM_Stats <- as.data.table(MA_YM_Stats[order(MA_YM_Stats$ManagedAreaName,
                                               MA_YM_Stats$Year,
```

```
MA_YM_Stats$Month,
                                               MA_YM_Stats$ShellType,
                                               MA_YM_Stats$SizeClass,
                                               MA YM Stats$HabitatType), ])
# Writes summary statistics to file
fwrite(MA_YM_Stats, paste0(out_dir,"/Shell_Height/Oyster_SH_MA_MMYY_Stats.txt"),
       sep="|")
# Removes variable storing data to improve computer memory
rm(MA YM Stats)
# Create summary statistics for each managed area based on Year intervals
MA_Y_Stats <- oysterraw[oysterraw$nyrpar=="ShellHeight_mm",] %>%
   group_by(AreaID, ManagedAreaName, Year, nyrpar, LiveDate_Qualifier,
            SizeClass, HabitatClassification) %>%
   dplyr::summarize(N_Data=length(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Min=min(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Max=max(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Median=median(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Mean=mean(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             StandardDeviation=sd(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Programs=paste(sort(unique(ProgramName), decreasing=FALSE),
                              collapse=', '),
             ProgramIDs=paste(sort(unique(ProgramID), decreasing=FALSE),
                              collapse=', '))
setnames(MA_Y_Stats, c("nyrpar", "LiveDate_Qualifier",
                        "HabitatClassification"),
         c("ParameterName", "ShellType", "HabitatType"))
MA_Y_Stats$ShellType[MA_Y_Stats$ShellType=="Exact"] <- "Live Oyster Shells"
MA_Y_Stats$ShellType[MA_Y_Stats$ShellType=="Estimate"] <- "Dead Oyster Shells"
\# Puts the data in order based on ManagedAreaName then Year
MA_Y_Stats <- as.data.table(MA_Y_Stats[order(MA_Y_Stats$ManagedAreaName,
                                             MA Y Stats $Year,
                                             MA_Y_Stats$ShellType,
                                             MA_Y_Stats$SizeClass,
                                             MA_Y_Stats$HabitatType), ])
# Writes summary statistics to file
fwrite(MA Y Stats, paste0(out dir,"/Shell Height/Oyster SH MA Yr Stats.txt"),
       sep="|")
# Removes variable storing data to improve computer memory
rm(MA_Y_Stats)
# Create summary statistics for each managed area based on Month intervals.
MA M Stats <- oysterraw[oysterraw$nyrpar=="ShellHeight mm",] %>%
   group by (AreaID, ManagedAreaName, Month, nyrpar,
            LiveDate_Qualifier, SizeClass,
            HabitatClassification) %>%
   dplyr::summarize(N_Data=length(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Min=min(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Max=max(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Median=median(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Mean=mean(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             StandardDeviation=sd(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Programs=paste(sort(unique(ProgramName), decreasing=FALSE),
```

```
collapse=', '),
             ProgramIDs=paste(sort(unique(ProgramID), decreasing=FALSE),
                              collapse=', '))
setnames(MA_M_Stats, c("nyrpar", "LiveDate_Qualifier",
                        "HabitatClassification"),
         c("ParameterName", "ShellType", "HabitatType"))
MA_M_Stats$ShellType[MA_M_Stats$ShellType=="Exact"] <- "Live Oyster Shells"
MA_M_Stats$ShellType[MA_M_Stats$ShellType=="Estimate"] <- "Dead Oyster Shells"
# Puts the data in order based on ManagedAreaName then Month
MA M Stats <- as.data.table(MA M Stats[order(MA M Stats$ManagedAreaName,
                                             MA M Stats $ Month,
                                             MA_M_Stats$ShellType,
                                             MA_M_Stats$SizeClass,
                                             MA_M_Stats$HabitatType), ])
# Writes summary statistics to file
fwrite(MA_M_Stats, paste0(out_dir,"/Shell_Height/Oyster_SH_MA_Mo_Stats.txt"),
       sep="|")
# Removes variable storing data to improve computer memory
rm(MA_M_Stats)
# Create summary overall statistics for each managed area.
MA_Ov_Stats <- oysterraw[oysterraw$nyrpar=="ShellHeight_mm",] %>%
  group_by(AreaID, ManagedAreaName, nyrpar,
            LiveDate_Qualifier, SizeClass,
            HabitatClassification) %>%
   dplyr::summarize(N Years=length(unique(
     LiveDate[!is.na(LiveDate) & !is.na(ShellHeight mm)])),
             SufficientData=ifelse(N_Years>=5, TRUE, FALSE),
             EarliestLiveDate=min(LiveDate[!is.na(ShellHeight_mm)]),
             LatestLiveDate=max(LiveDate[!is.na(ShellHeight_mm)]),
             LastSampleDate=max(SampleDate),
             N_Data=length(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Min=min(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Max=max(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Median=median(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Mean=mean(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             StandardDeviation=sd(ShellHeight_mm[!is.na(ShellHeight_mm)]),
             Programs=paste(sort(unique(ProgramName), decreasing=FALSE),
                              collapse=', '),
             ProgramIDs=paste(sort(unique(ProgramID), decreasing=FALSE),
                              collapse=', '))
setnames(MA_Ov_Stats, c("nyrpar", "LiveDate_Qualifier",
                        "HabitatClassification"),
         c("ParameterName", "ShellType", "HabitatType"))
MA_Ov_Stats$ShellType[MA_Ov_Stats$ShellType=="Exact"] <- "Live Oyster Shells"
MA_Ov_Stats$ShellType[MA_Ov_Stats$ShellType=="Estimate"] <- "Dead Oyster Shells"
# Puts the data in order based on ManagedAreaName
MA_Ov_Stats <- as.data.table(MA_Ov_Stats[order(MA_Ov_Stats$ManagedAreaName,
                                               MA_Ov_Stats$ShellType,
                                               MA_Ov_Stats$SizeClass,
                                               MA_Ov_Stats$HabitatType), ])
# Replaces blank ProgramIDs with NA (missing values)
```

```
MA_Ov_Stats$ProgramIDs <- replace(MA_Ov_Stats$ProgramIDs,
                                  MA_Ov_Stats$ProgramIDs=="", NA)
MA_Ov_Stats$Programs <- replace(MA_Ov_Stats$Programs,
                                  MA_Ov_Stats$Programs=="", NA)
# Write overall statistics to file
fwrite(MA_Ov_Stats, paste0(out_dir,"/Shell_Height/Oyster_SH_MA_Overall_Stats.txt"),
       sep="|")
# Removes variable storing data to improve computer memory
rm(MA Ov Stats)
##Percent Live
# Create summary statistics for each managed area based on Year and Month
# intervals.
MA YM Stats <- oysterraw[oysterraw$nyrpar=="PercentLive pct",] %>%
   group_by(AreaID, ManagedAreaName, Year, Month, nyrpar,
            LiveDate Qualifier, SizeClass, HabitatClassification) %>%
   dplyr::summarize(N_Data=length(PercentLive_pct[!is.na(PercentLive_pct)]),
             Min=min(PercentLive_pct[!is.na(PercentLive_pct)]),
             Max=max(PercentLive_pct[!is.na(PercentLive_pct)]),
             Median=median(PercentLive_pct[!is.na(PercentLive_pct)]),
             Mean=mean(PercentLive_pct[!is.na(PercentLive_pct)]),
             StandardDeviation=sd(PercentLive_pct[!is.na(PercentLive_pct)]),
             Programs=paste(sort(unique(ProgramName), decreasing=FALSE),
                              collapse=', '),
             ProgramIDs=paste(sort(unique(ProgramID), decreasing=FALSE),
                              collapse=', '))
setnames(MA_YM_Stats, c("nyrpar", "LiveDate_Qualifier",
                        "HabitatClassification"),
         c("ParameterName", "ShellType", "HabitatType"))
MA_YM_Stats$ShellType[MA_YM_Stats$ShellType=="Exact"] <- "Live Oyster Shells"
MA YM Stats$ShellType[MA YM Stats$ShellType=="Estimate"] <- "Dead Oyster Shells"
# Puts the data in order based on ManagedAreaName, Year, then Month
MA_YM_Stats <- as.data.table(MA_YM_Stats[order(MA_YM_Stats$ManagedAreaName,
                                               MA_YM_Stats$Year,
                                               MA_YM_Stats$Month,
                                               MA_YM_Stats$ShellType,
                                               MA_YM_Stats$SizeClass,
                                               MA_YM_Stats$HabitatType), ])
# Writes summary statistics to file
fwrite(MA_YM_Stats, paste0(out_dir,"/Percent_Live/Oyster_PrcLive_MA_MMYY_Stats.txt"),
       sep="|")
# Removes variable storing data to improve computer memory
rm(MA_YM_Stats)
# Create summary statistics for each managed area based on Year intervals
MA_Y_Stats <- oysterraw[oysterraw$nyrpar=="PercentLive_pct",] %>%
   group_by(AreaID, ManagedAreaName, Year, nyrpar, LiveDate_Qualifier,
            SizeClass, HabitatClassification) %>%
   dplyr::summarize(N_Data=length(PercentLive_pct[!is.na(PercentLive_pct)]),
             Min=min(PercentLive pct[!is.na(PercentLive pct)]),
             Max=max(PercentLive_pct[!is.na(PercentLive_pct)]),
             Median=median(PercentLive_pct[!is.na(PercentLive_pct)]),
             Mean=mean(PercentLive_pct[!is.na(PercentLive_pct)]),
```

```
StandardDeviation=sd(PercentLive_pct[!is.na(PercentLive_pct)]),
             Programs=paste(sort(unique(ProgramName), decreasing=FALSE),
                              collapse=', '),
             ProgramIDs=paste(sort(unique(ProgramID), decreasing=FALSE),
                              collapse=', '))
setnames(MA_Y_Stats, c("nyrpar", "LiveDate_Qualifier",
                        "HabitatClassification"),
         c("ParameterName", "ShellType", "HabitatType"))
MA_Y_Stats$ShellType[MA_Y_Stats$ShellType=="Exact"] <- "Live Oyster Shells"
MA_Y_Stats$ShellType[MA_Y_Stats$ShellType=="Estimate"] <- "Dead Oyster Shells"
# Puts the data in order based on ManagedAreaName then Year
MA_Y_Stats <- as.data.table(MA_Y_Stats[order(MA_Y_Stats$ManagedAreaName,
                                             MA Y Stats$Year,
                                             MA_Y_Stats$ShellType,
                                             MA_Y_Stats$SizeClass,
                                             MA_Y_Stats$HabitatType), ])
# Writes summary statistics to file
fwrite(MA_Y_Stats, paste0(out_dir,"/Percent_Live/Oyster_PrcLive_MA_Yr_Stats.txt"),
       sep="|")
# Removes variable storing data to improve computer memory
rm(MA_Y_Stats)
# Create summary statistics for each managed area based on Month intervals.
MA_M_Stats <- oysterraw[oysterraw$nyrpar=="PercentLive_pct",] %>%
   group by(AreaID, ManagedAreaName, Month, nyrpar,
            LiveDate Qualifier, SizeClass,
            HabitatClassification) %>%
   dplyr::summarize(N_Data=length(PercentLive_pct[!is.na(PercentLive_pct)]),
             Min=min(PercentLive_pct[!is.na(PercentLive_pct)]),
             Max=max(PercentLive_pct[!is.na(PercentLive_pct)]),
             Median=median(PercentLive_pct[!is.na(PercentLive_pct)]),
             Mean=mean(PercentLive_pct[!is.na(PercentLive_pct)]),
             StandardDeviation=sd(PercentLive_pct[!is.na(PercentLive_pct)]),
             Programs=paste(sort(unique(ProgramName), decreasing=FALSE),
                              collapse=', '),
             ProgramIDs=paste(sort(unique(ProgramID), decreasing=FALSE),
                              collapse=', '))
setnames(MA_M_Stats, c("nyrpar", "LiveDate_Qualifier",
                        "HabitatClassification"),
         c("ParameterName", "ShellType", "HabitatType"))
MA_M_Stats$ShellType[MA_M_Stats$ShellType=="Exact"] <- "Live Oyster Shells"
MA_M_Stats$ShellType[MA_M_Stats$ShellType=="Estimate"] <- "Dead Oyster Shells"
# Puts the data in order based on ManagedAreaName then Month
MA_M_Stats <- as.data.table(MA_M_Stats[order(MA_M_Stats$ManagedAreaName,
                                             MA_M_Stats$Month,
                                             MA_M_Stats$ShellType,
                                             MA_M_Stats$SizeClass,
                                             MA_M_Stats$HabitatType), ])
# Writes summary statistics to file
fwrite(MA_M_Stats, paste0(out_dir,"/Percent_Live/Oyster_PrcLive_MA_Mo_Stats.txt"),
       sep="|")
# Removes variable storing data to improve computer memory
rm(MA_M_Stats)
```

```
# Create summary overall statistics for each managed area.
MA_Ov_Stats <- oysterraw[oysterraw$nyrpar=="PercentLive_pct",] %>%
   group by(AreaID, ManagedAreaName, nyrpar,
           LiveDate Qualifier, SizeClass,
           HabitatClassification) %>%
   dplyr::summarize(N Years=length(unique(
     LiveDate[!is.na(LiveDate) & !is.na(PercentLive_pct)])),
            SufficientData=ifelse(N Years>=5, TRUE, FALSE),
            EarliestLiveDate=min(LiveDate[!is.na(PercentLive pct)]),
            LatestLiveDate=max(LiveDate[!is.na(PercentLive_pct)]),
            LastSampleDate=max(SampleDate),
            N_Data=length(PercentLive_pct[!is.na(PercentLive_pct)]),
            Min=min(PercentLive_pct[!is.na(PercentLive_pct)]),
            Max=max(PercentLive_pct[!is.na(PercentLive_pct)]),
            Median=median(PercentLive_pct[!is.na(PercentLive_pct)]),
            Mean=mean(PercentLive_pct[!is.na(PercentLive_pct)]),
            StandardDeviation=sd(PercentLive_pct[!is.na(PercentLive_pct)]),
            Programs=paste(sort(unique(ProgramName), decreasing=FALSE),
                              collapse=', '),
            ProgramIDs=paste(sort(unique(ProgramID), decreasing=FALSE),
                              collapse=', '))
setnames(MA_Ov_Stats, c("nyrpar", "LiveDate_Qualifier",
                        "HabitatClassification"),
         c("ParameterName", "ShellType", "HabitatType"))
MA_Ov_Stats$ShellType[MA_Ov_Stats$ShellType=="Exact"] <- "Live Oyster Shells"
MA_Ov_Stats$ShellType[MA_Ov_Stats$ShellType=="Estimate"] <- "Dead Oyster Shells"
# Puts the data in order based on ManagedAreaName
MA_Ov_Stats <- as.data.table(MA_Ov_Stats[order(MA_Ov_Stats$ManagedAreaName,
                                               MA_Ov_Stats$ShellType,
                                               MA_Ov_Stats$SizeClass,
                                               MA_Ov_Stats$HabitatType), ])
# Replaces blank ProgramIDs with NA (missing values)
MA_Ov_Stats$ProgramIDs <- replace(MA_Ov_Stats$ProgramIDs,
                                 MA_Ov_Stats$ProgramIDs=="", NA)
MA_Ov_Stats$Programs <- replace(MA_Ov_Stats$Programs,</pre>
                                  MA Ov Stats$Programs=="", NA)
# Write overall statistics to file
fwrite(MA_Ov_Stats, paste0(out_dir,"/Percent_Live/Oyster_PrcLive_MA_Overall_Stats.txt"),
       sep="|")
# Removes variable storing data to improve computer memory
rm(MA_Ov_Stats)
# LiveDate Threshold -----
oysterraw <- oysterraw[oysterraw$LiveDate>=1960,]
for(i in unique(oysterraw$ManagedAreaName)){
      oysterraw[ManagedAreaName==i & !is.na(LiveDate), `:=`
                (RelYear=(LiveDate-min(LiveDate))+1)]
```

Functions

Sets universal them for plots and creates functions that are used to create figures and data files from data subsets and GLMM models. The functions are:

- 1. diagnosticplots
- Creates panel of summary plots for each analysis
- 2. meplots
- Creates and saves plots for density and live percent
- 3. modresults
- Takes input data and model results for density and live percent.
- Adds model results to compilation variable.
- Send data and model results to diagnostic plots and meplots
- 4. meplotssh
- Creates and saves plots for shell height
- 5. modresultssh
- Takes input data and model results for shell height.
- Adds model results to compilation variable.
- Send data and model results to diagnostic plots and meplotssh

```
plot_theme <- theme_bw() +</pre>
  theme(panel.grid.major = element_blank(),
        panel.grid.minor = element_blank(),
        text=element_text(family="Arial"),
        plot.title=element text(hjust=0.5, size=12, color="#314963"),
        plot.subtitle=element_text(hjust=0.5, size=10, color="#314963"),
        legend.title=element text(size=10),
        legend.text = element_text(hjust = 0),
        axis.title.x = element_text(size=10, margin = margin(t = 5, r = 0,
                                                               b = 10, 1 = 0)),
        axis.title.y = element_text(size=10, margin = margin(t = 0, r = 10,
                                                               b = 0, 1 = 0)),
        axis.text=element_text(size=10),
        axis.text.x=element_text(angle = -45, hjust = 0))
plot_jitter <- position_jitter(width = 0.1, height = 0.1, seed=42)</pre>
#Function to save diagnostic plots
diagnosticplots <- function(model, indicator, managedarea, sizeclass="",
                            historical=FALSE){
  ind <- case_when(str_detect(indicator, "ercent") ~ "Pct",</pre>
                   str_detect(indicator, "ensity") ~ "Den",
                   str detect(indicator, "^S|^s") ~ "SH")
  ma <- managedarea
  if(sizeclass != ""){
    size <- case when(str detect(sizeclass, "25") &</pre>
                        str_detect(sizeclass, "75") ~ "25to75",
                      str_detect(sizeclass, "35") &
                         str_detect(sizeclass, "75") ~ "35to75",
```

```
str_detect(sizeclass, "25")==FALSE &
                       str_detect(sizeclass, "75") ~ "o75", TRUE ~ "raw")
  sizelab <- case_when(str_detect(sizeclass, "25") &</pre>
                          str detect(sizeclass, "75") ~ "25-75mm",
                        str_detect(sizeclass, "35") &
                          str_detect(sizeclass, "75") ~ "35-75mm",
                        str_detect(sizeclass, "25")==FALSE &
                          str detect(sizeclass, "75") ~ "\u2265 75mm",
                        TRUE ~ "raw")
}
#Save diagnostic plot(s) of chains
diag <- plot(model, plot=FALSE)</pre>
title <- textGrob(pasteO(ma, " (", ind, " ", sizelab, ")"),
                  just="left",
                  gp=gpar(fontsize=10))
diag[[1]] <- gtable_add_rows(</pre>
 diag[[1]],
 heights=grobHeight(title)+unit(5, "mm"),
 pos=0
)
diag[[1]] <- gtable_add_grob(</pre>
 diag[[1]],
 title,
 clip="off",
 1, 1, 1, 1)
if(class(try(diag[[2]], silent=TRUE)) != "try-error"){
 diag[[2]] <- gtable_add_rows(</pre>
    diag[[2]],
    heights=grobHeight(title)+unit(5, "mm"),
   pos=0
 )
}
if(class(try(diag[[3]], silent=TRUE)) != "try-error"){
 diag[[3]] <- gtable_add_rows(</pre>
    diag[[3]],
    heights=grobHeight(title)+unit(5, "mm"),
    pos=0
 )
}
#save chains plots
jpeg(filename=paste0("output/Figures/", ind, "_AllDates_GLMM_", ma,
                      "_PDistandMChains_", ifelse(sizeclass != "",
                                                   paste0(size, "_"), ""),
                      ifelse(historical==TRUE, "hist_", "_"),
                      Sys.Date(), ".png"),
     width=6,
```

```
height=ifelse(length(diag)==1, 6, ifelse(length(diag)==2, 12, 18)),
       units="in", quality=100, res=300)
  print(grid.arrange(grobs=diag, ncol=1))
  dev.off()
  #Save posterior predictive check plot
  postpc <- tryCatch(pp_check(model),</pre>
                      error=function(e) NA)
 k <- 1001
  while(is.na(postpc)==TRUE & k <= 1000){</pre>
    postpc <- tryCatch(pp_check(model),</pre>
                        error=function(e) NA)
   k <- k+1
  if(!is.na(postpc)){
    postpc <- postpc +</pre>
      labs(title=paste0(ind, "_AllDates_GLMM_", ma, "_PPcheck_",
                         ifelse(sizeclass != "", paste0(size, "_"), ""),
                         ifelse(historical==TRUE, "_hist_", "_"),
                         Sys.Date(), ".png"))
    ma_short <- MA_All[ManagedAreaName==ma, Abbreviation]</pre>
    ggsave(paste0("output/Figures/", ind, "_AllDates_GLMM_", ma_short, "_PPcheck_",
                  ifelse(sizeclass != "", paste0(size, "_"), ""),
                   ifelse(historical==TRUE, "_hist_", "_"), Sys.Date(),
                   ".png"),
           postpc,
           width=6.
           height=6,
           units="in",
           dpi=300)
 }
 print(paste0("Plots saved."))
}
# Save marginal effects plots
meplots <- function(models, data, indicator, managedarea, sizeclass="",
                    zoom=FALSE) {
  ind <- case_when(str_detect(indicator, "ercent") ~ "Pct",</pre>
                   str_detect(indicator, "ensity") ~ "Den",
                    str_detect(indicator, "^S|^s") ~ "SH")
 ma <- managedarea
  if(sizeclass != ""){
    size <- case_when(str_detect(sizeclass, "25") &</pre>
                         str_detect(sizeclass, "75") ~ "25to75",
                       str_detect(sizeclass, "35") &
```

```
str_detect(sizeclass, "75") ~ "35to75",
                     str_detect(sizeclass, "25")==FALSE &
                       str_detect(sizeclass, "75") ~ "o75", TRUE ~ "raw")
  sizelab <- case_when(str_detect(sizeclass, "25") &</pre>
                          str_detect(sizeclass, "75") ~ "25-75mm",
                        str_detect(sizeclass, "35") &
                          str_detect(sizeclass, "75") ~ "35-75mm",
                        str detect(sizeclass, "25")==FALSE &
                          str_detect(sizeclass, "75") ~ "\u2265 75mm",
                        TRUE ~ "raw")
}
if(ind=="Den"){
  nyrs <- max(data$LiveDate)-min(data$LiveDate)+1</pre>
  maxyr <- max(data$LiveDate)</pre>
  minyr <- min(data$LiveDate)</pre>
  if(grepl("Natural", unique(data$MA_plotlab))==TRUE){
    type <- "Natural"</pre>
  } else{
    type <- "Restored"</pre>
  yrdiff <- unique(data$YearDiff)</pre>
  # Creates break intervals for plots based on number of years of data
  if(nyrs > = 40){
    # Set breaks to every 10 years if more than 40 years of data
    brk <- 10
  }else if(nyrs<40 & nyrs>=20){
    # Set breaks to every 5 years if between 40 and 20 years of data
    brk <- 5
  }else if(nyrs<20 & nyrs>=12){
    # Set breaks to every 3 years if between 20 and 12 years of data
    brk <- 3
  }else if(nyrs<12 & nyrs>=8){
    # Set breaks to every 2 years if between 12 and 8 years of data
    brk <- 2
  }else if(nyrs<8){</pre>
    # Set breaks to every year if less than 8 years of data
    brk <- 1
  }else if(nyrs<3){</pre>
    brk <- 1
    maxyr <- maxyr + 1</pre>
    minyr <- minyr - 1
  yrlist <- seq(minyr,maxyr,brk)</pre>
  # nbreaks <- ifelse(nyrs < 11, nyrs+1, 12)</pre>
  # breaks <- if(minyr==0){
      c(minyr, round(minyr+c(1:(nbreaks-2))*((nyrs/nbreaks) +
  #
                                                  (nyrs/nbreaks)/nbreaks)),
        maxyr)+1
  # } else{
```

```
c(minyr, round(minyr+c(1:(nbreaks-2))*((nyrs/nbreaks) +
#
                                               (nyrs/nbreaks)/nbreaks)),
#
      maxyr)
# }
denplots <- plot(conditional_effects(models[[1]], re_formula=NULL),</pre>
                 plot=FALSE)
plot1 <- ggplot() +</pre>
  {if("meanDen_int" %in% colnames(data)){
    geom_point(data=data, aes(x=LiveDate,
                               y=meanDen_int), position=plot_jitter,
               shape=21, size=2, color="#333333", fill="#cccccc",
               alpha=1, inherit.aes=FALSE)
  } else{
    geom_point(data=data, aes(x=LiveDate,
                               y=Density_m2), position=plot_jitter,
               shape=21, size=2, color="#333333", fill="#cccccc",
               alpha=1, inherit.aes=FALSE)
  }
  } +
  list(geom_ribbon(data=denplots$RelYear$data,
                   aes(x=RelYear+yrdiff, y=Density_m2,
                       ymin=lower__, ymax=upper__),
                   fill="#000099", alpha=0.1, inherit.aes=FALSE),
       geom_line(data=denplots$RelYear$data,
                 aes(x=RelYear+yrdiff,
                      y=estimate__),
                 color="#000099", lwd=0.75, inherit.aes=FALSE)) +
  scale_x_continuous(limits=c(minyr-0.25, maxyr+0.25),
                     breaks=yrlist) +
  plot_theme +
  {if("meanDen_int" %in% colnames(data)){
    labs(title="Oyster Density",
         subtitle=managedarea,
         x="Year",
         y=bquote('Estimated density ('*~m^{-2}*')'))
    labs(title="Oyster Density",
         subtitle=managedarea,
         x="Year",
         y=bquote('Density ('*~m^{-2}*')'))
  }}
# labs(title="Oyster Density",
#
       subtitle=managedarea,
#
       x="Year",
       y=ifelse("meanDen_int" %in% colnames(data),
#
#
                             "Estimated density (square meters)",
#
                             bquote('Richness (species/100'*~m^{2}*')')))
ma_short <- MA_All[ManagedAreaName==ma, Abbreviation]</pre>
```

```
ggsave(paste0("output/Density/Figures/Oyster_Dens_GLMM_", ma_short, "_", type,
                ifelse(sizeclass != "", pasteO(size), "_raw"), ".png"),
         plot1,
         width=8.
         height=4,
         units="in",
         dpi=200)
}
#Marginal effects plot including random effects for percent live
if(ind=="Pct"){
 nyrs <- max(data$LiveDate)-min(data$LiveDate)+1</pre>
 maxyr <- max(data$LiveDate)</pre>
 minyr <- min(data$LiveDate)</pre>
 yrdiff <- unique(data$YearDiff)</pre>
  # Creates break intervals for plots based on number of years of data
 if(nyrs>=40){
    # Set breaks to every 10 years if more than 40 years of data
    brk <- 10
 }else if(nyrs<40 & nyrs>=20){
    # Set breaks to every 5 years if between 40 and 20 years of data
    brk <- 5
 }else if(nyrs<20 & nyrs>=12){
    # Set breaks to every 3 years if between 20 and 12 years of data
    brk <- 3
 }else if(nyrs<12 & nyrs>=8){
    # Set breaks to every 2 years if between 12 and 8 years of data
    brk <- 2
 }else if(nyrs<8){</pre>
    # Set breaks to every year if less than 8 years of data
    brk <- 1
 }else if(nyrs<3){</pre>
    brk <- 1
    maxyr <- maxyr + 1</pre>
    minyr <- minyr - 1
 }
 yrlist <- seq(minyr,maxyr,brk)</pre>
  # nbreaks <- ifelse(nyrs < 11, nyrs+1, 12)</pre>
  # breaks <- if(minyr==0){
     c(minyr, round(minyr+c(1:(nbreaks-2))*((nyrs/nbreaks) +
                                                  (nyrs/nbreaks)/nbreaks)),
  #
        maxyr)+1
  # } else{
  #
     c(minyr, round(minyr+c(1:(nbreaks-2))*((nyrs/nbreaks) +
  #
                                                  (nyrs/nbreaks)/nbreaks)),
  #
        maxyr)
  # }
 set.seed(987)
 pctplots <- plot(conditional_effects(models[[1]], re_formula=NULL),</pre>
                    plot=FALSE)
```

```
plot1 <- ggplot() +</pre>
  geom_point(data=data, aes(x=LiveDate,
                            y=100*PercentLive_dec), position=plot_jitter,
             shape=21, size=2, color="#333333", fill="#cccccc",
             alpha=1, inherit.aes=FALSE) +
  {if(names(pctplots$RelYear$data[2])=="PercentLive_dec"){
    list(geom_ribbon(data=pctplots$RelYear$data,
                     aes(x=RelYear+yrdiff,
                          y=100*PercentLive_dec, ymin=100*lower__,
                         ymax=100*upper__), fill="#000099",
                     alpha=0.1, inherit.aes=FALSE),
         geom_line(data=pctplots$RelYear$data,
                   aes(x=RelYear+yrdiff,
                       y=100*estimate__), color="#000099",
                   lwd=0.75, inherit.aes=FALSE))
  } else{
    list(geom_ribbon(data=pctplots$RelYear$data,
                     aes(x=RelYear+yrdiff,
                         y=100*LiveObs, ymin=100*lower__,
                         ymax=100*upper__), fill="#000099",
                     alpha=0.1, inherit.aes=FALSE),
         geom_line(data=pctplots$RelYear$data,
                   aes(x=RelYear+yrdiff,
                       y=100*estimate__), color="#000099",
                   lwd=0.75, inherit.aes=FALSE))
  }} +
  scale_x_continuous(limits=c(minyr-0.25, maxyr+0.25),
                     breaks=yrlist) +
  plot theme +
  theme(legend.text=element_text(size=10),
        legend.title=element_text(size=10)) +
    if(managedarea=="Lemon Bay Aquatic Preserve"){
      labs(title="Percent of Live vs. Dead Oysters",
           subtitle=managedarea,
           x="Year",
           y="Live vs. dead (%)")
      labs(title="Oyster Percent Live Cover",
           subtitle=managedarea,
           x="Year",
           y="Live cover (%)")
  }
ma_short <- MA_All[ManagedAreaName==ma, Abbreviation]</pre>
ggsave(paste0("output/Percent_Live/Figures/Oyster_PrcLive_GLMM_", ma_short,
              "_raw.png"),
       plot1,
       width=8,
       height=4,
       units="in",
```

```
dpi=200)
#Plot of modeled mean percent live
if("Region" %in% names(pctplots)){
  meanPct <- pctplots$Region$data</pre>
  setnames(meanPct, "effect1__", "Region")
  meanpctplot <- ggplot(meanPct, aes(x=Region, y=estimate__,</pre>
                                       ymin=lower__, ymax=upper__)) +
    geom_pointinterval(fill="black", size=3,
                        fatten_point=4, shape=21,
                        color="black") +
    labs(title="Oyster Percent Live Cover",
         subtitle=managedarea,
         y="Live cover (%)",
         fill=NULL) +
    plot_theme +
    theme(legend.text=element_text(size=10),
          legend.title=element_text(size=10))
  ma_short <- MA_All[ManagedAreaName==ma, Abbreviation]</pre>
  ggsave(paste0("output/Percent_Live/Figures/Oyster_PrcLive_GLMM_", ma_short,
                 "_raw_MeanRes.png"),
         meanpctplot,
         width=8,
         height=4,
         units="in",
         dpi=200)
}
#Plot of RelYear * Region interaction
if("RelYear:Region" %in% names(pctplots)){
  pctplots$RelYear$data$RelYear <-</pre>
    pctplots$RelYear$data$RelYear-
    (min(pctplots$RelYear$data$RelYear)-1)
  RelYrbyRegion <- pctplots$`RelYear:Region`</pre>
  intplot <- RelYrbyRegion +</pre>
    geom_point(data=data, aes(x=RelYear-(min(RelYear)-1),
                               y=PercentLive_dec,
                               fill=Region),
               alpha=0.5, shape=21, size=3, color="black",
               inherit.aes=FALSE) +
    scale_x_continuous(breaks=breaks,
                        labels=c(yrlist[breaks])) +
    labs(title=ma,
         x="Year",
         y="Proportion live",
         fill="Region") +
    plot_theme +
    theme(legend.text=element_text(size=12),
```

```
legend.title=element_text(size=13),
              legend.position="none") +
        facet_wrap(~ Region, ncol=3, scales="free")
      ma_short <- MA_All[ManagedAreaName==ma, Abbreviation]</pre>
      ggsave(paste0("output/Percent_Live/Figures/Oyster_PrcLive_GLMM_", ma_short,
                    " raw.png"),
             intplot,
             width=10,
             height=10,
             units="in",
             dpi=300)
    }
 }
}
# Create model results tables and save diagnostic plots
modresults <- function(datafile, models, indicator, meplotzoom=FALSE){</pre>
  for(m in seq_along(models)){
    modelobj <- models[[m]]</pre>
    sizeclass <- ifelse(str_detect(modelobj$file, "25to75|seed"),</pre>
                         "25-75mm",
                        ifelse(str_detect(modelobj$file, "35to75|seed"),
                                "35-75mm",
                                ifelse(str_detect(modelobj$file,
                                                  "o75|market"),
                                       ">75mm", "NA")))
    oyres_i <- setDT(broom.mixed::tidy(modelobj))</pre>
    #tidy() does not like that parameter values have underscores for
    #some reason, so the resulting table is incomplete
    if(nrow(oyres_i[effect=="fixed", ])-nrow(summary(modelobj)$fixed)==-1){
      missingrow <- data.table(effect="fixed",
                                component="cond",
                                #not sure what "cond" means in the tidy summary.
                                group=NA,
                                term=rownames(summary(modelobj)$fixed)[2],
                                estimate=summary(modelobj)$fixed$Estimate[2],
                                std.error=summary(modelobj)$fixed$Est.Error[2],
                                conf.low=summary(modelobj)$fixed$`1-95% CI`[2],
                                conf.high=summary(modelobj)$fixed$`u-95% CI`[2])
      oyres_i <- rbind(oyres_i, missingrow) %>% arrange(effect, group)
    oyres_i[, `:=` (indicator=indicator,
                    managed_area=unique(datafile$ManagedAreaName),
                    habitat_class=unique(datafile$HabitatClassification),
                    size_class=sizeclass,
                    live_date_qual=ifelse(
                      str_detect(modelobj$file, "_hist"),
                      "Estimate", "Exact"),
                    n_programs=if(
```

```
class(try(datafile$LiveDate_Qualifier)) !=
                       "try-error"){
                      length(
                         unique(
                           datafile[LiveDate_Qualifier==
                                      ifelse(
                                        str_detect(
                                          modelobj$file,
                                          " hist"),
                                        "Estimate".
                                        "Exact"),
                                    ProgramID]))
                    } else{length(unique(datafile[, ProgramID]))},
                    programs=if(class(try(
                       datafile$LiveDate_Qualifier)) != "try-error"){
                      list(unique(datafile[LiveDate_Qualifier==
                                               ifelse(
                                                 str_detect(
                                                   modelobj$file,
                                                   " hist"),
                                                 "Estimate",
                                                 "Exact"),
                                            ProgramID]))
                    } else{list(unique(datafile[, ProgramID]))},
                    filename=modelob;$file)]
    oysterresults <<- rbind(oysterresults, oyres_i)</pre>
    # Save diagnostic plots
    #diagnosticplots(modelobj, indicator,
    #unique(datafile$ManagedAreaName), sizeclass,
    #ifelse(str_detect(modelobj$file, "_hist"), TRUE, FALSE))
  }
  # Save marginal effects plots
  meplots(models, datafile, indicator, unique(datafile$ManagedAreaName),
          sizeclass, meplotzoom)
}
# Marginal effects plots for shell height (attempt to combine models into one plot)
meplotssh <- function(models1, data1, sizeclass1="", models2, data2,</pre>
                       sizeclass2="", managedarea, indicator, zoom=FALSE){
  ind <- case_when(str_detect(indicator, "ercent") ~ "Pct",</pre>
                   str_detect(indicator, "ensity") ~ "Den",
                   str detect(indicator, "^S|^s") ~ "SH")
  ma <- managedarea
  sizeclass1 <- unique(data1$SizeClass)</pre>
  sizeclass2 <- unique(data2$SizeClass)</pre>
  # Set size labels
  if(sizeclass1 != ""){
    size1 <- case_when(</pre>
      str_detect(sizeclass1, "25") & str_detect(sizeclass1, "75") ~ "25to75",
      str_detect(sizeclass1, "35") & str_detect(sizeclass1, "75") ~ "35to75",
      str_detect(sizeclass1, "25") == FALSE & str_detect(sizeclass1, "75") ~ "075",
```

```
TRUE ~ "raw")
  sizelab1 <- case_when(</pre>
    str_detect(sizeclass1, "25") & str_detect(sizeclass1, "75") ~ "25-75mm",
    str_detect(sizeclass1, "35") & str_detect(sizeclass1, "75") ~ "35-75mm",
    str_detect(sizeclass1, "25") == FALSE & str_detect(sizeclass1, "75") ~ "\u2265 75mm",
    TRUE ~ "raw")
}
if(sizeclass2 != ""){
 size2 <- case when(</pre>
    str_detect(sizeclass2, "25") & str_detect(sizeclass2, "75") ~ "25to75",
    str_detect(sizeclass2, "35") & str_detect(sizeclass2, "75") ~ "35to75",
    str_detect(sizeclass2, "25") == FALSE & str_detect(sizeclass2, "75") ~ "075",
    TRUE ~ "raw")
 sizelab2 <- case_when(</pre>
    str_detect(sizeclass2, "25") & str_detect(sizeclass2, "75") ~ "25-75mm",
    str_detect(sizeclass2, "35") & str_detect(sizeclass2, "75") ~ "35-75mm",
    str_detect(sizeclass2, "25") == FALSE & str_detect(sizeclass2, "75") ~ "\u2265 75mm",
    TRUE ~ "raw")
}
#Marginal effects plot including random effects
## Hist plot settings
y_max <- round(max(data2[!is.na(ShellHeight_mm), ShellHeight_mm]), -0)+1
y_breaks <- seq(0, 300, 50)</pre>
y_{labs} \leftarrow seq(0, 300, 50)
y_{minor} \leftarrow seq(0, 300, 25)
ylim_upper <- ceiling(y_max/25)*25</pre>
yrdiff1 <- unique(data1$YearDiff)</pre>
yrdiff2 <- unique(data2$YearDiff)</pre>
# function to set year breaks, type == "hist" or "live"
set_breaks <- function(type, data1, data2){</pre>
 ldq <- ifelse(type=="hist", "Estimate", "Exact")</pre>
 maxyr <- max(data1[!is.na(LiveDate) & LiveDate_Qualifier==ldq, LiveDate],</pre>
                data2[!is.na(LiveDate) & LiveDate_Qualifier==ldq, LiveDate])
 minyr <- min(data1[!is.na(LiveDate) & LiveDate_Qualifier==ldq, LiveDate],</pre>
                data2[!is.na(LiveDate) & LiveDate_Qualifier==ldq, LiveDate])
 nyrs <- (maxyr)-(minyr)+1</pre>
  # Creates break intervals for plots based on number of years of data
 if(nyrs > = 40){
    # Set breaks to every 10 years if more than 30 years of data
    brk <- 10
 }else if(nyrs<40 & nyrs>=20){
    # Set breaks to every 5 years if between 30 and 15 years of data
 }else if(nyrs<20 & nyrs>=12){
    # Set breaks to every 3 years if between 15 and 9 years of data
    brk <- 3
 }else if(nyrs<12 & nyrs>=8){
    # Set breaks to every 2 years if between 9 and 6 years of data
```

```
brk <- 2
  }else if(nyrs<8 & nyrs>=3){
    # Set breaks to every year if less than 6 years of data
    brk <- 1
  }else if(nyrs<3){</pre>
    brk <- 1
    maxyr <- maxyr + 1</pre>
    minyr <- minyr - 1
  }
  return(list("seq" = seq(minyr,maxyr,brk),"maxyr" = maxyr, "minyr" = minyr))
yrlist_hist <- set_breaks(type = "hist", data1 = data1, data2 = data2)[["seq"]]</pre>
maxyr_hist <- set_breaks(type = "hist", data1 = data1, data2 = data2)[["maxyr"]]</pre>
minyr_hist <- set_breaks(type = "hist", data1 = data1, data2 = data2)[["minyr"]]</pre>
yrlist_live <- set_breaks(type = "live", data1 = data1, data2 = data2)[["seq"]]</pre>
maxyr_live <- set_breaks(type = "live", data1 = data1, data2 = data2)[["maxyr"]]</pre>
minyr_live <- set_breaks(type = "live", data1 = data1, data2 = data2)[["minyr"]]
## Check data for Exact and Estimate
n hist1 <- nrow(data1[data1$LiveDate Qualifier=="Estimate" &
                          !is.na(data1$ShellHeight mm),])
n_live1 <- nrow(data1[data1$LiveDate_Qualifier=="Exact" &</pre>
                         !is.na(data1$ShellHeight_mm),])
n_hist2 <- nrow(data2[data2$LiveDate_Qualifier=="Estimate" &</pre>
                         !is.na(data2$ShellHeight mm),])
n_live2 <- nrow(data2[data2$LiveDate_Qualifier=="Exact" &
                         !is.na(data2$ShellHeight_mm),])
set.seed(987)
if(!is.null(models1)){
  liveplot_1 <- plot(conditional_effects(models1[[1]], re_formula=NULL), plot=FALSE)</pre>
if(!is.null(models2)){
  liveplot_2 <- plot(conditional_effects(models2[[1]], re_formula=NULL), plot=FALSE)</pre>
# Set boolean values for whether liveplot162 are available
liveplot1_avail <- class(try(liveplot_1, silent=TRUE)) != "try-error"</pre>
liveplot2_avail <- class(try(liveplot_2, silent=TRUE)) != "try-error"</pre>
# Set ribbon transparency value
a ribb \leftarrow 0.2
# Set size and shapes for plots
p_shape <- c("size2"=24, "size1"=21)</pre>
sizelab <- c("size2"=sizelab2, "size1"=sizelab1)</pre>
check <- NA
check1 <- NA
check2 <- NA
if(exists("present1")){
```

```
check1 <- c("size1"="#00374f")
} else{
  check1 <- c("size1"="#FFFFFF")</pre>
}
if(exists("present2")){
  check2 <- c("size2"="#0094b0")</pre>
} else{
  check2 <- c("size2"="#FFFFFF")</pre>
p_color <- c(check2, check1)</pre>
# Initial plots to set legends
plot_leg <- ggplot() +</pre>
 {if(liveplot1_avail){
    list(geom_ribbon(data=liveplot_1$RelYear$data,
                     aes(x=RelYear+yrdiff1, y=ShellHeight_mm,
                          ymin=lower__, ymax=upper__,
                          fill="size1"), alpha=a_ribb),
         geom_line(data=liveplot_1$RelYear$data,
                   aes(x=RelYear+yrdiff1, y=estimate__, color="size1"),
                   1wd=0.75)
 }} +
  {if(liveplot2_avail){
    list(geom ribbon(data=liveplot 2$RelYear$data,
                      aes(x=RelYear+yrdiff2, y=ShellHeight_mm,
                          ymin=lower__, ymax=upper__,
                          fill="size2"), alpha=a_ribb),
         geom_line(data=liveplot_2$RelYear$data,
                   aes(x=RelYear+yrdiff2, y=estimate__, color="size2"),
                   1wd=0.75)
 }} +
  geom_point(data=data1[!is.na(RelYear) & !is.na(LiveDate), ],
             aes(x=LiveDate, y=ShellHeight_mm, shape="size1"),
             position=plot_jitter, size=2, color="#333333", fill="#ccccc",
             alpha=0.2, inherit.aes=FALSE) +
  geom_point(data=data2[!is.na(RelYear) & !is.na(LiveDate), ],
             aes(x=LiveDate, y=ShellHeight_mm, shape="size2"),
             position=plot_jitter, size=2, color="#333333", fill="#ccccc",
             alpha=0.2, inherit.aes=FALSE) +
 plot_theme +
  theme(legend.position="right") +
  scale shape manual(name="Shell heights",
                      values=p shape,
                     labels=sizelab) +
  scale_color_manual(name="Shell heights",
                      values=p_color,
                     labels=sizelab) +
  scale_fill_manual(name="Shell heights",
                    values=p_color,
                    labels=sizelab)
leg <-get_legend(plot_leg)</pre>
```

```
rm(plot_leg)
# Dead oyster shell plot
plot1 <- ggplot() +
  geom_hline(yintercept=75, size=1, color="grey") +
  {if(n hist1>0){
    geom_point(data=data1[!is.na(RelYear) &
                            !is.na(LiveDate) &
                            LiveDate Qualifier=="Estimate", ],
               aes(x=LiveDate, y=ShellHeight_mm, shape="size1"),
               position=plot_jitter, size=2, color="#333333", fill="#ccccc",
               alpha=0.2, inherit.aes=FALSE)
 }} +
  {if(n_hist2>0){
    geom_point(data=data2[!is.na(RelYear) & !is.na(LiveDate) &
                            LiveDate_Qualifier=="Estimate", ],
               aes(x=LiveDate, y=ShellHeight_mm, shape="size2"),
               position=plot_jitter, size=2, color="#333333", fill="#ccccc",
               alpha=0.2, inherit.aes=FALSE)
 }} +
  scale_x_continuous(limits=c(minyr_hist-0.25, maxyr_hist+0.25),
                     breaks=yrlist_hist) +
  scale_y_continuous(breaks=y_breaks,
                     labels=y_labs, minor_breaks=y_minor) +
 plot theme +
  theme(plot.subtitle=element text(hjust=0, size=10, color="#314963"),
        legend.position="none",
 ) +
 labs(subtitle="Dead Oyster Shells",
       x="Estimated year",
       y="Shell height (mm)") +
  scale_shape_manual(name="Shell heights",
                     values=c("size1"=21, "size2"=24),
                     labels=c(sizelab1, sizelab2)) +
  scale_color_manual(name="Shell heights",
                     values=c("size1"="#00374f", "size2"="#0094b0"),
                     labels=c(sizelab1, sizelab2)) +
  scale_fill_manual(name="Shell heights",
                    values=c("size1"="#00374f", "size2"="#0094b0"),
                    labels=c(sizelab1, sizelab2)) +
  coord_cartesian(ylim=c(0, ylim_upper))
# Live oyster shell plot
plot2 <- ggplot() +</pre>
 geom_hline(yintercept=75, size=1, color="grey") +
  {if(n_live1>0){
    geom_point(data=data1[!is.na(RelYear) & !is.na(LiveDate) &
                            LiveDate_Qualifier=="Exact", ],
               aes(x=LiveDate, y=ShellHeight_mm, shape="size1"),
               position=plot_jitter, size=2, color="#333333", fill="#ccccc",
               alpha=0.2, inherit.aes=FALSE)
 }} +
  \{if(n_1ive2>0)\}
```

```
geom_point(data=data2[!is.na(RelYear) & !is.na(LiveDate) &
                            LiveDate_Qualifier=="Exact", ],
               aes(x=LiveDate, y=ShellHeight_mm, shape="size2"),
               position=plot_jitter, size=2, color="#333333", fill="#cccccc",
               alpha=0.2, inherit.aes=FALSE)
 }} +
  {if(liveplot1_avail){
    list(geom ribbon(data=liveplot 1$RelYear$data,
                     aes(x=RelYear+yrdiff1, y=ShellHeight mm,
                         ymin=lower__, ymax=upper__, fill="size1"),
                     alpha=a_ribb),
         geom_line(data=liveplot_1$RelYear$data,
                   aes(x=RelYear+yrdiff1, y=estimate__, color="size1"),
                   1wd=0.75)
 }} +
  {if(liveplot2_avail){
    list(geom_ribbon(data=liveplot_2$RelYear$data,
                     aes(x=RelYear+yrdiff2, y=ShellHeight_mm,
                         ymin=lower__, ymax=upper__, fill="size2"),
                     alpha=a_ribb),
         geom_line(data=liveplot_2$RelYear$data,
                   aes(x=RelYear+yrdiff2, y=estimate__, color="size2"),
 }} +
  scale x continuous(limits=c(minyr live-0.25, maxyr live+0.25),
                     breaks=yrlist live) +
  scale_y_continuous(breaks=y_breaks,
                     labels=y_labs, minor_breaks=y_minor) +
 plot theme +
  theme(plot.subtitle=element_text(hjust=0, size=10, color="#314963"),
        legend.position="none",
        axis.text.y=element_blank(), #remove y-axis labels
        axis.ticks.y=element_blank(), #remove y-axis ticks
        axis.title.y=element_blank() #removes y-axis title
  ) +
  labs(subtitle="Live Oyster Shells",
       x="Year",
      v="Shell height (mm)") +
  scale_shape_manual(name="Shell heights",
                     values=c("size1"=21, "size2"=24),
                     labels=c(sizelab1, sizelab2)) +
  scale_color_manual(name="Shell heights",
                     values=c("size1"="#00374f", "size2"="#0094b0"),
                     labels=c(sizelab1, sizelab2)) +
  scale_fill_manual(name="Shell heights",
                    values=c("size1"="#00374f", "size2"="#0094b0"),
                    labels=c(sizelab1, sizelab2)) +
  coord_cartesian(ylim=c(0, ylim_upper))
# Set plot title
plot_title <- ggplot() +</pre>
 labs(title="Oyster Size Class", subtitle=ma) +
 plot_theme +
```

```
theme(plot.subtitle=element_text(hjust=0.5, size=10, color="#314963"),
          panel.border=element blank(),
          panel.grid.major=element_blank(),
          panel.grid.minor=element_blank(), axis.line=element_blank())
  plot_comb <- ggarrange(plot1, plot2, leg, nrow=1,</pre>
                          widths=c(0.46, 0.39, 0.15))
  plot_comb <- ggarrange(plot_title, plot_comb, ncol=1,</pre>
                          heights=c(0.125, 0.875))
  ma_abrev <- MA_All[ManagedAreaName==ma, Abbreviation]</pre>
  ggsave(paste0("output/Shell_Height/Figures/Oyster_SH_GLMM_", ma_abrev, ".png"),
         plot_comb,
         width=8,
         height=4,
         units="in",
         dpi=200,
         bg="white")
 rm(liveplot_1, liveplot_2)
# Marginal effects plots for shell height (attempt to combine models into one plot)
meplotssh <- function(models1, data1, sizeclass1="", models2, data2,</pre>
                       sizeclass2="", managedarea, indicator, zoom=FALSE){
  ind <- case when(str detect(indicator, "ercent") ~ "Pct",</pre>
                    str_detect(indicator, "ensity") ~ "Den",
                    str_detect(indicator, "^S|^s") ~ "SH")
 ma <- managedarea
  sizeclass1 <- unique(data1$SizeClass)</pre>
  sizeclass2 <- unique(data2$SizeClass)</pre>
  # Set size labels
  if(sizeclass1 != ""){
    size1 <- case_when(</pre>
      str_detect(sizeclass1, "25") & str_detect(sizeclass1, "75") ~ "25to75",
      str_detect(sizeclass1, "35") & str_detect(sizeclass1, "75") ~ "35to75",
      str_detect(sizeclass1, "25") == FALSE & str_detect(sizeclass1, "75") ~ "075",
      TRUE ~ "raw")
    sizelab1 <- case_when(</pre>
      str_detect(sizeclass1, "25") & str_detect(sizeclass1, "75") ~ "25-75mm",
      str detect(sizeclass1, "35") & str detect(sizeclass1, "75") ~ "35-75mm",
      str_detect(sizeclass1, "25") == FALSE & str_detect(sizeclass1, "75") ~ "\u2265 75mm",
      TRUE ~ "raw")
  if(sizeclass2 != ""){
    size2 <- case_when(</pre>
      str_detect(sizeclass2, "25") & str_detect(sizeclass2, "75") ~ "25to75",
      str_detect(sizeclass2, "35") & str_detect(sizeclass2, "75") ~ "35to75",
      str_detect(sizeclass2, "25") == FALSE & str_detect(sizeclass2, "75") ~ "075",
      TRUE ~ "raw")
    sizelab2 <- case_when(</pre>
```

```
str_detect(sizeclass2, "25") & str_detect(sizeclass2, "75") ~ "25-75mm",
    str_detect(sizeclass2, "35") & str_detect(sizeclass2, "75") ~ "35-75mm",
    str_detect(sizeclass2, "25") == FALSE & str_detect(sizeclass2, "75") ~ "\u2265 75mm",
    TRUE ~ "raw")
}
#Marginal effects plot including random effects
## Hist plot settings
y max <- round(max(data2[!is.na(ShellHeight mm), ShellHeight mm]), -0)+1
y_breaks \leftarrow seq(0, 300, 50)
y_{labs} \leftarrow seq(0, 300, 50)
y_{minor} \leftarrow seq(0, 300, 25)
ylim_upper <- ceiling(y_max/25)*25</pre>
yrdiff1 <- unique(data1$YearDiff)</pre>
yrdiff2 <- unique(data2$YearDiff)</pre>
# function to set year breaks, type == "hist" or "live"
set_breaks <- function(type, data1, data2){</pre>
 ldq <- ifelse(type=="hist", "Estimate", "Exact")</pre>
 maxyr <- max(data1[!is.na(LiveDate) & LiveDate_Qualifier==ldq, LiveDate],</pre>
                data2[!is.na(LiveDate) & LiveDate_Qualifier==ldq, LiveDate])
 minyr <- min(data1[!is.na(LiveDate) & LiveDate_Qualifier==ldq, LiveDate],</pre>
                data2[!is.na(LiveDate) & LiveDate_Qualifier==ldq, LiveDate])
 nyrs <- (maxyr)-(minyr)+1</pre>
  # Creates break intervals for plots based on number of years of data
 if(nyrs > = 40){
    # Set breaks to every 10 years if more than 30 years of data
    brk <- 10
 }else if(nyrs<40 & nyrs>=20){
    # Set breaks to every 5 years if between 30 and 15 years of data
 }else if(nyrs<20 & nyrs>=12){
    # Set breaks to every 3 years if between 15 and 9 years of data
    brk <- 3
 }else if(nyrs<12 & nyrs>=8){
    # Set breaks to every 2 years if between 9 and 6 years of data
    brk <- 2
 }else if(nyrs<8 & nyrs>=3){
    # Set breaks to every year if less than 6 years of data
    brk <- 1
 }else if(nyrs<3){</pre>
   brk <- 1
    maxyr <- maxyr + 1</pre>
    minyr <- minyr - 1
 return(list("seq" = seq(minyr,maxyr,brk),"maxyr" = maxyr, "minyr" = minyr))
yrlist_hist <- set_breaks(type = "hist", data1 = data1, data2 = data2)[["seq"]]</pre>
maxyr_hist <- set_breaks(type = "hist", data1 = data1, data2 = data2)[["maxyr"]]</pre>
```

```
minyr_hist <- set_breaks(type = "hist", data1 = data1, data2 = data2)[["minyr"]]
yrlist_live <- set_breaks(type = "live", data1 = data1, data2 = data2)[["seq"]]</pre>
maxyr_live <- set_breaks(type = "live", data1 = data1, data2 = data2)[["maxyr"]]</pre>
minyr_live <- set_breaks(type = "live", data1 = data1, data2 = data2)[["minyr"]]
## Check data for Exact and Estimate
n_hist1 <- nrow(data1[data1$LiveDate_Qualifier=="Estimate" &</pre>
                          !is.na(data1$ShellHeight mm),])
n_live1 <- nrow(data1[data1$LiveDate_Qualifier=="Exact" &</pre>
                          !is.na(data1$ShellHeight mm),])
n_hist2 <- nrow(data2[data2$LiveDate_Qualifier=="Estimate" &</pre>
                          !is.na(data2$ShellHeight_mm),])
n_live2 <- nrow(data2[data2$LiveDate_Qualifier=="Exact" &</pre>
                          !is.na(data2$ShellHeight_mm),])
set.seed(987)
if(!is.null(models1)){
  liveplot_1 <- plot(conditional_effects(models1[[1]], re_formula=NULL), plot=FALSE)</pre>
if(!is.null(models2)){
  liveplot_2 <- plot(conditional_effects(models2[[1]], re_formula=NULL), plot=FALSE)</pre>
# Set boolean values for whether liveplot162 are available
liveplot1 avail <- class(try(liveplot 1, silent=TRUE)) != "try-error"</pre>
liveplot2_avail <- class(try(liveplot_2, silent=TRUE)) != "try-error"</pre>
# Set ribbon transparency value
a ribb <- 0.2
# Set size and shapes for plots
p_shape <- c("size2"=24, "size1"=21)</pre>
sizelab <- c("size2"=sizelab2, "size1"=sizelab1)</pre>
check <- NA
check1 <- NA
check2 <- NA
if(exists("present1")){
  check1 <- c("size1"="#00374f")
} else{
  check1 <- c("size1"="#FFFFFF")</pre>
if(exists("present2")){
  check2 <- c("size2"="#0094b0")</pre>
} else{
  check2 <- c("size2"="#FFFFFF")</pre>
p_color <- c(check2, check1)</pre>
# Initial plots to set legends
plot_leg <- ggplot() +</pre>
```

```
{if(liveplot1_avail){
    list(geom_ribbon(data=liveplot_1$RelYear$data,
                     aes(x=RelYear+yrdiff1, y=ShellHeight_mm,
                         ymin=lower__, ymax=upper__,
                         fill="size1"), alpha=a_ribb),
         geom_line(data=liveplot_1$RelYear$data,
                   aes(x=RelYear+yrdiff1, y=estimate__, color="size1"),
                   1wd=0.75)
 }} +
  {if(liveplot2 avail){
    list(geom_ribbon(data=liveplot_2$RelYear$data,
                     aes(x=RelYear+yrdiff2, y=ShellHeight_mm,
                         ymin=lower__, ymax=upper__,
                         fill="size2"), alpha=a_ribb),
         geom_line(data=liveplot_2$RelYear$data,
                   aes(x=RelYear+yrdiff2, y=estimate__, color="size2"),
                   1wd=0.75)
 }} +
  geom_point(data=data1[!is.na(RelYear) & !is.na(LiveDate), ],
             aes(x=LiveDate, y=ShellHeight_mm, shape="size1"),
             position=plot_jitter, size=2, color="#333333", fill="#ccccc",
             alpha=0.2, inherit.aes=FALSE) +
  geom_point(data=data2[!is.na(RelYear) & !is.na(LiveDate), ],
             aes(x=LiveDate, y=ShellHeight_mm, shape="size2"),
             position=plot jitter, size=2, color="#333333", fill="#ccccc",
             alpha=0.2, inherit.aes=FALSE) +
 plot_theme +
  theme(legend.position="right") +
  scale_shape_manual(name="Shell heights",
                     values=p_shape,
                     labels=sizelab) +
  scale_color_manual(name="Shell heights",
                     values=p_color,
                     labels=sizelab) +
  scale_fill_manual(name="Shell heights",
                    values=p_color,
                    labels=sizelab)
leg <-get_legend(plot_leg)</pre>
rm(plot_leg)
# Dead oyster shell plot
plot1 <- ggplot() +</pre>
 geom_hline(yintercept=75, size=1, color="grey") +
 {if(n_hist1>0){
    geom_point(data=data1[!is.na(RelYear) &
                            !is.na(LiveDate) &
                            LiveDate_Qualifier=="Estimate", ],
               aes(x=LiveDate, y=ShellHeight_mm, shape="size1"),
               position=plot_jitter, size=2, color="#333333", fill="#ccccc",
               alpha=0.2, inherit.aes=FALSE)
 }} +
  {if(n_hist2>0){
```

```
geom_point(data=data2[!is.na(RelYear) & !is.na(LiveDate) &
                            LiveDate_Qualifier=="Estimate", ],
               aes(x=LiveDate, y=ShellHeight_mm, shape="size2"),
               position=plot_jitter, size=2, color="#333333", fill="#cccccc",
               alpha=0.2, inherit.aes=FALSE)
 }} +
  scale_x_continuous(limits=c(minyr_hist-0.25, maxyr_hist+0.25),
                     breaks=yrlist hist) +
  scale_y_continuous(breaks=y_breaks,
                     labels=y labs, minor breaks=y minor) +
 plot_theme +
  theme(plot.subtitle=element_text(hjust=0, size=10, color="#314963"),
        legend.position="none",
  ) +
 labs(subtitle="Dead Oyster Shells",
       x="Estimated year",
       y="Shell height (mm)") +
  scale_shape_manual(name="Shell heights",
                     values=c("size1"=21, "size2"=24),
                     labels=c(sizelab1, sizelab2)) +
  scale_color_manual(name="Shell heights",
                     values=c("size1"="#00374f", "size2"="#0094b0"),
                     labels=c(sizelab1, sizelab2)) +
  scale_fill_manual(name="Shell heights",
                    values=c("size1"="#00374f", "size2"="#0094b0").
                    labels=c(sizelab1, sizelab2)) +
  coord_cartesian(ylim=c(0, ylim_upper))
# Live oyster shell plot
plot2 <- ggplot() +</pre>
  geom_hline(yintercept=75, size=1, color="grey") +
  \{if(n live1>0)\}
    geom_point(data=data1[!is.na(RelYear) & !is.na(LiveDate) &
                            LiveDate_Qualifier=="Exact", ],
               aes(x=LiveDate, y=ShellHeight_mm, shape="size1"),
               position=plot_jitter, size=2, color="#333333", fill="#ccccc",
               alpha=0.2, inherit.aes=FALSE)
 }} +
 \{if(n_1ive2>0)\}
    geom point(data=data2[!is.na(RelYear) & !is.na(LiveDate) &
                            LiveDate_Qualifier=="Exact", ],
               aes(x=LiveDate, y=ShellHeight_mm, shape="size2"),
               position=plot_jitter, size=2, color="#333333", fill="#ccccc",
               alpha=0.2, inherit.aes=FALSE)
 }} +
  {if(liveplot1_avail){
    list(geom_ribbon(data=liveplot_1$RelYear$data,
                     aes(x=RelYear+yrdiff1, y=ShellHeight_mm,
                         ymin=lower__, ymax=upper__, fill="size1"),
                     alpha=a_ribb),
         geom_line(data=liveplot_1$RelYear$data,
                   aes(x=RelYear+yrdiff1, y=estimate__, color="size1"),
```

```
}} +
  {if(liveplot2_avail){
    list(geom_ribbon(data=liveplot_2$RelYear$data,
                     aes(x=RelYear+yrdiff2, y=ShellHeight_mm,
                         ymin=lower__, ymax=upper__, fill="size2"),
                     alpha=a ribb),
         geom_line(data=liveplot_2$RelYear$data,
                   aes(x=RelYear+yrdiff2, y=estimate__, color="size2"),
                   1wd=0.75)
 }} +
  scale_x_continuous(limits=c(minyr_live-0.25, maxyr_live+0.25),
                     breaks=yrlist_live) +
  scale y continuous(breaks=y breaks,
                     labels=y_labs, minor_breaks=y_minor) +
 plot theme +
  theme(plot.subtitle=element_text(hjust=0, size=10, color="#314963"),
        legend.position="none",
        axis.text.y=element_blank(), #remove y-axis labels
        axis.ticks.y=element_blank(), #remove y-axis ticks
        axis.title.y=element_blank() #removes y-axis title
  ) +
 labs(subtitle="Live Oyster Shells",
       x="Year",
       y="Shell height (mm)") +
  scale shape manual(name="Shell heights",
                     values=c("size1"=21, "size2"=24),
                     labels=c(sizelab1, sizelab2)) +
  scale_color_manual(name="Shell heights",
                     values=c("size1"="#00374f", "size2"="#0094b0"),
                     labels=c(sizelab1, sizelab2)) +
  scale_fill_manual(name="Shell heights",
                    values=c("size1"="#00374f", "size2"="#0094b0"),
                    labels=c(sizelab1, sizelab2)) +
  coord_cartesian(ylim=c(0, ylim_upper))
# Set plot title
plot_title <- ggplot() +</pre>
 labs(title="Oyster Size Class", subtitle=ma) +
 plot_theme +
  theme(plot.subtitle=element_text(hjust=0.5, size=10, color="#314963"),
        panel.border=element_blank(),
        panel.grid.major=element_blank(),
        panel.grid.minor=element blank(), axis.line=element blank())
plot_comb <- ggarrange(plot1, plot2, leg, nrow=1,</pre>
                       widths=c(0.46, 0.39, 0.15))
plot_comb <- ggarrange(plot_title, plot_comb, ncol=1,</pre>
                       heights=c(0.125, 0.875))
ma_abrev <- MA_All[ManagedAreaName==ma, Abbreviation]</pre>
ggsave(paste0("output/Shell_Height/Figures/Oyster_SH_GLMM_", ma_abrev, ".png"),
```

```
plot_comb,
         width=8,
         height=4,
         units="in",
         dpi=200,
         bg="white")
 rm(liveplot_1, liveplot_2)
# Create model results tables and save diagnostic plots
modresultssh <- function(datafile1, models1, datafile2, models2, indicator,</pre>
                         meplotzoom=FALSE){
  datafile1$SizeClass[datafile1$SizeClass=="25to75mm" &
                        datafile1$MA_plotlab==
                         "St. Martins Marsh Aquatic Preserve_Natural"] <-
    "35-75mm"
  sizeclass1 <- unique(datafile1$SizeClass)</pre>
  for(m in seq_along(models1)){
    modelobj <- models1[[m]]</pre>
    oyres i <- setDT(broom.mixed::tidy(modelobj))</pre>
    #tidy() does not like that parameter values have underscores
    #for some reason, so the resulting table is incomplete
    if(nrow(oyres_i[effect=="fixed", ])-nrow(summary(modelobj)$fixed)==-1){
      missingrow <- data.table(effect="fixed",</pre>
                                component="cond",
                                #not sure what "cond" means in the tidy summary.
                                term=rownames(summary(modelobj)$fixed)[2],
                                estimate=summary(modelobj)$fixed$Estimate[2],
                                std.error=summary(modelobj)$fixed$Est.Error[2],
                                conf.low=summary(modelobj)$fixed$`1-95% CI`[2],
                                conf.high=summary(modelobj)$fixed$`u-95% CI`[2])
      oyres_i <- rbind(oyres_i, missingrow) %>% arrange(effect, group)
    }
    setDT(oyres i)
    oyres_i[, `:=` (indicator=indicator,
                    managed_area=unique(datafile1$ManagedAreaName),
                    habitat_class=unique(datafile1$HabitatClassification),
                    size_class=sizeclass1,
                    live_date_qual=ifelse(
                      str_detect(
                        modelobj$file, "_hist"), "Estimate",
                      "Exact"),
                    n_programs=if(class(
                      try(datafile1$LiveDate_Qualifier))!="try-error"){
                      length(unique(
                         datafile1[LiveDate_Qualifier==
                                     ifelse(str_detect(
                                       modelobj$file, "_hist"),
                                       "Estimate", "Exact"),
```

```
ProgramID]))
                  } else{length(unique(datafile1[, ProgramID]))},
                  programs=if(class(try(
                    datafile1$LiveDate_Qualifier)) != "try-error"){
                    list(unique(
                      datafile1[LiveDate Qualifier==
                                   ifelse(
                                     str detect(
                                       modelobj$file,
                                       " hist"),
                                     "Estimate",
                                     "Exact"),
                                 ProgramID]))
                  } else{list(unique(datafile1[, ProgramID]))},
                  filename=modelobj$file)]
  oysterresults <<- rbind(oysterresults, oyres_i)</pre>
  # Save diagnostic plots
  #diagnosticplots(modelobj, indicator,
  #unique(datafile$ManagedAreaName), sizeclass,
  #ifelse(str_detect(modelobj$file, "_hist"), TRUE, FALSE))
}
datafile2$SizeClass[datafile2$SizeClass=="25to75mm" &
                      datafile2$MA plotlab==
                      "St. Martins Marsh Aquatic Preserve_Natural"] <- "35-75mm"
sizeclass2 <- unique(datafile2$SizeClass)</pre>
for(m in seq_along(models2)){
 modelobj <- models2[[m]]</pre>
  oyres_i <- setDT(broom.mixed::tidy(modelobj))</pre>
  #tidy() does not like that parameter values have underscores for
  #some reason, so the resulting table is incomplete
  if(nrow(oyres_i[effect=="fixed", ])-nrow(summary(modelobj)$fixed)==-1){
    missingrow <- data.table(effect="fixed",
                             component="cond",
                              #not sure what "cond" means in the tidy summary.
                             group=NA,
                             term=rownames(summary(modelobj)$fixed)[2],
                             estimate=summary(modelobj)$fixed$Estimate[2],
                             std.error=summary(modelobj)$fixed$Est.Error[2],
                              conf.low=summary(modelobj)$fixed$`1-95% CI`[2],
                             conf.high=summary(modelobj)$fixed$`u-95% CI`[2])
    oyres_i <- rbind(oyres_i, missingrow) %>% arrange(effect, group)
 }
  oyres_i <- oyres_i %>%
    mutate(
      indicator = indicator,
      managed_area = unique(datafile2$ManagedAreaName),
      habitat_class = unique(datafile2$HabitatClassification),
```

```
size_class = sizeclass2,
      live_date_qual = if_else(
        str_detect(modelobj$file, "_hist"), "Estimate", "Exact"
      n_programs = if (class(try(datafile2$LiveDate_Qualifier)) != "try-error") {
        datafile2 %>%
          filter(LiveDate_Qualifier == if_else(str_detect(modelobj$file, "_hist"), "Estimate", "Exact
          pull(ProgramID) %>%
          unique() %>%
          length()
      } else {
        datafile2 %>%
          pull(ProgramID) %>%
          unique() %>%
          length()
      },
      programs = if (class(try(datafile2$LiveDate_Qualifier)) != "try-error") {
        list(datafile2 %>%
               filter(LiveDate_Qualifier == if_else(str_detect(modelobj$file, "_hist"), "Estimate", "
               pull(ProgramID) %>%
               unique())
      } else {
        list(datafile2 %>% pull(ProgramID) %>% unique())
      },
      filename = modelobj$file
 oysterresults <<- rbind(oysterresults, oyres_i)</pre>
  # Save diagnostic plots
  #diagnosticplots(modelobj, indicator,
  #unique(datafile$ManagedAreaName), sizeclass,
  #ifelse(str_detect(modelobj$file, "_hist"), TRUE, FALSE))
}
# Save marginal effects plots
meplotssh(models1, datafile1, sizeclass1, models2, datafile2, sizeclass2,
          unique(datafile1$ManagedAreaName), indicator, meplotzoom)
```

Oyster Shell Height Analysis

Subsets data for that which is related to shell height. The data is further subset for each managed area and shell size class. Appropriate GLMMs are called from file for each shell size class, or are created if they don't exist. Data and models are then sent to the modresultssh function to create figures and save data.

```
# Parallel, num cores setup
ncores <- 4
nchains <- 4
threads <- 10
iter <- 3000
warmup <- 1000

# At least 5 years of data are required in order to run model analyses</pre>
```

```
# Function checks N years of data, returns T or F
suff_years <- function(data){length(unique(data$Year))>=5}
#summarize shell height data
sh_all_sum <- summarySE(oysterraw[!is.na(ShellHeight_mm), ],</pre>
                        measurevar='ShellHeight_mm',
                        groupvars=c('ManagedAreaName', 'LiveDate_Qualifier',
                                    'LiveDate'))
## Apalachicola Bay Aquatic Preserve_Natural ------
#Exclude the five samples that don't have counts less than the "NumberMeasured"
#value for the corresponding program (see variable exploration graphs in the
#25to75mm section for the rationale and graphs for this step.)
numValves <- unique(oysterraw[, c("ProgramID", "RelYear", "counts",</pre>
                                   "QuadIdentifier", "Subtidal", "QuadSize_m2",
                                   "LiveDate_Qualifier", "NumberMeasured_n")])
exclude_samps <- subset(numValves, numValves$NumberMeasured_n=="20" &
                          numValves$counts > 19)$QuadIdentifier
ab_sho25 <- oysterraw[!is.na(ShellHeight_mm) &
                         ShellHeight_mm >= 25 &
                         MA_plotlab=="Apalachicola Bay Aquatic Preserve_Natural" &
                         QuadIdentifier %in% setdiff(
                           oysterraw[!is.na(ShellHeight mm) &
                                        ManagedAreaName==
                                        "Apalachicola Bay Aquatic Preserve",
                                      QuadIdentifier], exclude_samps), ]
saveRDS(ab_sho25, paste0('output/model_results/data/ab_sho25_', Sys.Date(), '.rds'))
### abap-25 to 75mm -----
ab_sh25to75 <- ab_sho25[ShellHeight_mm < 75, ]
saveRDS(ab_sh25to75, paste0('output/model_results/data/ab_sh25to75_', Sys.Date(), '.rds'))
if(suff_years(subset(ab_sh25to75, ab_sh25to75$LiveDate_Qualifier!="Estimate"))){
  ab_sh25to75_glmm <- brm(formula=ShellHeight_mm | trunc(lb=25, ub=75) ~
                            RelYear+QuadSize_m2+(1 | UniversalReefID),
                          data=subset(ab_sh25to75, ab_sh25to75$LiveDate_Qualifier!="Estimate"),
                          family=gaussian, cores=ncores,
                          control=list(adapt_delta=0.995, max_treedepth=20),
                          iter=iter, warmup=warmup, chains=nchains, thin=3, seed=5699,
                          backend="rstan", threads=threading(threads),
                          file="output/model_results/GLMMs/ab_sh25to75_glmm4b.rds")
 models1 <- list(ab_sh25to75_glmm)</pre>
} else {models1 <- NULL}</pre>
# Create model results tables and save diagnostic plots
data1 <- ab_sh25to75</pre>
```

```
### ABAP->75mm ----
ab sho75 <- ab sho25[ShellHeight mm >= 75, ]
saveRDS(ab sho75, paste0('output/model results/data/ab sho75 ', Sys.Date(), '.rds'))
if(suff years(subset(ab sho75, ab sho75$LiveDate Qualifier!="Estimate"))){
  ab sho75 glmm <- brm(formula=ShellHeight mm | trunc(1b=75, ub=250) ~
                         RelYear+(1 | UniversalReefID),
                       data=subset(ab_sho75,
                                   ab_sho75$LiveDate_Qualifier!="Estimate"),
                       family=gaussian, cores=ncores,
                       control= list(adapt_delta=0.995, max_treedepth=20),
                       iter=iter, warmup=warmup, chains=nchains, thin=3, seed=3639,
                       backend="rstan", threads=threading(threads),
                       file="output/model_results/GLMMs/ab_sho75_glmm4b.rds")
 models2 <- list(ab_sho75_glmm)</pre>
} else {models2 <- NULL}</pre>
# Create model results tables and save diagnostic plots and marginal effects plots
data2 <- ab sho75
modresultssh(data1, models1, data2, models2, "Size class", meplotzoom=FALSE)
## Apalachicola National Estuarine Research Reserve_Natural ------
an_sho25 <- oysterraw[!is.na(ShellHeight_mm) &
                        !is.na(LiveDate) &
                        ShellHeight_mm >= 25 &
                        MA_plotlab==
                        "Apalachicola National Estuarine Research Reserve_Natural" &
                        QuadIdentifier %in%
                        setdiff(oysterraw[!is.na(ShellHeight_mm) &
                                            ManagedAreaName==
                                            "Apalachicola National Estuarine Research Reserve",
                                          QuadIdentifier], exclude samps), ]
saveRDS(an_sho25, paste0('output/model_results/data/an_sho25_', Sys.Date(), '.rds'))
### ANERR-25 to 75mm -----
an_sh25to75 <- subset(an_sho25, an_sho25$ShellHeight_mm < 75)</pre>
saveRDS(an_sh25to75, paste0('output/model_results/data/an_sh25to75_',
                            Sys.Date(), '.rds'))
if(suff_years(subset(an_sh25to75, an_sh25to75$LiveDate_Qualifier!="Estimate"))){
  an_sh25to75_glmm <- brm(formula=ShellHeight_mm | trunc(lb=25, ub=75) ~
                            RelYear+QuadSize_m2+(1 | UniversalReefID),
                          data=subset(an_sh25to75, an_sh25to75$LiveDate_Qualifier!="Estimate"),
                          family=gaussian, cores=ncores,
```

```
control=list(adapt_delta=0.995, max_treedepth=20),
                         iter=iter, warmup=warmup, chains=nchains, thin=3, seed=5699,
                         backend="rstan", threads=threading(threads),
                         file="output/model_results/GLMMs/an_sh25to75_glmm4b.rds")
 models1 <- list(an sh25to75 glmm)</pre>
} else {models1 <- NULL}</pre>
# Create model results tables and save diagnostic plots and marginal effects plots
data1 <- an_sh25to75</pre>
### ANERR->75mm -------
an_sho75 <- an_sho25[ShellHeight_mm >= 75, ]
saveRDS(an_sho75, paste0('output/model_results/data/an_sho75_', Sys.Date(), '.rds'))
if(suff_years(subset(an_sho75, an_sho75$LiveDate_Qualifier!="Estimate"))){
 an_sho75_glmm <- brm(formula=ShellHeight_mm | trunc(lb=75, ub=250) ~
                        RelYear+(1 | UniversalReefID),
                      data=subset(an_sho75, an_sho75$LiveDate_Qualifier!=
                                   "Estimate"), family=gaussian, cores=ncores,
                      control= list(adapt_delta=0.995, max_treedepth=20),
                      iter=iter, warmup=warmup, chains=nchains, thin=3, seed=3639,
                      backend="rstan", threads=threading(threads),
                      file="output/model results/GLMMs/an sho75 glmm4b.rds")
 models2 <- list(an sho75 glmm)</pre>
} else {models2 <- NULL}</pre>
# Create model results tables and save diagnostic plots and marginal effects plots
data2 <- an_sho75
#modresults(data, models, "Size class", meplotzoom=TRUE)
modresultssh(data1, models1, data2, models2, "Size class", meplotzoom=FALSE)
## Estero Bay Aquatic Preserve_Natural -----
eb_sho25 <- oysterraw[!is.na(ShellHeight_mm) &
                       ShellHeight_mm >= 25 &
                       MA_plotlab=="Estero Bay Aquatic Preserve_Natural", ]
saveRDS(eb_sho25, paste0('output/model_results/data/eb_sho25_', Sys.Date(), '.rds'))
### EBAP-25 to 75mm ------
eb_sh25to75 <- subset(eb_sho25, eb_sho25$ShellHeight_mm < 75)
eb_sh25to75 <- eb_sh25to75[!is.na(eb_sh25to75$QuadSize_m2),]</pre>
saveRDS(eb_sh25to75, paste0('output/model_results/data/eb_sh25to75_',
                           Sys.Date(), '.rds'))
# Does not update models because this subset of data have NA for QuadSize_m2 values
if(suff_years(subset(eb_sh25to75,eb_sh25to75$LiveDate_Qualifier=="Exact"))){
```

```
eb_sh25to75_glmm <- brm(formula=ShellHeight_mm ~
                           RelYear+QuadSize m2+(0+RelYear | UniversalReefID),
                         data=subset(eb_sh25to75,
                                    eb_sh25to75$LiveDate_Qualifier=="Exact"),
                         family=gaussian, cores=ncores,
                         control= list(adapt_delta=0.995, max_treedepth=20),
                         iter=iter, warmup=warmup, chains=nchains, thin=3, seed=6881,
                         backend="rstan", threads=threading(threads),
                         file="output/model_results/GLMMs/eb_sh25to75_glmm5.rds")
 models1 <- list(eb_sh25to75_glmm)</pre>
} else {models1 <- NULL}</pre>
# Create model results tables and save diagnostic plots and marginal effects plots
data1 <- eb_sh25to75</pre>
### EBAP->75mm -----
eb_sho75 <- eb_sho25[ShellHeight_mm >= 75, ]
saveRDS(eb_sho75, paste0('output/model_results/data/eb_sho75_', Sys.Date(), '.rds'))
if(suff_years(subset(eb_sho75, eb_sho75$LiveDate_Qualifier=="Exact"))){
 eb_sho75_glmm <- brm(formula=ShellHeight_mm ~
                       RelYear+(1 | UniversalReefID),
                      data=subset(eb_sho75, eb_sho75$LiveDate_Qualifier=="Exact"),
                      family=gaussian, cores=ncores,
                      control=list(adapt_delta=0.995, max_treedepth=20), iter=iter,
                      warmup=warmup, chains=nchains, thin=3, seed=3138,
                      backend="rstan", threads=threading(threads),
                      file="output/model_results/GLMMs/eb_sho75_glmm4.rds")
 models2 <- list(eb_sho75_glmm)</pre>
} else {models2 <- NULL}</pre>
# Create model results tables and save diagnostic plots and marginal effects plots
data2 <- eb_sho75
modresultssh(data1, models1, data2, models2, "Size class", meplotzoom=FALSE)
## Guana River Marsh Aquatic Preserve Natural -----
grm_sho25 <- oysterraw[!is.na(ShellHeight_mm) &</pre>
                        ShellHeight mm >= 25 &
                        MA plotlab==
                        "Guana River Marsh Aquatic Preserve_Natural", ]
saveRDS(grm_sho25, paste0('output/model_results/data/grm_sho25_',
                         Sys.Date(), '.rds'))
### GRMAP-25 to 75mm -----
grm_sh25to75 <- subset(grm_sho25, grm_sho25$ShellHeight_mm < 75)
```

```
saveRDS(grm_sh25to75, paste0('output/model_results/data/grm_sh25to75_',
                             Sys.Date(), '.rds'))
if(suff_years(subset(grm_sh25to75, grm_sh25to75$LiveDate_Qualifier=="Exact"))){
  # NumberMeasred_n contains NA, removed from formula
  grm sh25to75 glmm <- brm(formula=ShellHeight mm | trunc(1b=25, ub=75) ~
                             RelYear+(1 | UniversalReefID),
                           data=subset(grm_sh25to75,
                                       grm_sh25to75$LiveDate_Qualifier=="Exact"),
                           family=gaussian, cores=ncores,
                           control= list(adapt_delta=0.8, max_treedepth=10),
                           iter=iter, warmup=warmup, chains=nchains, thin=3,
                           seed=3457, backend="rstan", threads=threading(threads),
                           file="output/model_results/GLMMs/grm_sh25to75_glmm4.rds")
 models1 <- list(grm_sh25to75_glmm)</pre>
} else {models1 <- NULL}</pre>
# Create model results tables and save diagnostic plots and marginal effects plots
data1 <- grm_sh25to75</pre>
### GRMAP->75mm ------
grm_sho75 <- grm_sho25[ShellHeight_mm >= 75, ]
saveRDS(grm_sho75, paste0('output/model_results/data/grm_sho75_',
                          Sys.Date(), '.rds'))
if(suff_years(subset(grm_sho75, grm_sho75$LiveDate_Qualifier=="Exact"))){
  grm_sho75_glmm <- brm(formula=ShellHeight_mm | trunc(lb=75, ub=250) ~
                          RelYear+(1 | UniversalReefID),
                        data=subset(grm_sho75,
                                    grm_sho75$LiveDate_Qualifier=="Exact"),
                        family=gaussian, cores=ncores,
                        control= list(adapt_delta=0.8, max_treedepth=10),
                        iter=iter, warmup=warmup, chains=nchains, thin=3,
                        seed=4352, backend="rstan", threads=threading(threads),
                        file="output/model results/GLMMs/grm sho75 glmm4.rds")
 models2 <- list(grm_sho75_glmm)</pre>
} else {models2 <- NULL}</pre>
# Create model results tables and save diagnostic plots and marginal effects plots
data2 <- grm_sho75</pre>
modresultssh(data1, models1, data2, models2, "Size class", meplotzoom=FALSE)
## Guana Tolomato Matanzas National Estuarine Research Reserve_Natural ------
gtmn_sho25 <- oysterraw[!is.na(ShellHeight_mm) &
                          ShellHeight_mm >= 25 &
                          MA_plotlab==
                          "Guana Tolomato Matanzas National Estuarine Research Reserve_Natural", ]
```

```
saveRDS(gtmn_sho25, paste0('output/model_results/data/gtmn_sho25_',
                          Sys.Date(), '.rds'))
### GTMNERR-25 to 75mm -------------
gtmn_sh25to75 <- subset(gtmn_sho25, gtmn_sho25$ShellHeight_mm < 75)
saveRDS(gtmn sh25to75, paste0('output/model results/data/gtmn sh25to75 ',
                             Sys.Date(), '.rds'))
if(suff_years(subset(gtmn_sh25to75, gtmn_sh25to75$LiveDate_Qualifier != "Estimate"))){
 gtmn_sh25to75_glmm <- brm(formula=ShellHeight_mm | trunc(lb=25, ub=75) ~
                             RelYear+(1 | UniversalReefID),
                           data=subset(gtmn_sh25to75,
                                       gtmn_sh25to75$LiveDate_Qualifier != "Estimate"),
                           family=gaussian, cores=ncores,
                           control=list(adapt_delta=0.8, max_treedepth=20),
                           iter=iter, warmup=warmup, chains=nchains, thin=3,
                           seed=7844, backend="rstan", threads=threading(threads),
                           file="output/model_results/GLMMs/gtmn_sh25to75_glmm5.rds")
 models1 <- list(gtmn_sh25to75_glmm)</pre>
} else {models1 <- NULL}</pre>
# Create model results tables and save diagnostic plots and marginal effects plots
data1 <- gtmn_sh25to75
### GTMNERR->75mm ------------
gtmn_sho75 <- gtmn_sho25[ShellHeight_mm >= 75,]
saveRDS(gtmn_sho75, paste0('output/model_results/data/gtmn_sho75_',
                          Sys.Date(), '.rds'))
if(suff_years(subset(gtmn_sho75, gtmn_sho75$LiveDate_Qualifier != "Estimate"))){
 gtmn_sho75_glmm <- brm(formula=ShellHeight_mm | trunc(lb=75, ub=250) ~</pre>
                          RelYear + (1 | UniversalReefID),
                        data=subset(gtmn sho75,
                                    gtmn_sho75$LiveDate_Qualifier != "Estimate"),
                        family=gaussian,
                        cores=ncores, control=list(adapt_delta=0.995, max_treedepth=20),
                        iter=iter, warmup=warmup, chains=nchains, thin=3,
                        seed=5332, backend="rstan", threads=threading(threads),
                        file="output/model_results/GLMMs/gtmn_sho75_glmm6.rds")
 models2 <- list(gtmn_sho75_glmm)</pre>
} else {models2 <- NULL}</pre>
# Create model results tables and save diagnostic plots and marginal effects plots
data2 <- gtmn_sho75</pre>
modresultssh(data1, models1, data2, models2, "Size class", meplotzoom=FALSE)
## Indian River-Vero Beach to Ft. Pierce Aquatic Preserve_Natural ------
irvbfp_sho25 <- oysterraw[!is.na(ShellHeight_mm) &</pre>
```

```
ShellHeight_mm >= 25 &
                            MA_plotlab==
                            "Indian River-Vero Beach to Ft. Pierce Aquatic Preserve Natural", ]
saveRDS(irvbfp_sho25, paste0('output/model_results/data/irvbfp_sho25_',
                             Sys.Date(), '.rds'))
### IRVBFPAP-25 to 75mm -----
irvbfp_sh25to75 <- subset(irvbfp_sho25, irvbfp_sho25$ShellHeight_mm < 75)</pre>
saveRDS(irvbfp_sh25to75, paste0('output/model_results/data/irvbfp_sh25to75_',
                                Sys.Date(), '.rds'))
if(suff_years(subset(irvbfp_sh25to75, irvbfp_sh25to75$LiveDate_Qualifier != "Estimate"))){
  irvbfp_sh25to75_glmm <- brm(formula=ShellHeight_mm | trunc(1b=25, ub=75) ~</pre>
                                RelYear+(1 | UniversalReefID),
                              data=subset(irvbfp_sh25to75,
                                          irvbfp_sh25to75$LiveDate_Qualifier != "Estimate"),
                              family=gaussian, cores=ncores,
                              control=list(adapt_delta=0.8, max_treedepth=20),
                              iter=iter, warmup=warmup, chains=nchains, thin=3,
                              seed=7844, backend="rstan", threads=threading(threads),
                              file="output/model_results/GLMMs/irvbfp_sh25to75_glmm.rds")
 models1 <- list(irvbfp_sh25to75_glmm)</pre>
} else {models1 <- NULL}</pre>
# Create model results tables and save diagnostic plots and marginal effects plots
data1 <- irvbfp_sh25to75</pre>
### IRVBFPAP->75mm ------
irvbfp_sho75 <- irvbfp_sho25[ShellHeight_mm >= 75, ]
saveRDS(irvbfp_sho75, paste0('output/model_results/data/irvbfp_sho75_',
                             Sys.Date(), '.rds'))
if(suff_years(subset(irvbfp_sho75, irvbfp_sho75$LiveDate_Qualifier != "Estimate"))){
  irvbfp_sho75_glmm <- brm(formula=ShellHeight_mm | trunc(lb=75, ub=250) ~</pre>
                             RelYear + (1 | UniversalReefID),
                           data=subset(irvbfp sho75,
                                       irvbfp_sho75$LiveDate_Qualifier != "Estimate"),
                           family=gaussian,
                           cores=ncores, control=list(adapt_delta=0.995, max_treedepth=20),
                           iter=iter, warmup=warmup, chains=nchains, thin=3,
                           seed=5332, backend="rstan", threads=threading(threads),
                           file="output/model_results/GLMMs/irvbfp_sho75_glmm.rds")
 models2 <- list(irvbfp_sho75_glmm)</pre>
} else {models2 <- NULL}</pre>
# Create model results tables and save diagnostic plots and marginal effects plots
```

```
data2 <- irvbfp_sho75</pre>
modresultssh(data1, models1, data2, models2, "Size class", meplotzoom=FALSE)
## Lemon Bay Aquatic Preserve Natural -----
lb_sho25 <- oysterraw[!is.na(ShellHeight_mm) &</pre>
                       ShellHeight mm >= 25 &
                       MA_plotlab=="Lemon Bay Aquatic Preserve_Natural", ]
saveRDS(1b_sho25, paste0('output/model_results/data/1b_sho25_',
                        Sys.Date(), '.rds'))
### LBAP-25 to 75mm ------
lb_sh25to75 <- subset(lb_sho25, lb_sho25$ShellHeight_mm < 75)</pre>
saveRDS(lb_sh25to75, paste0('output/model_results/data/lb_sh25to75_', Sys.Date(), '.rds'))
if(suff_years(subset(1b_sh25to75, lb_sh25to75$LiveDate_Qualifier != "Estimate"))){
 lb_sh25to75_glmm <- brm(formula=ShellHeight_mm | trunc(lb=25, ub=75) ~
                           RelYear+(1 | UniversalReefID),
                         data=subset(lb sh25to75,
                                    lb_sh25to75$LiveDate_Qualifier != "Estimate"),
                         family=gaussian, cores=ncores,
                         control=list(adapt_delta=0.8, max_treedepth=20),
                         iter=iter, warmup=warmup, chains=nchains, thin=3,
                         seed=7844, backend="rstan", threads=threading(threads),
                         file="output/model_results/GLMMs/lb_sh25to75_glmm.rds")
 models1 <- list(lb_sh25to75_glmm)</pre>
} else {models1 <- NULL}</pre>
# Create model results tables and save diagnostic plots and marginal effects plots
data1 <- lb_sh25to75</pre>
### LBAP->75mm ------
lb_sho75 <- lb_sho25[ShellHeight_mm >= 75, ]
saveRDS(lb_sho75, paste0('output/model_results/data/lb_sho75_', Sys.Date(), '.rds'))
if(suff_years(subset(lb_sho75, lb_sho75$LiveDate_Qualifier != "Estimate"))){
 lb_sho75_glmm <- brm(formula=ShellHeight_mm | trunc(lb=25, ub=75) ~
                           RelYear+(1 | UniversalReefID),
                         data=subset(lb_sho75,
                                    lb_sho75$LiveDate_Qualifier != "Estimate"),
                         family=gaussian, cores=ncores,
                         control=list(adapt_delta=0.8, max_treedepth=20),
                         iter=iter, warmup=warmup, chains=nchains, thin=3,
                         seed=7844, backend="rstan", threads=threading(threads),
                         file="output/model_results/GLMMs/lb_sho75_glmm.rds")
 models2 <- list(lb_sho75_glmm)</pre>
```

```
#Important: note that time-averaging is not accounted for in the model fit for
#the data on shell height >75mm. The measurement error approach I was taking
#did not result in any models that converged, possibly because the combination
#of the data and degree of measurement error leads to multiple possible
#solutions. This means the model reported in this section makes the unrealistic
#assumption that the estimated sample ages are exactly correct.

# Create model results tables and save diagnostic plots and marginal effects plots
data2 <- lb_sho75
modresultssh(data1, models1, data2, models2, "Size class", meplotzoom=FALSE)</pre>
```

Density Analysis

Subsets data for that which is related to density. The data is further subset for each managed area. Appropriate GLMMs are called from file for each shell size class, or are created if they don't exist. Data and models are then sent to the modresults function to create figures and save data.

```
oysterraw$YearDiff <- oysterraw$LiveDate-oysterraw$RelYear</pre>
# #Make a collapsed version of the oysterraw table for density
oysterraw_den <- oysterraw[, c("ProgramID", "ProgramName", "LocationID",</pre>
                               "ProgramLocationID", "QuadIdentifier",
                                "ReefIdentifier", "LiveDate",
                                "LiveDate_Qualifier", "SampleDate", "Year",
                                "Month", "ManagedAreaName", "Region",
                                "SurveyMethod", "HabitatClassification",
                                "QuadSize_m2", "MADup", "Density_m2",
                                "Number_of_Oysters_Counted_Total_Count",
                                "Number_of_Oysters_Counted_Live_Count",
                                "Number_of_Oysters_Counted_Dead_Count",
                                "ObsIndex", "UniversalReefID",
                                "MA plotlab", "Subtidal", "RelYear", "YearDiff")]
oysterraw_den[!is.na(Density_m2), DensIndex := ObsIndex]
oysterraw den[!is.na(Number of Oysters Counted Total Count), NTotIndex := ObsIndex]
oysterraw_den[!is.na(Number_of_Oysters_Counted_Live_Count), NLiveIndex := ObsIndex]
oysterraw_den[!is.na(Number_of_Oysters_Counted_Dead_Count), NDeadIndex := ObsIndex]
oysterraw_den[, ObsIndex := NULL]
oysterraw_den <- unique(oysterraw_den)</pre>
oysterraw_den <- oysterraw_den %>%
  dplyr::group_by(ProgramID, ProgramName, LocationID, ProgramLocationID,
                  QuadIdentifier, ReefIdentifier, LiveDate,
                  LiveDate_Qualifier, SampleDate, Year, Month,
                  ManagedAreaName, Region, SurveyMethod,
                  HabitatClassification, QuadSize_m2, MADup, UniversalReefID,
                  MA plotlab, Subtidal) %>%
  tidyr::fill(Density_m2, Number_of_Oysters_Counted_Total_Count,
              Number_of_Oysters_Counted_Live_Count,
              Number of Oysters Counted Dead Count,
              DensIndex, NTotIndex, NLiveIndex, NDeadIndex) %>%
  tidyr::fill(Density_m2, Number_of_Oysters_Counted_Total_Count,
              Number_of_Oysters_Counted_Live_Count,
```

```
Number_of_Oysters_Counted_Dead_Count,
              DensIndex, NTotIndex, NLiveIndex, NDeadIndex,
              .direction='up') %>%
  dplyr::distinct()
oysterraw_den <- subset(oysterraw_den, !is.na(oysterraw_den$Density_m2) |
                          !is.na(oysterraw_den$Number_of_Oysters_Counted_Total_Count) |
                          !is.na(oysterraw_den$Number_of_Oysters_Counted_Live_Count) |
                          !is.na(oysterraw_den$Number_of_Oysters_Counted_Dead_Count) |
                          !is.na(oysterraw_den$DensIndex)
                          !is.na(oysterraw_den$NTotIndex)
                          !is.na(oysterraw_den$NLiveIndex) |
                          !is.na(oysterraw_den$NDeadIndex))
setDT(oysterraw_den)
#Remove NAs in Density_m2 column
oysterraw_den <- subset(oysterraw_den, !is.na(oysterraw_den$Density_m2))</pre>
#Summarize density data by managed area
den_all_sum <- summarySE(oysterraw_den, measurevar='Density_m2',</pre>
                         groupvars=c('ManagedAreaName', 'Year'))
## Raw density results -----
### Apalachicola Bay Aquatic Preserve Natural ------
ab_n <- subset(oysterraw_den,</pre>
               oysterraw_den$MA_plotlab==
                 "Apalachicola Bay Aquatic Preserve_Natural")
ab_n[, Density_m2 := as.integer(round(Density_m2))]
saveRDS(ab_n, paste0('output/model_results/data/ab_n_', Sys.Date(), '.rds'))
ab_den_glmm <- brm(formula=Density_m2 ~</pre>
                     RelYear+(0+RelYear | UniversalReefID), data=ab n,
                   family=negbinomial, cores=ncores,
                   control= list(adapt_delta=0.995, max_treedepth=20),
                   iter=iter,
                   warmup=warmup, chains=nchains, init=0, thin=3, seed=5512,
                   backend="rstan", threads=threading(threads),
                   file="output/model_results/GLMMs/ab_den_glmm9.rds")
# ab_den_qlmm <- update(ab_den_qlmm,
#
                        newdata = ab n,
#
                        family=neqbinomial, cores=ncores,
#
                        control= list(adapt_delta=0.995, max_treedepth=20),
#
                        iter=iter.
#
                        warmup=warmup, chains=nchains, init=0, thin=3, seed=5512,
#
                        backend="rstan", threads=threading(threads),
                        file="output/model_results/GLMMs/ab_den_qlmm9.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- ab_n
```

```
models <- list(ab_den_glmm)</pre>
modresults(data, models, "Density", meplotzoom=FALSE)
### Apalachicola National Estuarine Research Reserve_Natural ---
an_n <- subset(oysterraw_den,</pre>
               oysterraw den$MA plotlab==
                 "Apalachicola National Estuarine Research Reserve Natural")
an_n[, Density_m2 := as.integer(round(Density_m2))]
saveRDS(an_n, paste0('output/model_results/data/an_n_', Sys.Date(), '.rds'))
an_den_glmm <- brm(formula=Density_m2 ~</pre>
                     RelYear+Subtidal+(0+RelYear | UniversalReefID),
                   data=an_n, family=zero_inflated_negbinomial, cores=ncores,
                   control= list(adapt_delta=0.995, max_treedepth=20), iter=iter,
                   warmup=warmup, chains=nchains, init=0, thin=3, seed=4677,
                   backend="rstan", threads=threading(threads),
                   file="output/model_results/GLMMs/an_den_glmm11.rds")
# an_den_glmm <- update(an_den_glmm,
                        newdata = an_n,
#
                        family=zero_inflated_negbinomial, cores=ncores,
#
                        control= list(adapt_delta=0.995, max_treedepth=20), iter=iter,
#
                        warmup=warmup, chains=nchains, init=0, thin=3, seed=4677,
#
                        backend="rstan", threads=threading(threads),
                        file="output/model_results/GLMMs/an_den_glmm11.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- an n
models <- list(an_den_glmm)</pre>
modresults(data, models, "Density", meplotzoom=FALSE)
### Estero Bay Aquatic Preserve_Natural ------
eb_n <- subset(oysterraw_den,
               oysterraw_den$MA_plotlab=="Estero Bay Aquatic Preserve_Natural")
eb_n[, Density_m2 := as.integer(round(Density_m2))]
saveRDS(eb_n, paste0('output/model_results/data/eb_n_', Sys.Date(), '.rds'))
eb_den_glmm <- brm(formula=Density_m2 ~</pre>
                     RelYear+(1 | UniversalReefID), data=eb_n,
                   family=zero inflated negbinomial, cores=ncores,
                   control= list(adapt_delta=0.995, max_treedepth=20), iter=iter,
                   warmup=warmup, chains=nchains, init=0, thin=3, seed=1298,
                   backend="rstan", threads=threading(threads),
                   file="output/model_results/GLMMs/eb_den_glmm10.rds")
# eb_den_qlmm <- update(eb_den_qlmm,
#
                        newdata = eb_n,
#
                        family=zero_inflated_neqbinomial, cores=ncores,
#
                        control= list(adapt_delta=0.995, max_treedepth=20), iter=iter,
                        warmup=warmup, chains=nchains, init=0, thin=3, seed=1298,
```

```
backend="rstan", threads=threading(threads),
#
                        file="output/model results/GLMMs/eb den glmm10.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- eb_n
models <- list(eb den glmm)</pre>
modresults(data, models, "Density", meplotzoom=FALSE)
### Guana River Marsh Aquatic Preserve Natural -----
grm_n <- subset(oysterraw_den,</pre>
                oysterraw_den$MA_plotlab==
                  "Guana River Marsh Aquatic Preserve_Natural")
grm_n[, Density_m2 := as.integer(round(Density_m2))]
saveRDS(grm_n, paste0('output/model_results/data/grm_n_',
                      Sys.Date(), '.rds'))
grm_den_glmm <- brm(formula=Density_m2 ~</pre>
                      RelYear+(1 | UniversalReefID), data=grm_n,
                    family=zero_inflated_negbinomial, cores=2,
                    control= list(adapt_delta=0.995, max_treedepth=20),
                    iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
                    seed=9875, backend="rstan", threads=threading(threads),
                    file="output/model_results/GLMMs/grm_den_glmm6.rds")
# grm_den_glmm <- update(grm_den_glmm,</pre>
                         newdata = qrm n,
#
                         family=zero_inflated_negbinomial, cores=2,
#
                         control= list(adapt_delta=0.995, max_treedepth=20),
#
                         iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
                         seed=9875, backend="rstan", threads=threading(threads),
#
#
                         file="output/model_results/GLMMs/qrm_den_qlmm6.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- grm_n
models <- list(grm_den_glmm)</pre>
modresults(data, models, "Density", meplotzoom=FALSE)
### Guana Tolomato Matanzas National Estuarine Research Reserve_Natural ----
gtmn_n <- subset(oysterraw_den,</pre>
                 oysterraw_den$MA_plotlab==
                   "Guana Tolomato Matanzas National Estuarine Research Reserve Natural")
gtmn n[, Density m2 := as.integer(round(Density m2))]
saveRDS(gtmn_n, paste0('output/model_results/data/gtmn_n_', Sys.Date(), '.rds'))
gtmn_den_glmm <- brm(formula=Density_m2 ~</pre>
                       RelYear+(1 | UniversalReefID),
                     data=gtmn_n, family=zero_inflated_negbinomial, cores=ncores,
                     control= list(adapt_delta=0.995, max_treedepth=20),
                     iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
                     seed=3647, backend="rstan", threads=threading(threads),
                     file="output/model_results/GLMMs/gtmn_den_glmm18.rds")
```

```
# gtmn_den_glmm <- update(gtmn_den_glmm,
#
                           newdata = qtmn n,
#
                           family=zero_inflated_negbinomial, cores=ncores,
#
                           control= list(adapt delta=0.995, max treedepth=20),
#
                           iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
#
                           seed=3647, backend="rstan", threads=threading(threads),
                           file="output/model_results/GLMMs/gtmn_den_glmm18.rds")
#
# Create model results tables and save diagnostic plots and marginal effects plots
data <- gtmn_n
models <- list(gtmn_den_glmm)</pre>
modresults(data, models, "Density", meplotzoom=FALSE)
### Indian River-Vero Beach to Ft. Pierce Aquatic Preserve_Natural --
irvb_n <- subset(oysterraw_den,</pre>
                 oysterraw_den$MA_plotlab==
                   "Indian River-Vero Beach to Ft. Pierce Aquatic Preserve_Natural")
irvb_n[, Density_m2 := as.integer(round(Density_m2))]
saveRDS(irvb_n, paste0('output/model_results/data/irvb_n_',
                       Sys.Date(), '.rds'))
irvb den glmm <- brm(formula=Density m2 ~</pre>
                       RelYear+(0+RelYear | UniversalReefID),
                     data=irvb n,
                     family=negbinomial, cores=ncores,
                     control= list(adapt_delta=0.995, max_treedepth=20),
                     iter=iter,
                     warmup=warmup, chains=nchains, init=0, thin=3, seed=5512,
                     backend="rstan", threads=threading(threads),
                     file="output/model_results/GLMMs/irvb_den_glmm9.rds")
# irvb_den_glmm <- update(irvb_den_glmm,</pre>
#
                          newdata = irvb_n,
#
                           family=negbinomial, cores=ncores,
#
                           control= list(adapt_delta=0.995, max_treedepth=20),
#
                           iter=iter,
#
                           warmup=warmup, chains=nchains, init=0, thin=3, seed=5512,
#
                           backend="rstan", threads=threading(threads),
#
                           file="output/model results/GLMMs/irvb den glmm9.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- irvb_n
models <- list(irvb_den_glmm)</pre>
modresults(data, models, "Density", meplotzoom=FALSE)
### Jensen Beach to Jupiter Inlet Aquatic Preserve_Natural ----
jbji_n <- subset(oysterraw_den,</pre>
                 oysterraw_den$MA_plotlab==
                   "Jensen Beach to Jupiter Inlet Aquatic Preserve_Natural")
jbji_n[, Density_m2 := as.integer(round(Density_m2))]
saveRDS(jbji_n, paste0('output/model_results/data/jbji_n_',
```

```
Sys.Date(), '.rds'))
jbji_den_glmm <- brm(formula=Density_m2 ~</pre>
                       RelYear+(0+RelYear | UniversalReefID),
                     data=jbji_n,
                     family=negbinomial, cores=ncores,
                     control= list(adapt_delta=0.995, max_treedepth=20),
                     iter=iter,
                     warmup=warmup, chains=nchains, init=0, thin=3, seed=5512,
                     backend="rstan", threads=threading(threads),
                     file="output/model_results/GLMMs/jbji_den_glmm9.rds")
# jbji_den_glmm <- update(irvb_den_glmm,</pre>
                          newdata = jbji_n,
#
                          family=neqbinomial, cores=ncores,
#
                          control= list(adapt_delta=0.995, max_treedepth=20),
#
                          iter=iter.
#
                          warmup=warmup, chains=nchains, init=0, thin=3, seed=5512,
#
                          backend="rstan", threads=threading(threads),
#
                          file="output/model_results/GLMMs/jbji_den_qlmm9.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- jbji_n
models <- list(jbji_den_glmm)</pre>
modresults(data, models, "Density", meplotzoom=FALSE)
### Lemon Bay Aquatic Preserve Natural -----
lb_n <- subset(oysterraw_den,</pre>
               oysterraw_den$MA_plotlab=="Lemon Bay Aquatic Preserve_Natural")
lb_n[, Density_m2 := as.integer(round(Density_m2))]
saveRDS(lb_n, paste0('output/model_results/data/lb_n_', Sys.Date(), '.rds'))
lb_den_glmm <- brm(formula=Density_m2 ~</pre>
                     RelYear+(1 | ReefIdentifier), data=lb_n,
                   family=zero_inflated_negbinomial, cores=2,
                   control= list(adapt_delta=0.995, max_treedepth=20),
                   iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
                   seed=4612, backend="rstan", threads=threading(threads),
                   file="output/model results/GLMMs/lb den glmm6.rds")
# lb_den_glmm <- update(lb_den_glmm,
#
                        newdata = lb n,
#
                        family=zero_inflated_negbinomial, cores=2,
#
                        control= list(adapt_delta=0.995, max_treedepth=20),
#
                        iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
#
                        seed=4612, backend="rstan", threads=threading(threads),
                        file="output/model_results/GLMMs/lb_den_qlmm6.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- lb_n
models <- list(lb_den_glmm)</pre>
modresults(data, models, "Density", meplotzoom=FALSE)
```

```
### Pine Island Sound Aquatic Preserve_Natural -----
oysterraw_den[str_detect(MA_plotlab, "Pine Island Sound"), `:=`
              (MA plotlab=ifelse(str detect(ProgramLocationID, "Reference")
                                   str_detect(ProgramLocationID, "Control"),
                                 "Pine Island Sound Aquatic Preserve Natural",
                                 "Pine Island Sound Aquatic Preserve_Restored"),
                HabitatClassification=ifelse(str detect(ProgramLocationID,
                                                         "Reference") |
                                               str_detect(ProgramLocationID,
                                                           "Control"),
                                              "Natural", "Restored"))]
pis_n <- subset(oysterraw_den,</pre>
                oysterraw_den$MA_plotlab==
                  "Pine Island Sound Aquatic Preserve_Natural")
pis_n[, `:=` (Density_m2=as.integer(round(Density_m2)),
              Treatment=ifelse(UniversalReefID==170711,
                               "Reference", "Control"))]
saveRDS(pis_n, paste0('output/model_results/data/pis_n_', Sys.Date(), '.rds'))
pis den glmm <- brm(formula=Density m2 ~
                      RelYear+(0+RelYear | UniversalReefID),
                    data=pis_n, family=zero_inflated_negbinomial, cores=ncores,
                    control= list(adapt_delta=0.995, max_treedepth=20),
                    iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
                    seed=5243, backend="rstan", threads=threading(threads),
                    file="output/model_results/GLMMs/pis_den_glmm9.rds")
# pis_den_glmm <- update(pis_den_glmm,</pre>
                         newdata = pis_n,
#
                         family=zero_inflated_neqbinomial, cores=ncores,
#
                         control= list(adapt_delta=0.995, max_treedepth=20),
#
                         iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
                         seed=5243, backend="rstan", threads=threading(threads),
#
                         file="output/model_results/GLMMs/pis_den_glmm9.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- pis n
models <- list(pis_den_glmm)</pre>
modresults(data, models, "Density", meplotzoom=FALSE)
### Pine Island Sound Aquatic Preserve_Restored ------
pisr_n <- subset(oysterraw_den,</pre>
                 oysterraw_den$MA_plotlab==
                   "Pine Island Sound Aquatic Preserve_Restored")
pisr_n[, `:=` (Density_m2=as.integer(round(Density_m2)),
               Treatment=ifelse(UniversalReefID==170711,
                                "Reference", "Control"))]
saveRDS(pisr_n, paste0('output/model_results/data/pisr_n_', Sys.Date(), '.rds'))
pisr_den_glmm <- brm(formula=Density_m2 ~</pre>
```

```
RelYear+QuadSize_m2, data=pisr_n,
                     family=zero_inflated_negbinomial,
                     prior=set prior("uniform(0,5)", class="b", lb=0, ub=5),
                     cores=ncores, control= list(adapt_delta=0.995, max_treedepth=20),
                     iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
                     seed=8441, backend="rstan", threads=threading(threads),
                     file="output/model results/GLMMs/pisr den glmm12.rds")
# pisr_den_glmm <- update(pisr_den_glmm,</pre>
                          newdata = pisr_n,
#
                          family=zero_inflated_neqbinomial,
#
                          prior=set_prior("uniform(0,5)", class="b", lb=0, ub=5),
#
                          cores=ncores, control= list(adapt_delta=0.995, max_treedepth=20),
#
                          iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
#
                          seed=8441, backend="rstan", threads=threading(threads),
#
                          file="output/model_results/GLMMs/pisr_den_qlmm12.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- pisr_n
models <- list(pisr_den_glmm)</pre>
modresults(data, models, "Density", meplotzoom=FALSE)
PI_R <- nrow(subset(oysterresults,
                    oysterresults$managed_area==
                      "Pine Island Sound Aquatic Preserve" &
                      oysterresults$indicator=="Density" &
                      oysterresults$habitat_class=="Restored"))
oysterresults$group[is.na(oysterresults$group)] <- NA
if(PI_R>0){
  oysterresults$group[oysterresults$managed_area==
                        "Pine Island Sound Aquatic Preserve" &
                        oysterresults$indicator=="Density" &
                        oysterresults$habitat_class=="Restored"] <-
    c(NA, NA, NA)
  oysterresults$term[oysterresults$managed_area==
                       "Pine Island Sound Aquatic Preserve" &
                       oysterresults$indicator=="Density" &
                       oysterresults$habitat_class=="Restored"] <-</pre>
    c("(Intercept)", "RelYear", "QuadSize_m2")
}
```

Percent Live Analysis

Subsets data for that which is related to percent live. The data is further subset for each managed area. Appropriate GLMMs are called from file for each shell size class, or are created if they don't exist. Data and models are then sent to the modresults function to create figures and save data.

```
"LiveDate_Qualifier", "SampleDate", "Year",
                               "Month", "ManagedAreaName", "Region",
                               "SurveyMethod", "PercentLiveMethod",
                               "HabitatClassification", "QuadSize m2", "MADup",
                               "PercentLive pct",
                               "Number_of_Oysters_Counted_Total_Count",
                               "Number of Oysters Counted Live Count",
                               "Number_of_Oysters_Counted_Dead_Count",
                               "ObsIndex", "UniversalReefID",
                               "MA_plotlab", "Subtidal", "RelYear", "YearDiff")]
oysterraw_pct[!is.na(PercentLive_pct), PctIndex := ObsIndex]
oysterraw_pct[!is.na(Number_of_Oysters_Counted_Total_Count),
              NTotIndex := ObsIndex]
oysterraw_pct[!is.na(Number_of_Oysters_Counted_Live_Count),
              NLiveIndex := ObsIndex]
oysterraw_pct[!is.na(Number_of_Oysters_Counted_Dead_Count),
              NDeadIndex := ObsIndex]
oysterraw_pct[, ObsIndex := NULL]
oysterraw_pct <- unique(oysterraw_pct)</pre>
oysterraw pct <- oysterraw pct %>%
  dplyr::group_by(ProgramID, ProgramName, ProgramLocationID, QuadIdentifier,
                  ReefIdentifier, LiveDate, LiveDate_Qualifier, SampleDate,
                  Year, Month, ManagedAreaName, Region, SurveyMethod,
                  PercentLiveMethod, HabitatClassification, QuadSize_m2,
                  MADup, UniversalReefID, MA plotlab, Subtidal,
                  RelYear) %>%
  tidyr::fill(PercentLive_pct, Number_of_Oysters_Counted_Total_Count,
              Number_of_Oysters_Counted_Live_Count,
              Number_of_Oysters_Counted_Dead_Count,
              PctIndex, NTotIndex, NLiveIndex, NDeadIndex) %>%
  tidyr::fill(PercentLive_pct, Number_of_Oysters_Counted_Total_Count,
              Number of Oysters Counted Live Count,
              Number_of_Oysters_Counted_Dead_Count,
              PctIndex, NTotIndex, NLiveIndex, NDeadIndex,
              .direction='up') %>%
  dplyr::distinct()
oysterraw pct <- subset(oysterraw pct, !is.na(oysterraw pct$PercentLive pct) |
                          !is.na(oysterraw_pct$Number_of_Oysters_Counted_Total_Count) |
                          !is.na(oysterraw_pct$Number_of_Oysters_Counted_Live_Count) |
                          !is.na(oysterraw_pct$Number_of_Oysters_Counted_Dead_Count) |
                          !is.na(oysterraw_pct$PctIndex) |
                          !is.na(oysterraw_pct$NTotIndex) |
                          !is.na(oysterraw_pct$NLiveIndex) |
                          !is.na(oysterraw_pct$NDeadIndex))
setDT(oysterraw_pct)
#Calculate PercentLive_pct values for some ProgramIDs where it is missing.
#Couldn't include at the start of the script because need to use the counts columns
#rather than the QuadSize_m2 column which is filled for the whole combined table.
oysterraw_pct[ProgramID==972 | ProgramID==4014 | ProgramID==4044,
              PercentLive_pct :=
```

```
(Number_of_Oysters_Counted_Live_Count/
                    (Number_of_Oysters_Counted_Live_Count+
                      Number_of_Oysters_Counted_Dead_Count) * 100)]
#Filter NAs for PercentLive_pct (these are related to 1) programs that do
#counts to measure density, but do not estimate percent live and
#2) Programs that are listed as measuring percent live by a Point-intercept
#method, which cannot be calculated from counts.
oysterraw_pct <- oysterraw_pct[!is.na(PercentLive_pct), ]</pre>
#Add column of decimal versions of percent live values
oysterraw_pct[, PercentLive_dec := PercentLive_pct/100]
#Summarize percent live values
pct_all_sum <- summarySE(oysterraw_pct, measurevar='PercentLive_pct',</pre>
                         groupvars=c('ManagedAreaName', 'Year', 'PercentLiveMethod'))
## Apalachicola Bay Aquatic Preserve_Natural ------
abap_p <- subset(oysterraw_pct,</pre>
                 oysterraw_pct$MA_plotlab==
                   "Apalachicola Bay Aquatic Preserve_Natural")
saveRDS(abap_p, paste0('output/model_results/data/abap_p_', Sys.Date(), '.rds'))
abap_p_binom <- data.table(ProgramID=character(), ProgramLocationID=character(),</pre>
                           QuadIdentifier=character(), Year=integer(),
                           ManagedAreaName=character(),
                           PercentLiveMethod=character(),
                           UniversalReefID=factor(), Region=character(),
                           MA_plotlab=character(), RelYear=integer(),
                           PercentLive_pct=numeric(), LiveObs=logical())
for(i in 1:nrow(abap_p)){
  dat_i <- abap_p[i, c("ProgramID", "ProgramLocationID", "QuadIdentifier",</pre>
                        "Year", "ManagedAreaName", "PercentLiveMethod",
                       "UniversalReefID", "Region", "MA_plotlab", "RelYear",
                       "PercentLive_pct")]
 dat_1 <- purr::map_dfr(seq_len(round(dat_i$PercentLive_pct[1], digits=0)),</pre>
                           ~dat_i[, LiveObs := 1])
  dat_nl <- purrr::map_dfr(seq_len((100-round(dat_i$PercentLive_pct[1]),</pre>
                                               digits=0))),
                           ~dat_i[, LiveObs := 0])
 dat <- rbind(dat_1, dat_nl)</pre>
 abap_p_binom <- rbind(abap_p_binom, dat)</pre>
saveRDS(abap_p_binom,
        paste0('output/model_results/data/abap_p_binom_',
               Sys.Date(), '.rds'))
abap_pct_glmm <- brm(formula=LiveObs ~ RelYear+(1 | UniversalReefID),
                     data=abap_p_binom, family=bernoulli, cores=ncores,
                     control= list(adapt_delta=0.995,
                                    max_treedepth=20),
```

```
iter=iter, warmup=warmup, chains=nchains, init=0,
                      thin=3,
                      seed=4331, backend="rstan",
                      threads=threading(threads),
                     file="output/model results/GLMMs/abap pct glmm3.rds")
# abap_pct_glmm <- update(abap_pct_glmm,</pre>
                           newdata = abap_p_binom,
#
                           family=bernoulli, cores=ncores,
#
                           control= list(adapt_delta=0.995,
#
                                         max_treedepth=20),
#
                           iter=iter, warmup=warmup, chains=nchains, init=0,thin=3,
#
                           seed=4331, backend="rstan", threads=threading(threads),
#
                           file="output/model_results/GLMMs/abap_pct_qlmm3.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- abap_p
models <- list(abap_pct_glmm)</pre>
modresults(data, models, "Percent live", meplotzoom=FALSE)
## Apalachicola National Estuarine Research Reserve Natural -----
anerr_p <- subset(oysterraw_pct,</pre>
                  oysterraw_pct$MA_plotlab==
                     "Apalachicola National Estuarine Research Reserve_Natural")
saveRDS(anerr_p, paste0('output/model_results/data/anerr_p_', Sys.Date(), '.rds'))
anerr_p_binom <- data.table(ProgramID=character(), ProgramLocationID=character(),</pre>
                             QuadIdentifier=character(), Year=integer(),
                             ManagedAreaName=character(),
                             PercentLiveMethod=character(),
                             UniversalReefID=factor(), Region=character(),
                             MA_plotlab=character(), RelYear=integer(),
                             PercentLive_pct=numeric(), LiveObs=logical())
for(i in 1:nrow(anerr p)){
  dat_i <- anerr_p[i, c("ProgramID", "ProgramLocationID", "QuadIdentifier",</pre>
                         "Year", "ManagedAreaName", "PercentLiveMethod",
                         "UniversalReefID", "Region", "MA plotlab", "RelYear",
                         "PercentLive pct")]
 dat_1 <- purrr::map_dfr(seq_len(round(dat_i$PercentLive_pct[1], digits=0)),</pre>
                           ~dat_i[, LiveObs := 1])
  dat_nl <- purrr::map_dfr(seq_len((100-round(dat_i$PercentLive_pct[1],</pre>
                                                digits=0))),
                            ~dat_i[, LiveObs := 0])
  dat <- rbind(dat_1, dat_nl)</pre>
  anerr_p_binom <- rbind(anerr_p_binom, dat)</pre>
saveRDS(anerr_p_binom,
        paste0('output/model_results/data/anerr_p_binom_',
               Sys.Date(), '.rds'))
anerr_pct_glmm <- brm(formula=LiveObs ~ RelYear+(1 | UniversalReefID),</pre>
                      data=anerr_p_binom, family=bernoulli, cores=ncores,
```

```
control= list(adapt_delta=0.995,
                                     max treedepth=20),
                       iter=iter, warmup=warmup, chains=nchains, init=0,
                       seed=4331, backend="rstan",
                       threads=threading(threads),
                       file="output/model_results/GLMMs/anerr_pct_glmm3.rds")
# anerr_pct_glmm <- update(anerr_pct_glmm,</pre>
#
                            newdata = anerr_p_binom,
#
                            family=bernoulli, cores=ncores,
#
                            control= list(adapt_delta=0.995,
#
                                           max treedepth=20),
#
                            iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
#
                            seed=4331, backend="rstan", threads=threading(threads),
#
                            file="output/model_results/GLMMs/anerr_pct_glmm3.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- anerr_p</pre>
models <- list(anerr_pct_glmm)</pre>
modresults(data, models, "Percent live", meplotzoom=FALSE)
## Guana River Marsh Aquatic Preserve Natural ----
grm p <- subset(oysterraw pct,</pre>
                oysterraw_pct$MA_plotlab==
                   "Guana River Marsh Aquatic Preserve Natural")
saveRDS(grm_p, paste0('output/model_results/data/grm_p_', Sys.Date(), '.rds'))
grm_p_binom <- data.table(ProgramID=character(), ProgramLocationID=character(),</pre>
                           QuadIdentifier=character(), Year=integer(),
                           ManagedAreaName=character(),
                           PercentLiveMethod=character(),
                           UniversalReefID=factor(), Region=character(),
                           MA_plotlab=character(), RelYear=integer(),
                           PercentLive_pct=numeric(), LiveObs=logical())
for(i in 1:nrow(grm_p)){
  dat_i <- grm_p[i, c("ProgramID", "ProgramLocationID", "QuadIdentifier",</pre>
                       "Year", "ManagedAreaName", "PercentLiveMethod",
                       "UniversalReefID", "Region", "MA_plotlab", "RelYear",
                       "PercentLive_pct")]
 dat_1 <- purrr::map_dfr(seq_len(round(dat_i$PercentLive_pct[1], digits=0)),</pre>
                           ~dat i[, LiveObs := 1])
  dat_nl <- purrr::map_dfr(seq_len((100-round(dat_i$PercentLive_pct[1],</pre>
                                                digits=0))),
                            ~dat_i[, LiveObs := 0])
  dat <- rbind(dat_1, dat_n1)</pre>
  grm_p_binom <- rbind(grm_p_binom, dat)</pre>
saveRDS(grm_p_binom, paste0('output/model_results/data/grm_p_binom_',
                             Sys.Date(), '.rds'))
grm_pct_glmm <- brm(formula=LiveObs ~ RelYear+(1 | UniversalReefID),</pre>
```

```
data=grm_p_binom, family=bernoulli, cores=ncores,
                    control= list(adapt_delta=0.995, max_treedepth=20),
                    iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
                    seed=4331, backend="rstan", threads=threading(threads),
                    file="output/model results/GLMMs/grm pct glmm3.rds")
# grm_pct_glmm <- update(grm_pct_glmm,</pre>
                         newdata = grm_p_binom,
                          family=bernoulli, cores=ncores,
#
#
                          control= list(adapt_delta=0.995, max_treedepth=20),
#
                          iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
#
                          seed=4331, backend="rstan", threads=threading(threads),
                         file="output/model_results/GLMMs/grm_pct_glmm3.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- grm_p
models <- list(grm_pct_glmm)</pre>
modresults(data, models, "Percent live", meplotzoom=FALSE)
## Guana Tolomato Matanzas National Estuarine Research
## Reserve Natural -----
gtm_p <- subset(oysterraw_pct,</pre>
                oysterraw_pct$MA_plotlab==
                  "Guana Tolomato Matanzas National Estuarine Research Reserve_Natural")
saveRDS(gtm_p, paste0('output/model_results/data/gtm_p_', Sys.Date(), '.rds'))
gtm_p_binom <- data.table(ProgramID=character(), ProgramLocationID=character(),</pre>
                           QuadIdentifier=character(), Year=integer(),
                          ManagedAreaName=character(),
                          PercentLiveMethod=character(),
                          UniversalReefID=factor(), Region=character(),
                          MA_plotlab=character(), RelYear=integer(),
                           PercentLive_pct=numeric(), LiveObs=logical())
for(i in 1:nrow(gtm_p)){
  dat_i <- gtm_p[i, c("ProgramID", "ProgramLocationID", "QuadIdentifier",</pre>
                       "Year", "ManagedAreaName", "PercentLiveMethod",
                      "UniversalReefID", "Region", "MA_plotlab", "RelYear",
                      "PercentLive pct")]
  dat_1 <- purrr::map_dfr(seq_len(round(dat_i$PercentLive_pct[1], digits=0)),</pre>
                           ~dat_i[, LiveObs := 1])
  dat_nl <- purrr::map_dfr(seq_len((100-round(dat_i$PercentLive_pct[1],</pre>
                                               digits=0))),
                            ~dat_i[, LiveObs := 0])
  dat <- rbind(dat_1, dat_nl)</pre>
 gtm_p_binom <- rbind(gtm_p_binom, dat)</pre>
saveRDS(gtm_p_binom,
        paste0('output/model_results/data/gtm_p_binom_',
               Sys.Date(), '.rds'))
gtm_pct_glmm <- brm(formula=LiveObs ~ RelYear+(1 | UniversalReefID),</pre>
                    data=gtm_p_binom, family=bernoulli, cores=ncores,
```

```
control= list(adapt_delta=0.995,
                                   max_treedepth=20),
                    iter=iter, warmup=warmup, chains=nchains, init=0,
                    seed=4331, backend="rstan",
                    threads=threading(threads),
                    file="output/model_results/GLMMs/gtm_pct_glmm3.rds")
# gtm_pct_glmm <- update(gtm_pct_glmm,</pre>
#
                          newdata = gtm_p_binom,
#
                          family=bernoulli, cores=ncores,
#
                          control= list(adapt_delta=0.995,
#
                                        max_treedepth=20),
#
                          iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
#
                          seed=4331, backend="rstan", threads=threading(threads),
#
                          file="output/model_results/GLMMs/gtm_pct_glmm3.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- gtm_p
models <- list(gtm_pct_glmm)</pre>
modresults(data, models, "Percent live", meplotzoom=FALSE)
## Indian River-Vero Beach to Ft. Pierce Aquatic Preserve_Natural ----
irvb p <- subset(oysterraw pct,</pre>
                 oysterraw pct$MA plotlab==
                   "Indian River-Vero Beach to Ft. Pierce Aquatic Preserve_Natural")
saveRDS(irvb_p, paste0('output/model_results/data/irvb_p_', Sys.Date(), '.rds'))
irvb_p_binom <- data.table(ProgramID=character(), ProgramLocationID=character(),</pre>
                            QuadIdentifier=character(), Year=integer(),
                            ManagedAreaName=character(),
                            PercentLiveMethod=character(),
                            UniversalReefID=factor(), Region=character(),
                            MA_plotlab=character(), RelYear=integer(),
                            PercentLive_pct=numeric(), LiveObs=logical())
for(i in 1:nrow(irvb_p)){
  dat_i <- irvb_p[i, c("ProgramID", "ProgramLocationID", "QuadIdentifier",</pre>
                        "Year", "ManagedAreaName", "PercentLiveMethod",
                        "UniversalReefID", "Region", "MA_plotlab", "RelYear",
                        "PercentLive_pct")]
 dat_1 <- purrr::map_dfr(seq_len(round(dat_i$PercentLive_pct[1], digits=0)),</pre>
                           ~dat i[, LiveObs := 1])
  dat_nl <- purrr::map_dfr(seq_len((100-round(dat_i$PercentLive_pct[1],</pre>
                                                digits=0))),
                            ~dat_i[, LiveObs := 0])
  dat <- rbind(dat_1, dat_nl)</pre>
  irvb_p_binom <- rbind(irvb_p_binom, dat)</pre>
saveRDS(irvb_p_binom, paste0('output/model_results/data/irvb_p_binom_',
                              Sys.Date(), '.rds'))
```

```
irvb_pct_glmm <- brm(formula=LiveObs ~ RelYear+(1 | UniversalReefID),</pre>
                     data=irvb_p_binom, family=bernoulli, cores=ncores,
                     control= list(adapt_delta=0.995, max_treedepth=20),
                     iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
                     seed=4331, backend="rstan", threads=threading(threads),
                     file="output/model results/GLMMs/irvb pct glmm3.rds")
# irvb_pct_glmm <- update(irvb_pct_glmm,</pre>
                           newdata = irvb_p_binom,
#
#
                           family=bernoulli, cores=ncores,
#
                           control= list(adapt_delta=0.995, max_treedepth=20),
#
                           iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
                           seed=4331, backend="rstan", threads=threading(threads),
#
                           file="output/model_results/GLMMs/irvb_pct_qlmm3.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- irvb_p
models <- list(irvb_pct_glmm)</pre>
modresults(data, models, "Percent live", meplotzoom=FALSE)
## Jensen Beach to Jupiter Inlet Aquatic Preserve_Natural -----
jbji_p <- subset(oysterraw_pct,</pre>
                 oysterraw pct$MA plotlab==
                   "Jensen Beach to Jupiter Inlet Aquatic Preserve Natural")
saveRDS(jbji_p, paste0('output/model_results/data/jbji_p_', Sys.Date(), '.rds'))
jbji_p_binom <- data.table(ProgramID=character(), ProgramLocationID=character(),
                            QuadIdentifier=character(), Year=integer(),
                            ManagedAreaName=character(),
                            PercentLiveMethod=character(),
                            UniversalReefID=factor(), Region=character(),
                            MA_plotlab=character(), RelYear=integer(),
                            PercentLive_pct=numeric(), LiveObs=logical())
for(i in 1:nrow(jbji_p)){
  dat_i <- jbji_p[i, c("ProgramID", "ProgramLocationID", "QuadIdentifier",</pre>
                        "Year", "ManagedAreaName", "PercentLiveMethod",
                        "UniversalReefID", "Region", "MA_plotlab", "RelYear",
                        "PercentLive_pct")]
 dat_1 <- purrr::map_dfr(seq_len(round(dat_i$PercentLive_pct[1], digits=0)),</pre>
                           ~dat_i[, LiveObs := 1])
 dat_nl <- purrr::map_dfr(seq_len((100-round(dat_i$PercentLive_pct[1]),</pre>
                                                digits=0))),
                            ~dat_i[, LiveObs := 0])
  dat <- rbind(dat_1, dat_nl)</pre>
  jbji_p_binom <- rbind(jbji_p_binom, dat)</pre>
saveRDS(jbji_p_binom, paste0('output/model_results/data/jbji_p_binom_',
                              Sys.Date(), '.rds'))
jbji_pct_glmm <- brm(formula=LiveObs ~ RelYear+(1 | UniversalReefID),</pre>
                     data=jbji_p_binom, family=bernoulli, cores=ncores,
```

```
control= list(adapt_delta=0.995, max_treedepth=20),
                     iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
                     seed=4331, backend="rstan", threads=threading(threads),
                     file="output/model_results/GLMMs/jbji_pct_glmm3.rds")
# jbji_pct_glmm <- update(jbji_pct_glmm,</pre>
                          newdata = jbji_p_binom,
                          family=bernoulli, cores=ncores,
#
#
                          control= list(adapt delta=0.995, max treedepth=20),
#
                          iter=iter, warmup=warmup, chains=nchains, init=0, thin=3,
#
                          seed=4331, backend="rstan", threads=threading(threads),
#
                          file="output/model_results/GLMMs/jbji_pct_glmm3.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- jbji_p
models <- list(jbji_pct_glmm)</pre>
modresults(data, models, "Percent live", meplotzoom=FALSE)
## Lemon Bay Aquatic Preserve_Natural -----
lb_p <- subset(oysterraw_pct,</pre>
               oysterraw_pct$MA_plotlab=="Lemon Bay Aquatic Preserve_Natural")
lb_p[, PercentLive_dec := PercentLive_pct/100]
#PercentLiveMethod=="Percent" for Lemon Bay program(s) with sufficient data,
#so cannot be modeled as binomial
saveRDS(lb_p, paste0('output/model_results/data/lb_p_', Sys.Date(), '.rds'))
lb_pct_glmm <- brm(formula=PercentLive_dec ~</pre>
                     RelYear+(0+RelYear | ReefIdentifier),
                   data=subset(lb_p, lb_p$PercentLive_dec > 0 & lb_p$PercentLive_dec < 1),family=Beta,
                   cores=ncores, control= list(adapt_delta=0.995, max_treedepth=20),
                   iter=iter, warmup=warmup, chains=nchains, init=0, thin=3, seed=8465,
                   backend="rstan", threads=threading(threads),
                   file="output/model_results/GLMMs/lb_pct_glmm6.rds")
# lb_pct_qlmm <- update(lb_pct_qlmm,
#
                        newdata = subset(lb_p, lb_p$PercentLive_dec > 0),
#
                        family=Beta,
#
                         cores=ncores, control= list(adapt_delta=0.995, max_treedepth=20),
#
                         iter=iter, warmup=warmup, chains=nchains, init=0, thin=3, seed=8465,
                        backend="rstan", threads=threading(threads),
#
                        file="output/model_results/GLMMs/lb_pct_glmm6.rds")
# Create model results tables and save diagnostic plots and marginal effects plots
data <- lb_p
models <- list(lb_pct_glmm)</pre>
modresults(data, models, "Percent live", meplotzoom=FALSE)
```

Save & Export Results

The compiled model results variable is saved to a csv and rds file. The model results are evaluated to see if any models need to be reconsidered based on the rhat convergence being over 1.05. rhat values are written to file.

```
fwrite(oysterresults, paste0("output/GLMM_AllDates_ModelResults.csv"), sep=",")
saveRDS(oysterresults, paste0("output/GLMM_AllDates_ModelResults.rds"))
#Get Rhat values for all models to check which ones may need to be reparameterized
model_list <- unique(oysterresults$filename)</pre>
rhats_all <- data.table(filename=character(),</pre>
                         term=character(),
                         rhat=numeric())
rhats_sum <- data.table(filename=character(),</pre>
                         rhat=numeric())
for(mod in model_list){
  mod_i <- readRDS(mod)</pre>
  allrhat_i <- rhat(mod_i)</pre>
  sumrhat_i <- c(summary(mod_i)$fixed$Rhat, summary(mod_i)$spec_pars$Rhat)</pre>
  allrhat_model_i <- data.table(filename=mod,</pre>
                                 term=names(allrhat_i),
                                 rhat=allrhat_i)
  sumrhat_model_i <- data.table(filename=mod,</pre>
                                 rhat=sumrhat_i)
 rhats_all <- rbind(rhats_all, allrhat_model_i)</pre>
 rhats_sum <- rbind(rhats_sum, sumrhat_model_i)</pre>
}
rhats_all[, rhat_r := round(rhat, 2)]
rhats_sum[, rhat_r := round(rhat, 2)]
saveRDS(rhats_all, paste0("output/rhats_all_", Sys.Date(), ".rds"))
saveRDS(rhats_sum, paste0("output/rhats_sum_", Sys.Date(), ".rds"))
models_to_check_allrhat <- unique(rhats_all[rhat_r > 1.05, filename])
models_to_check_sumrhat <- unique(rhats_sum[rhat_r > 1.05, filename])
# Zip all figures
for(p in c("Density", "Percent_Live", "Shell_Height")){
  out_dir <- paste0("output/", p)</pre>
  fig_list <- list.files(paste0(out_dir, "/Figures"), pattern=".png", full=FALSE)
 filename <- paste0("Oyster", gsub("_", "", p), "Figures")
  setwd(paste0(out_dir, "/Figures"))
  zip(filename, files=fig_list)
  setwd(wd)
}
```

Oyster Figures

Density

Apalachicola Bay Aquatic Preserve

Natural

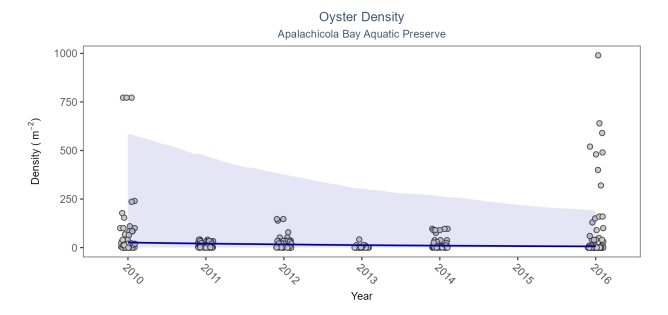


Figure 1: Figure for Oyster Density in Apalachicola Bay Aquatic Preserve

Table 1: Model results for Oyster Density - Natural

Shell Type	$Habitat\ Type$	Trend Status	Estimate	Standard Error	$Credible\ Interval$
Live Oyster Shells	Natural	Significantly decreasing trend	-0.24	0.11	-0.47 to -0.02

Apalachicola National Estuarine Research Reserve Natural

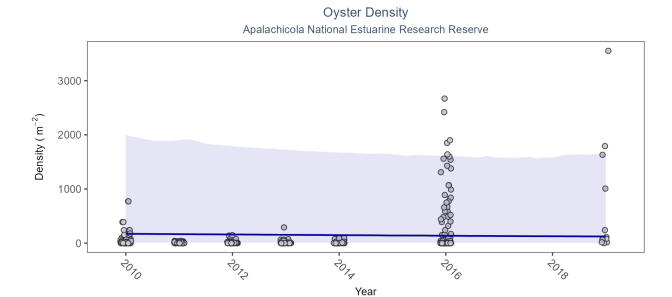


Figure 2: Figure for Oyster Density in Apalachicola National Estuarine Research Reserve

Table 2: Model results for Oyster Density - Natural

Shell Type	$Habitat\ Type$	Trend Status	Estimate	Standard Error	Credible Interval
Live Oyster Shells	Natural	No significant change	-0.04	0.1	-0.23 to 0.16

Estero Bay Aquatic Preserve

Natural

Oyster Density Estero Bay Aquatic Preserve

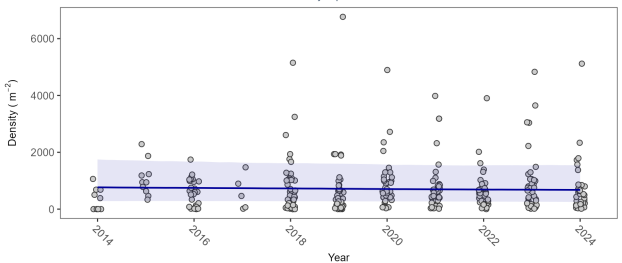


Figure 3: Figure for Oyster Density in Estero Bay Aquatic Preserve

Table 3: Model results for Oyster Density - Natural

Shell Type	$Habitat\ Type$	Trend Status	Estimate	$Standard\ Error$	$Credible\ Interval$
Live Oyster Shells	Natural	No significant change	-0.01	0.02	-0.06 to 0.03

Guana River Marsh Aquatic Preserve

Natural

Oyster Density Guana River Marsh Aquatic Preserve

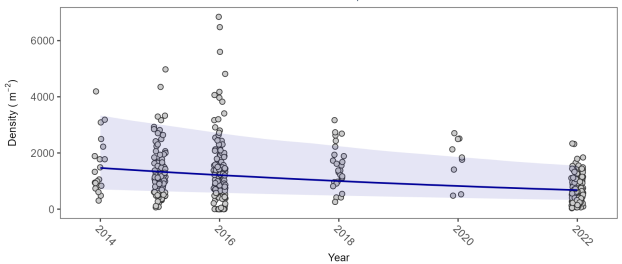


Figure 4: Figure for Oyster Density in Guana River Marsh Aquatic Preserve

Table 4: Model results for Oyster Density - Natural

$Shell\ Type$	$Habitat\ Type$	Trend Status	Estimate	$Standard\ Error$	$Credible\ Interval$
Live Oyster Shells	Natural	Significantly decreasing trend	-0.1	0.01	-0.12 to -0.07

Guana Tolomato Matanzas National Estuarine Research Reserve Natural



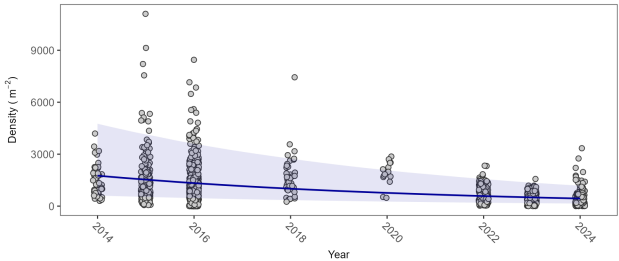


Figure 5: Figure for Oyster Density in Guana Tolomato Matanzas National Estuarine Research Reserve

Table 5: Model results for Oyster Density - Natural

Shell Type	Habitat Type	Trend Status	Estimate	Standard Error	Credible Interval
Live Oyster Shells	Natural	Significantly decreasing trend	-0.14	0.01	-0.15 to -0.12

Indian River-Vero Beach to Ft. Pierce Aquatic Preserve Natural



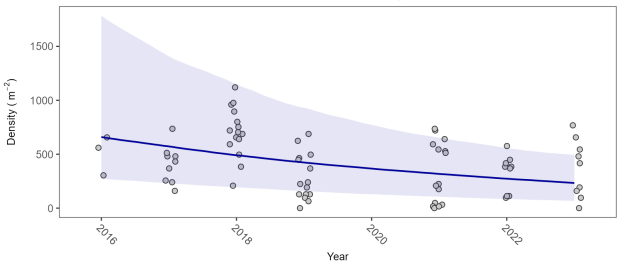


Figure 6: Figure for Oyster Density in Indian River-Vero Beach to Ft. Pierce Aquatic Preserve

Table 6: Model results for Oyster Density - Natural

Shell Type	Habitat Type	Trend Status	Estimate	Standard Error	Credible Interval
Live Oyster Shells	Natural	Significantly decreasing trend	-0.17	0.06	-0.29 to -0.05

Lemon Bay Aquatic Preserve

Natural

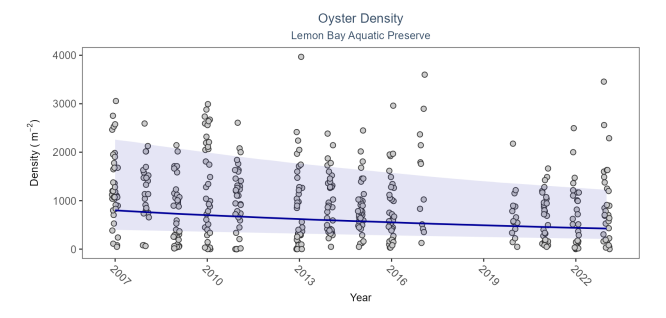


Figure 7: Figure for Oyster Density in Lemon Bay Aquatic Preserve

Table 7: Model results for Oyster Density - Natural

$Shell\ Type$	$Habitat\ Type$	Trend Status	Estimate	Standard Error	$Credible\ Interval$
Live Oyster Shells	Natural	Significantly decreasing trend	-0.04	0.01	-0.05 to -0.03

Pine Island Sound Aquatic Preserve

Natural

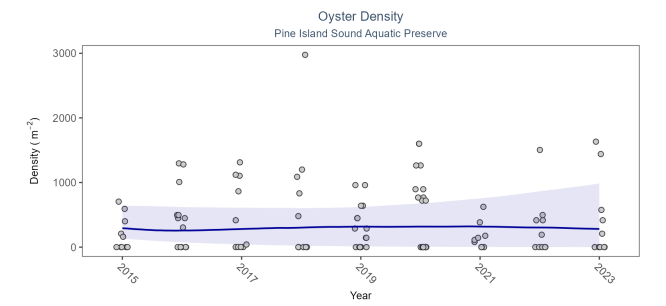


Figure 8: Figure for Oyster Density in Pine Island Sound Aquatic Preserve

Table 8: Model results for Oyster Density - Natural

Shell Type	$Habitat\ Type$	Trend Status	Estimate	Standard Error	Credible Interval
Live Oyster Shells	Natural	No significant change	-0.27	0.47	-1.22 to 0.72

Restored

Oyster Density Pine Island Sound Aquatic Preserve 2000 -1500 0 0 Density (m^{-2}) 0 8 1000 500 -0 660 000 ф Ф 0 08 0 0 - ∞ 707> 7075 2070 Year

Figure 9: Figure for Oyster Density in Pine Island Sound Aquatic Preserve

Table 9: Model results for Oyster Density - Restored

Shell Type	Habitat Type	Trend Status	Estimate	Standard Error	Credible Interval
Live Oyster Shells	Restored	Significantly increasing trend	0.02	0.02	0 to 0.08

Percent Live

Apalachicola Bay Aquatic Preserve

Natural

Oyster Percent Live Cover Apalachicola Bay Aquatic Preserve

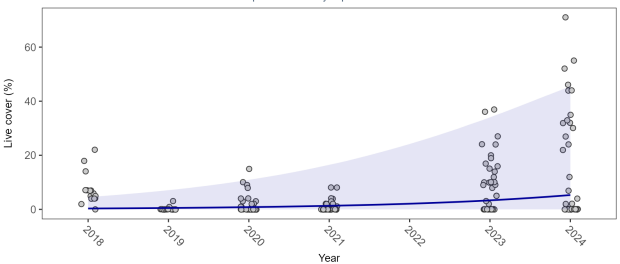


Figure 10: Figure for Oyster Percent Live in Apalachicola Bay Aquatic Preserve

Table 10: Model results for Oyster Percent Live - Natural

Shell Type	$Habitat\ Type$	Trend Status	Estimate	Standard Error	Credible Interval
Live Oyster Shells	Natural	Significantly increasing trend	0.48	0.02	0.44 to 0.53

Apalachicola National Estuarine Research Reserve Natural

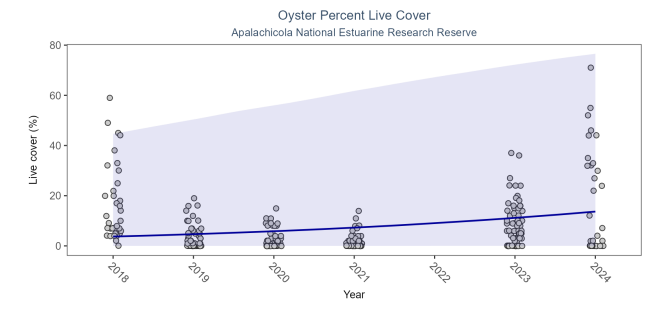


Figure 11: Figure for Oyster Percent Live in Apalachicola National Estuarine Research Reserve

Table 11: Model results for Oyster Percent Live - Natural

Shell Type	Habitat Type	Trend Status	Estimate	Standard Error	Credible Interval
Live Oyster Shells	Natural	Significantly increasing trend	0.23	0.01	0.21 to 0.26

Guana River Marsh Aquatic Preserve

Natural



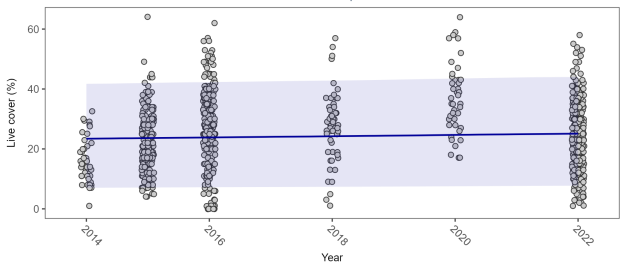


Figure 12: Figure for Oyster Percent Live in Guana River Marsh Aquatic Preserve

Table 12: Model results for Oyster Percent Live - Natural

Shell Type	Habitat Type	Trend Status	Estimate	Standard Error	Credible Interval
Live Oyster Shells	Natural	Significantly increasing trend	0.01	0	0 to 0.02

Guana Tolomato Matanzas National Estuarine Research Reserve Natural

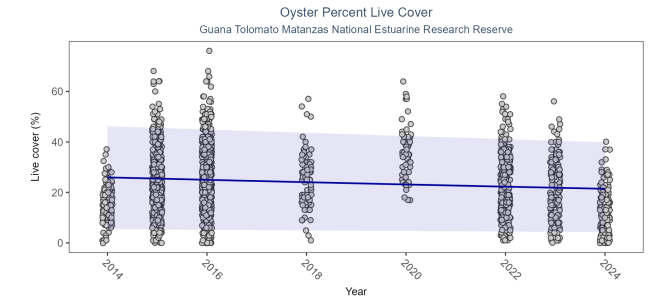


Figure 13: Figure for Oyster Percent Live in Guana Tolomato Matanzas National Estuarine Research Reserve

Table 13: Model results for Oyster Percent Live - Natural

Shell Type	Habitat Type	Trend Status	Estimate	Standard Error	Credible Interval
Live Oyster Shells	Natural	Significantly decreasing trend	-0.02	0	-0.03 to -0.02

Indian River-Vero Beach to Ft. Pierce Aquatic Preserve Natural



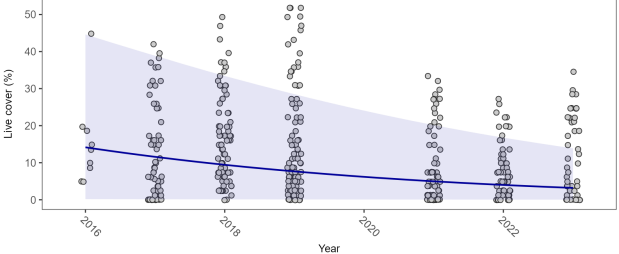


Figure 14: Figure for Oyster Percent Live in Indian River-Vero Beach to Ft. Pierce Aquatic Preserve

Table 14: Model results for Oyster Percent Live - Natural

Shell Type	Habitat Type	Trend Status	Estimate	Standard Error	Credible Interval
Live Oyster Shells	Natural	Significantly decreasing trend	-0.23	0.01	-0.25 to -0.21

Jensen Beach to Jupiter Inlet Aquatic Preserve Natural

Oyster Percent Live Cover Jensen Beach to Jupiter Inlet Aquatic Preserve

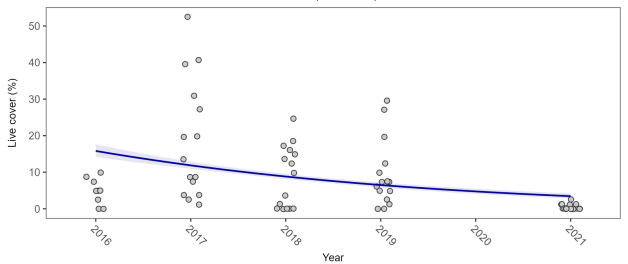


Figure 15: Figure for Oyster Percent Live in Jensen Beach to Jupiter Inlet Aquatic Preserve

Table 15: Model results for Oyster Percent Live - Natural

Shell Type	Habitat Type	Trend Status	Estimate	Standard Error	Credible Interval
Live Oyster Shells	Natural	Significantly decreasing trend	-0.33	0.03	-0.39 to -0.28

Lemon Bay Aquatic Preserve

Natural

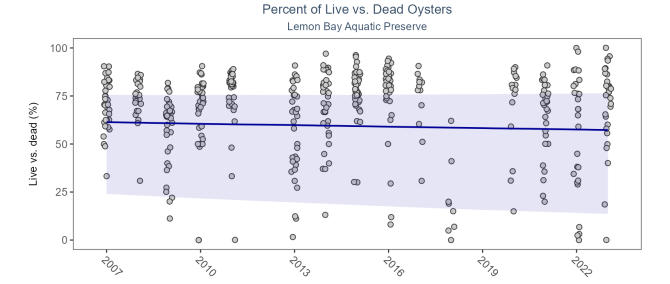


Figure 16: Figure for Oyster Percent Live in Lemon Bay Aquatic Preserve

Year

Table 16: Model results for Oyster Percent Live - Natural

Shell Type	$Habitat\ Type$	Trend Status	Estimate	$Standard\ Error$	$Credible\ Interval$
Live Oyster Shells	Natural	No significant change	-0.01	0.01	-0.03 to 0.01

Shell Height

Apalachicola National Estuarine Research Reserve

Natural



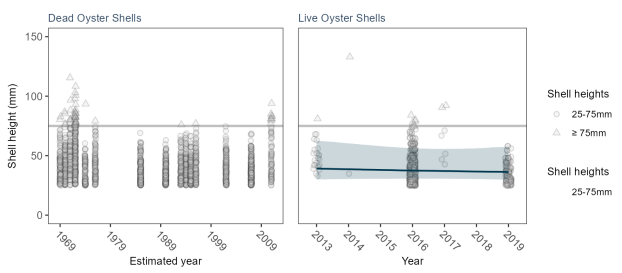


Figure 17: Figure for Oyster Shell Height in Apalachicola National Estuarine Research Reserve

Table 17: Model results for Oyster Shell Height - Natural

Shell Type	$Habitat\ Type$	Trend Status	Estimate	Standard Error	Credible Interval
Dead Oyster Shells	Natural	-	-	-	NA to NA
Dead Oyster Shells	Natural	-	-	-	NA to NA
Dead Oyster Shells	Natural	-	-	-	NA to NA
Live Oyster Shells	Natural	-	-	-	NA to NA
Live Oyster Shells	Natural	No significant change	-1.21	4.52	-10.4 to 7.72
Live Oyster Shells	Natural	-	-	-	NA to NA

Estero Bay Aquatic Preserve

Natural

Oyster Size Class Estero Bay Aquatic Preserve

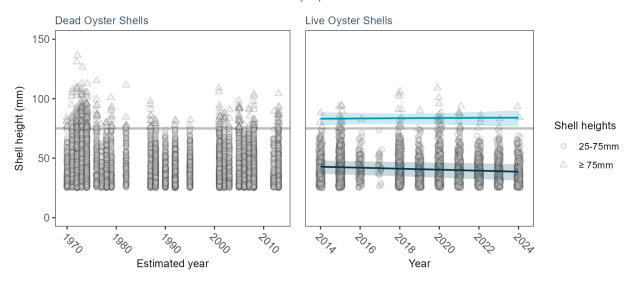


Figure 18: Figure for Oyster Shell Height in Estero Bay Aquatic Preserve

Table 18: Model results for Oyster Shell Height - Natural

Shell Type	Habitat Type	Trend Status	Estimate	Standard Error	Credible Interval
Dead Oyster Shells	Natural	-	-	-	NA to NA
Dead Oyster Shells	Natural	-	-	-	NA to NA
Dead Oyster Shells	Natural	-	-	-	NA to NA
Live Oyster Shells	Natural	No significant change	0.07	0.42	-0.75 to 0.89
Live Oyster Shells	Natural	Significantly decreasing trend	-0.43	0.10	-0.64 to -0.22
Live Oyster Shells	Natural	-	-	-	NA to NA

Guana River Marsh Aquatic Preserve

Natural



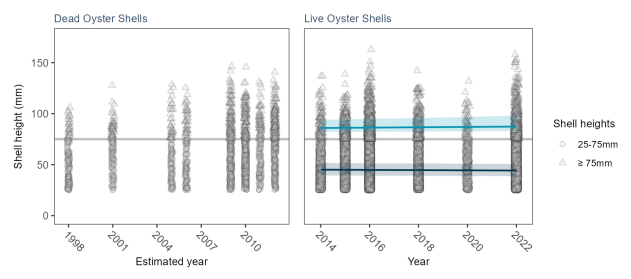


Figure 19: Figure for Oyster Shell Height in Guana River Marsh Aquatic Preserve

Table 19: Model results for Oyster Shell Height - Natural

Shell Type	Habitat Type	Trend Status	Estimate	Standard Error	Credible Interval
Dead Oyster Shells	Natural	-	-	-	NA to NA
Dead Oyster Shells	Natural	-	-	-	NA to NA
Dead Oyster Shells	Natural	-	-	-	NA to NA
Live Oyster Shells	Natural	Significantly increasing trend	1.16	0.54	0.13 to 2.25
Live Oyster Shells	Natural	No significant change	-0.91	0.56	-2.04 to 0.14
Live Oyster Shells	Natural	-	-	-	NA to NA

Guana Tolomato Matanzas National Estuarine Research Reserve Natural

Oyster Size Class Guana Tolomato Matanzas National Estuarine Research Reserve

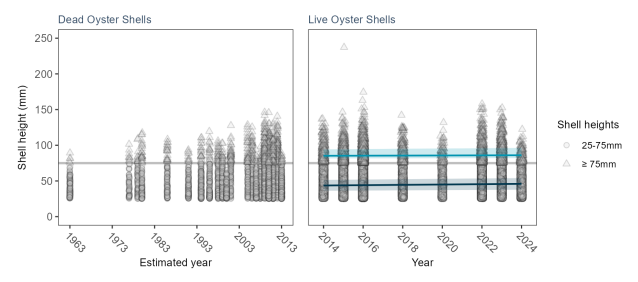


Figure 20: Figure for Oyster Shell Height in Guana Tolomato Matanzas National Estuarine Research Reserve

Table 20: Model results for Oyster Shell Height - Natural

Shell Type	Habitat Type	Trend Status	Estimate	Standard Error	Credible Interval
Dead Oyster Shells	Natural	-	-	-	NA to NA
Dead Oyster Shells	Natural	-	-	-	NA to NA
Dead Oyster Shells	Natural	-	-	-	NA to NA
Live Oyster Shells	Natural	No significant change	0.57	0.37	-0.14 to 1.26
Live Oyster Shells	Natural	Significantly increasing trend	1.55	0.25	1.07 to 2.07
Live Oyster Shells	Natural	-	-	-	NA to NA

Indian River-Vero Beach to Ft. Pierce Aquatic Preserve Natural

Oyster Size Class Indian River-Vero Beach to Ft. Pierce Aquatic Preserve

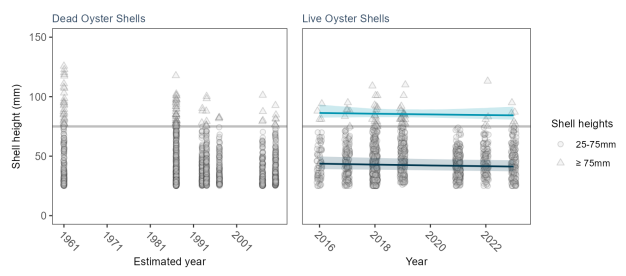


Figure 21: Figure for Oyster Shell Height in Indian River-Vero Beach to Ft. Pierce Aquatic Preserve

Table 21: Model results for Oyster Shell Height - Natural

Shell Type	Habitat Type	Trend Status	Estimate	Standard Error	Credible Interval
Dead Oyster Shells	Natural	-	-	-	NA to NA
Dead Oyster Shells	Natural	-	-	-	NA to NA
Dead Oyster Shells	Natural	-	-	-	NA to NA
Live Oyster Shells	Natural	No significant change	-1.21	2.98	-7.4 to 3.64
Live Oyster Shells	Natural	Significantly decreasing trend	-0.95	0.46	-1.9 to -0.07
Live Oyster Shells	Natural	-	-	-	NA to NA