

BITS: Baltic International Trawl Survey data processing summary

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General info

This document presents the cleaning code and summary of the Baltic Sea bottom trawl survey provided by DATRAS, ICES. It contains data from **1992** and up to **2020**.

Data cleaning in R

```
#####
#### R code to download and clean DATRAS data from ICES
#### URL: https://datras.ices.dk/
#### Coding: Aurore Maureaud + Juliano Palacios + Daniel van Denderen, December 2022
#####
rm(list=ls())

date <- "14DEC2022"

#####
#### LOAD LIBRARIES & options to decide in the code
#####
library(data.table)
```

```

library(tidyverse)
library(icesDatras)
library(wormrms)
library(curl)
library(urltools)
library(here) # for easy work around on multiple computers
library(taxize) # for getting correct species names
library(googledrive)

# load relevant functions
source("functions/write_clean_data.r")
source("functions/clean_taxa.R")
source("functions/get_length_weight_coeffs.R")
fishglob_data_columns <- read_excel("standard_formats/fishglob_data_columns.xlsx")

# should the last version of DATRAS be downloaded?
download_last_version <- FALSE

# should the data be loaded from the last version saved?
load_stored_datras <- FALSE

# should we save a new version of hh and hl?
save_hh_and_hl <- FALSE

# should we get the length-weight relationships from fishbase?
need_get_lw_rel <- FALSE

#####
##### LOAD FILES #####
#####

if(download_last_version == TRUE){
  last.year <- 2020

  # Haul info from Datras
  hh.ns <- getDATRAS(record='HH', survey='NS-IBTS', years=c(1967:last.year),
    quarters=c(1,3))
  hh.baltic <- getDATRAS(record='HH', survey='BITS', years=c(1991:last.year),
    quarters=c(1,4))
  hh.evhoe <- getDATRAS(record='HH', survey='EVHOE', years=c(1997:last.year),
    quarters=4)
  hh.cgfs <- getDATRAS(record='HH', survey='FR-CGFS', years=c(1998:last.year),
    quarters=4)
  hh.igfs <- getDATRAS(record='HH', survey='IE-IGFS', years=c(2003:last.year),
    quarters=4)
  hh.nigfs <- getDATRAS(record='HH', survey='NIGFS', years=c(2005:last.year),
    quarters=c(1:4))
  hh.pt <- getDATRAS(record='HH', survey='PT-IBTS', years=c(2002:last.year),
    quarters=c(3:4))
  hh.rock <- getDATRAS(record='HH', survey='ROCKALL', years=c(1999:2009),
    quarters=3)
  hh.scorock <- getDATRAS(record='HH', survey='SCOROC', years=c(2011:last.year),
    quarters=c(1:3))
}

```

```

            quarters=3)
hh.swc <- getDATRAS(record='HH', survey='SWC-IBTS', years=c(1985:2010),
                     quarters=c(1:4))
hh.scowcgfs <- getDATRAS(record='HH', survey='SCOWCGFS', years=c(2011:last.year),
                           quarters=c(1:4))

write.csv(hh.ns, file = "E:/fishglob data/Publicly available/DATRAS/hh.ns.csv",
          row.names = F)
write.csv(hh.baltic, file = "E:/fishglob data/Publicly available/DATRAS/hh.baltic.csv",
          row.names = F)
write.csv(hh.evhoe, file = "E:/fishglob data/Publicly available/DATRAS/hh.evhoe.csv",
          row.names = F)
write.csv(hh.cgfs, file = "E:/fishglob data/Publicly available/DATRAS/hh.cgfs.csv",
          row.names = F)
write.csv(hh.igfs, file = "E:/fishglob data/Publicly available/DATRAS/hh.igfs.csv",
          row.names = F)
write.csv(hh.nigfs, file = "E:/fishglob data/Publicly available/DATRAS/hh.nigfs.csv",
          row.names = F)
write.csv(hh.pt, file = "E:/fishglob data/Publicly available/DATRAS/hh.pt.csv",
          row.names = F)
write.csv(hh.rock, file = "E:/fishglob data/Publicly available/DATRAS/hh.rock.csv",
          row.names = F)
write.csv(hh.scorock, file = "E:/fishglob data/Publicly available/DATRAS/hh.scorock.csv",
          row.names = F)
write.csv(hh.swc, file = "E:/fishglob data/Publicly available/DATRAS/hh.swc.csv",
          row.names = F)
write.csv(hh.scowcgfs, file = "E:/fishglob data/Publicly available/DATRAS/
hh.scowcgfs.csv", row.names = F)

# Length info from DATRAS
hl.ns <- getDATRAS(record='HL', survey='NS-IBTS', years=c(1967:last.year),
                     quarters=c(1,3))
hl.baltic <- getDATRAS(record='HL', survey='BITS', years=c(1991:last.year),
                        quarters=c(1,4))
hl.evhoe <- getDATRAS(record='HL', survey='EVHOE', years=c(1997:last.year),
                       quarters=4)
hl.cgfs <- getDATRAS(record='HL', survey='FR-CGFS', years=c(1998:last.year),
                      quarters=4)
hl.igfs <- getDATRAS(record='HL', survey='IE-IGFS', years=c(2003:last.year),
                      quarters=4)
hl.nigfs <- getDATRAS(record='HL', survey='NIGFS', years=c(2005:last.year),
                       quarters=c(1:4))
hl.pt <- getDATRAS(record='HL', survey='PT-IBTS', years=c(2002:last.year),
                    quarters=c(3:4))
hl.rock <- getDATRAS(record='HL', survey='ROCKALL', years=c(1999:2009),
                      quarters=3)
hl.scorock <- getDATRAS(record='HL', survey='SCOROC', years=c(2011:last.year),
                         quarters=3)
hl.swc <- getDATRAS(record='HL', survey='SWC-IBTS', years=c(1985:2010),
                     quarters=c(1:4))
hl.scowcgfs <- getDATRAS(record='HL', survey='SCOWCGFS', years=c(2011:last.year),
                           quarters=c(1:4))

```

```

write.csv(hl.ns, file = "E:/fishglob data/Publicly available/DATRAS/hl.ns.csv",
          row.names = F)
write.csv(hl.baltic, file = "E:/fishglob data/Publicly available/DATRAS/hl.baltic.csv",
          row.names = F)
write.csv(hl.evhoe, file = "E:/fishglob data/Publicly available/DATRAS/hl.evhoe.csv",
          row.names = F)
write.csv(hl.cgfs, file = "E:/fishglob data/Publicly available/DATRAS/hl.cgfs.csv",
          row.names = F)
write.csv(hl.igfs, file = "E:/fishglob data/Publicly available/DATRAS/hl.igfs.csv",
          row.names = F)
write.csv(hl.nigfs, file = "E:/fishglob data/Publicly available/DATRAS/hl.nigfs.csv",
          row.names = F)
#write.csv(hl.pt, file = "E:/fishglob data/Publicly available/DATRAS/hl.pt.csv",
#row.names = F)
write.csv(hl.rock, file = "E:/fishglob data/Publicly available/DATRAS/hl.rock.csv",
          row.names = F)
write.csv(hl.scorock, file = "E:/fishglob data/Publicly available/DATRAS/hl.scorock.csv",
          row.names = F)
#write.csv(hl.swc, file = "E:/fishglob data/Publicly available/DATRAS/hl.swc.csv",
#row.names = F)
write.csv(hl.scowcgfs, file = "E:/fishglob data/Publicly available/DATRAS/hl.scowcgfs.csv",
          row.names = F)

}

if(load_stored_datras == TRUE){

  hh.ns <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.ns.csv")
  hh.baltic <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.baltic.csv")
  hh.evhoe <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.evhoe.csv")
  hh.cgfs <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.cgfs.csv")
  hh.igfs <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.igfs.csv")
  hh.nigfs <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.nigfs.csv")
  hh.pt <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.pt.csv")
  hh.rock <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.rock.csv")
  hh.scorock <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.scorock.csv")
  hh.swc <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.swc.csv")
  hh.scowcgfs <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.scowcgfs.csv")

  hl.ns <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.ns.csv")
  hl.baltic <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.baltic.csv")
  hl.evhoe <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.evhoe.csv")
  hl.cgfs <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.cgfs.csv")
  hl.igfs <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.igfs.csv")
  hl.nigfs <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.nigfs.csv")
  hl.pt <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.pt.csv") %>%
    dplyr::rename(Valid_Aphia = ValidAphiaID) %>%
    select(RecordType, Survey, Quarter, Country, Ship, Gear, SweepLngt, GearEx,
           DoorType, StNo, HaulNo, Year, SpecCodeType, SpecCode, SpecVal, Sex,
           TotalNo, CatIdentifier, NoMeas, SubFactor, SubWgt, CatCatchWgt, LngtCode,
           LngtClass, HLNoAtLngt, DevStage, LenMeasType, DateofCalculation,
           Valid_Aphia)
  hl.rock <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.rock.csv")
}

```

```

hl.scorock <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.scorock.csv")
hl.swc <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.swc.csv")%>%
  dplyr::rename(Valid_Aphia = ValidAphiaID) %>%
  select(RecordType, Survey, Quarter, Country, Ship, Gear, SweepLngt, GearEx,
         DoorType, StNo, HaulNo, Year, SpecCodeType, SpecCode, SpecVal, Sex,
         TotalNo, CatIdentifier, NoMeas, SubFactor, SubWgt, CatCatchWgt, LngtCode,
         LngtClass, HLNoAtLngt, DevStage, LenMeasType, DateofCalculation,
         Valid_Aphia)
hl.scowcgfs <- read.csv("E:/fishglob data/Publicly available/DATRAS/hl.scowcgfs.csv")

hh <- rbind(hh.ns, hh.baltic, hh.evhoe, hh.cgfs, hh.igfs, hh.nigfs, hh.pt, hh.rock,
            hh.scorock, hh.swc, hh.scowcgfs)

hl <- rbind(hl.ns, hl.baltic, hl.evhoe, hl.cgfs, hl.igfs, hl.nigfs, hl.pt, hl.rock,
            hl.scorock, hl.swc, hl.scowcgfs)

rm(hl.ns, hl.baltic, hl.evhoe, hl.cgfs, hl.igfs, hl.nigfs, hl.pt, hl.rock, hl.scorock,
    hl.swc, hl.scowcgfs,
    hh.ns, hh.baltic, hh.evhoe, hh.cgfs, hh.igfs, hh.nigfs, hh.pt, hh.rock, hh.scorock,
    hh.swc, hh.scowcgfs)
#rm(ca.ns, ca.baltic, ca.evhoe, ca.cgfs, ca.igfs, ca.nigfs, ca.pt, ca.rock,
#  ca.scorock, ca.swc, ca.scowcgfs)

}

if(save_hh_and_hl == TRUE){
  save(hh, file = paste0("E:/fishglob data/Publicly available/DATRAS/hh.",date,".RData"))
  save(hl, file = paste0("E:/fishglob data/Publicly available/DATRAS/hl.",date,".RData"))
}

#####
# Alternative
#####

# Juliano
hl <- fread("/Volumes/Enterprise/Data/FishGlob/Data/Raw/ices_hl.csv")
unique(hl$Survey)
hh <- fread("/Volumes/Enterprise/Data/FishGlob/Data/Raw/ices_hh.csv")
unique(hh$Survey)

# Aurore
load("/Volumes/Elements/fishglob data/Publicly available/DATRAS/hl.29JUL2021.RData")
load("/Volumes/Elements/fishglob data/Publicly available/DATRAS/hh.29JUL2021.RData")

#####
##### CREATE A UNIQUE HAUL ID #####
#####

hl$HaulID <- paste(hl$Survey, hl$Year, hl$Quarter, hl$Country, hl$Ship, hl$Gear, hl$StNo,
                     hl$HaulNo)
hh$HaulID <- paste(hh$Survey, hh$Year, hh$Quarter, hh$Country, hh$Ship, hh$Gear, hh$StNo,
                     hh$HaulNo)

```

```

# Is the HaulID unique?
hhn <- unique(hh$HaulID)
length(hhn)==nrow(hh)

# check which one is not
pb <- c()
for (i in 1:length(hhn)){
  j <- which(hh$HaulID==hhn[i])
  if(length(j)>1){pb <- hhn[i]}
}

# problem with one haul in NS-IBTS
hh <- hh %>%
  filter(HaulID!=pb)
hl <- hl %>%
  filter(HaulID!=pb)

# Only keep hauls where there is the length composition.
# 70273 hauls in hh and 70273 in hl
hh <- subset(hh, hh$HaulID %in% hl$HaulID)
hl <- subset(hl, hl$HaulID %in% hh$HaulID)

#####
##### MERGE HH and HL FILES
#####

haulidhl <- sort(unique(hl$HaulID))
haulidhh <- sort(unique(hh$HaulID))
identical(haulidhh, haulidhl)
rm(haulidhh, haulidhl)

# remove some columns in hl
hl$SweepLngt <- hl$SpecCodeType <- hl$SpecCode <- hl$Sex <- hl>DateofCalculation <- NULL
hl$RecordType <- hl$GearEx <- NULL

# remove some columns in hh
hh>DateofCalculation <- hh$ThClineDepth <- hh$ThermoCline <- hh$SwellHeight <- NULL
hh$SwellDir <- hh$WindSpeed <- hh$WindDir <- hh$BotCurSpeed <- NULL
hh$BotCurDir <- hh$SurCurSpeed <- hh$SurCurDir <- hh$SpeedWater <- hh$TowDir <- NULL
hh$WgtGroundRope <- hh$KiteDim <- hh$Buoyancy <- hh$Tickler <- NULL
hh$DoorWgt <- hh$DoorSurface <- hh$WarpDen <- hh$Warpdia <- hh$Warplngt <- NULL
hh$Rigging <- hh$HydroStNo <- hh$HaulLat <- hh$HaulLong <- hh$DayNight <- NULL
hh$Stratum <- hh$TimeShot <- hh$Day <- hh$RecordType <- hh$GearExp <- hh$DoorType <- NULL

#survey <- merge(hh, hl, by='HaulID', all.x=FALSE, all.y=TRUE)
survey <- right_join(hh, hl, by=c('HaulID', 'Survey', 'Quarter', 'Country', 'Ship',
                                    'Gear', 'StNo', 'HaulNo', 'Year'))
nrow(survey)==nrow(hl)

survey <- survey %>%
  dplyr::rename(SBT = BotTemp,
                SST = SurTemp,

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        Speed = GroundSpeed,
        AphiaID = Valid_Aphia)

#### Check if the HaulID is unique
#### Not the case for the baltic sea, a lot of duplicates!!!
#ids <- unique(hh$HaulID)
#pb <- vector()
#for(i in 1:length(ids)){
#  x <- which(hh$HaulID==ids[i])
#  if(length(x)>1){pb[length(pb)+1] <- ids[i]}
#}
#print(pb) # dim 0 ok!

#####
##### REMOVE INVALID DATA #####
#####
survey <- survey %>%
  filter(HaulVal %in% 'V', #Remove invalid hauls
         !is.na(AphiaID), # Remove invalid species records
         SpecVal %in% c(1,10,4,7),
         DataType %in% c('S','R','C'))

#####
##### RESCALE DATA INTO ABUNDANCE FOR THE HAUL DURATION AND ABUNDANCE AT LENGTH #####
#####
# If Data Type=='C', abundance at length already readjusted with time so get back the
# abundance for the actual duration of the haul.
# If data type=='R', abundance at length is multiplied by subfactor and adjusted to time
survey$CatCatchWgt = as.numeric(survey$CatCatchWgt)

survey <- survey %>%
  mutate(HLNoAtLngt = case_when(DataType=='C' ~ HLNoAtLngt*SubFactor*HaulDur/60,
                                 DataType %in% c('S','R') ~ HLNoAtLngt*SubFactor),
         TotalNo = case_when(DataType=='C' ~ TotalNo*HaulDur/60,
                               DataType %in% c('S','R') ~ TotalNo),
         CatCatchWgt = case_when(DataType=='C' ~ CatCatchWgt*HaulDur/60,
                                 DataType %in% c('S','R') ~ CatCatchWgt)) %>%
  select(-HaulVal, -DataType, -StdSpecRecCode, -SpecVal, -SubWgt, -SubFactor) %>%
  mutate(Survey = if_else(Survey=='SCOWCGFS', 'SWC-IBTS', Survey)) %>%
  mutate(Survey = if_else(Survey=='SCOROC', 'ROCKALL', Survey)) %>%
  filter(!(Survey=="NS-IBTS" & BySpecRecCode %in% c(0,2,3,4,5)),
         # remove hauls where not all species are recorded
         !(Survey=="BITS" & BySpecRecCode==0))

#####
##### GET THE SWEPT AREA in km2 #####
#####

source('cleaning.codes/source_DATRAS_wing_doorspread.R')

```

```

#####
##### GET CPUEs AND RIGHT COLUMNS NAMES
#####

# Remove data without length composition or negative values
xx <- subset(survey, HLNoAtLngt<0 | is.na(LngtClass))
no_length_hauls <- sort(unique(xx$HaulID)) # 10,178 hauls without length data

# Only keep abundances/weight
survey <- survey %>%
  #filter(!(HaulID %in% no_length_hauls)) %>% # remove hauls without length data
  mutate(numcpue = TotalNo/Area.swept, # abundance/km2
         wtcpue = CatCatchWgt/(Area.swept*1000), # weight in kg/km2
         numh = (TotalNo*60)/HaulDur, # abundance/hour
         wgth = CatCatchWgt*60/(HaulDur*1000), # weight in kg/h
         num = TotalNo, # raw number of individuals
         wgt = CatCatchWgt/1000, # raw weight in kg
         numlencpue = HLNoAtLngt/Area.swept, # abundance/km2 per length class
         numlenh = HLNoAtLngt*60/HaulDur, # abundance/h per length class
         Season = 'NA',
         Depth = replace(Depth, Depth<0, NA),
         SBT = replace(SBT, SBT<0, NA),
         SST = replace(SST, SST<0, NA),
         LngtClass = ifelse(LngtCode %in% c('.','0'), LngtClass*0.1, LngtClass)) %>%
  # fix unit of length class
  dplyr::rename(Length = LngtClass) %>%
  select(Survey, HaulID, StatRec, Year, Month, Quarter, Season, ShootLat, ShootLong,
         HaulDur, Area.swept, Gear, Depth, SBT, SST, AphiaID, CatIdentifier, numcpue,
         wtcpue, numh, wgth, num, wgt, Length, numlencpue, numlenh)
survey <- data.frame(survey)

#####
## fishglob taxa cleaning ##
#####

# Make AphiaID list per survey
aphia_datras <- survey %>%
  select(Survey, AphiaID) %>%
  dplyr::rename(survey = Survey,
                worms_id_datras = AphiaID) %>%
  distinct()

# Clean taxa north sea
ns_data <- aphia_datras %>% filter(survey=="NS-IBTS")
clean_ns <- clean_taxa(ns_data$worms_id_datras, input_survey = "NS-IBTS",
                       save=F, fishbase=TRUE)

# Clean taxa bay of biscay
evhoe_data <- aphia_datras %>% filter(survey=="EVHOE")
clean_evhoe <- clean_taxa(evhoe_data$worms_id_datras, input_survey = "EVHOE",
                           save=F, fishbase=TRUE)

```

```

# Clean taxa english channel
cgfs_data <- aphia_datras %>% filter(survey=="FR-CGFS")
clean_cgfs <- clean_taxa(cgfs_data$worms_id_datras, input_survey = "FR-CGFS",
                           save=F, fishbase=TRUE)

# Clean taxa baltic sea
bits_data <- aphia_datras %>% filter(survey=="BITS")
clean_bits <- clean_taxa(bits_data$worms_id_datras, input_survey = "BITS",
                           save=F, fishbase=TRUE)

# Clean taxa scottish sea
swc_data <- aphia_datras %>% filter(survey %in% c("SCOWCGFS", "SWC-IBTS"))
clean_swc <- clean_taxa(swc_data$worms_id_datras, input_survey = "SWC-IBTS",
                           save=F, fishbase=TRUE)

# Clean taxa rockall
rock_data <- aphia_datras %>% filter(survey %in% c("SCOROC", "ROCKALL"))
clean_rock <- clean_taxa(rock_data$worms_id_datras, input_survey = "ROCKALL",
                           save=F, fishbase=TRUE)

# Clean taxa irish sea
ir_data <- aphia_datras %>% filter(survey=="IE-IGFS")
clean_ir <- clean_taxa(ir_data$worms_id_datras, input_survey = "IE-IGFS",
                           save=F, fishbase=TRUE)

# Clean taxa northern ireland
nigfs_data <- aphia_datras %>% filter(survey=="NIGFS")
clean_nigfs <- clean_taxa(nigfs_data$worms_id_datras, input_survey = "NIGFS",
                           save=F, fishbase=TRUE)

# Clean taxa for portugal
pt_data <- aphia_datras %>% filter(survey=="PT-IBTS")
clean_pt <- clean_taxa(pt_data$worms_id_datras, input_survey = "PT-IBTS",
                           save=F, fishbase=TRUE)

clean_datras_taxa <- rbind(clean_bits, clean_cgfs, clean_evhoe, clean_ir, clean_nigfs,
                           clean_pt, clean_rock, clean_swc, clean_ns) %>%
  mutate(query = as.numeric(as.vector(query))) %>%
  distinct()

recoded_taxa <- c("Dipturus", "Liparis", "Chelon", "Mustelus", "Alosa", "Argentina",
                  "Callionymus", "Ciliata", "Gaidropsarus", "Sebastes", "Syngnathus",
                  "Pomatoschistus", "Gobius")

spp_to_recode <- c("Dipturus batis", "Dipturus flossada", "Dipturus batis-complex",
                  "Dipturus intermedia", "Liparis montagui", "Liparis liparis",
                  "Liparis liparis liparis", "Chelon aurata", "Chelon ramada",
                  "Mustelus mustelus/asterias", "Mustelus mustelus", "Mustelus asterias",
                  "Alosa alosa", "Alosa fallax", "Argentina silus", "Argentina sphyraena",
                  "Callionymus reticulatus", "Callionymus maculatus", "Ciliata mustela",
                  "Ciliata septentrionalis", "Gaidropsaurus macrophthalmus",
                  "Gaidropsaurus mediterraneus", "Gaidropsaurus vulgaris",
                  "Sebastes norvegicus", "Sebastes mentella", "Sebastes marinus",

```

```

    "Syngnathus rostellatus", "Syngnathus acus", "Syngnathus typhle",
    "Nerophis ophidion", "Pomatoschistus microps", "Pomatoschistus minutus",
    "Pomatoschistus pictus", "Gobius cobitis", "Gobius niger",
    "Leusueurigobius friesii", "Neogobius melanostomus")

alphaid <- get_wormsid(recoded_taxa)
alphaid <- tibble(taxa = recoded_taxa,
                  worms_id = alphaid[1:length(recoded_taxa)])
clean_manual_recoded <- clean_taxa(alphaid$worms_id, input_survey = "recoded",
                                      save = F, fishbase=TRUE)

clean_datras_taxa <- clean_datras_taxa %>%
  select(-survey) %>%
  mutate(SpecCode = ifelse(taxa %in% spp_to_recode, NA, SpecCode),
         rank = ifelse(taxa %in% spp_to_recode, "Genus", rank),
         #dipturus
         worms_id = ifelse(taxa %in% c("Dipturus batis", "Dipturus flossada",
                                         "Dipturus batis-complex", "Dipturus intermedia"),
                            105762, worms_id),
         taxa = ifelse(taxa %in% c("Dipturus batis", "Dipturus flossada",
                                    "Dipturus batis-complex", "Dipturus intermedia"),
                       "Dipturus", taxa),
         # liparis
         worms_id = ifelse(taxa %in% c("Liparis montagui", "Liparis liparis",
                                         "Liparis liparis liparis"), 126160, worms_id),
         taxa = ifelse(taxa %in% c("Liparis montagui", "Liparis liparis",
                                   "Liparis liparis liparis"), "Liparis", taxa),
         # chelon
         worms_id = ifelse(taxa %in% c("Chelon aurata", "Chelon ramada"), 126030, worms_id),
         taxa = ifelse(taxa %in% c("Chelon aurata", "Chelon ramada"), "Chelon", taxa),
         #mustelus
         worms_id = ifelse(taxa %in% c("Mustelus mustelus/asterias", "Mustelus mustelus",
                                         "Mustelus asterias"), 105732, worms_id),
         taxa = ifelse(taxa %in% c("Mustelus mustelus/asterias", "Mustelus mustelus",
                                   "Mustelus asterias"), "Mustelus", taxa),
         #alosa
         worms_id = ifelse(taxa %in% c("Alosa alosa", "Alosa fallax"), 125715, worms_id),
         taxa = ifelse(taxa %in% c("Alosa alosa", "Alosa fallax"), "Alosa", taxa),
         #argentina
         worms_id = ifelse(taxa %in% c("Argentina silus", "Argentina sphyraena"),
                           125885, worms_id),
         taxa = ifelse(taxa %in% c("Argentina silus", "Argentina sphyraena"),
                       "Argentina", taxa),
         # callionymus
         worms_id = ifelse(taxa %in% c("Callionymus reticulatus", "Callionymus maculatus"),
                           125930, worms_id),
         taxa = ifelse(taxa %in% c("Callionymus reticulatus", "Callionymus maculatus"),
                       "Callionymus", taxa),
         # ciliata
         worms_id = ifelse(taxa %in% c("Ciliata mustela", "Ciliata septentrionalis"),
                           125741, worms_id),
         taxa = ifelse(taxa %in% c("Ciliata mustela", "Ciliata septentrionalis"),
                       "Ciliata", taxa),

```

```

# gaidropsarus
worms_id = ifelse(taxa %in% c("Gaidropsaurus macrophthalmus",
                               "Gaidropsaurus mediterraneus",
                               "Gaidropsaurus vulgaris"),
                   125743,worms_id),
taxa = ifelse(taxa %in% c("Gaidropsaurus macrophthalmus",
                           "Gaidropsaurus mediterraneus",
                           "Gaidropsaurus vulgaris"), "Gaidropsarus",taxa),
# sebastes
worms_id = ifelse(taxa %in% c("Sebastes norvegicus","Sebastes mentella",
                               "Sebastes marinus"),
                   126175,worms_id),
taxa = ifelse(taxa %in% c("Sebastes norvegicus","Sebastes mentella",
                           "Sebastes marinus"),
               "Sebastes",taxa),
# syngnathus
worms_id = ifelse(taxa %in% c("Syngnathus rostellatus","Syngnathus acus",
                               "Syngnathus typhle","Nerophis ophidion"),
                   126227,worms_id),
taxa = ifelse(taxa %in% c("Syngnathus rostellatus","Syngnathus acus",
                           "Syngnathus typhle","Nerophis ophidion"),
               "Syngnathus",taxa),
# pomatosc
worms_id = ifelse(taxa %in% c("Pomatoschistus microps","Pomatoschistus minutus",
                               "Pomatoschistus pictus"), 125999,worms_id),
taxa = ifelse(taxa %in% c("Pomatoschistus microps","Pomatoschistus minutus",
                           "Pomatoschistus pictus"), "Pomatoschistus",taxa),
# gobius
worms_id = ifelse(taxa %in% c("Gobius cobitis","Gobius niger",
                               "Leusueurigobius friesii",
                               "Neogobius melanostomus"), 125988,worms_id),
taxa = ifelse(taxa %in% c("Gobius cobitis","Gobius niger",
                           "Leusueurigobius friesii",
                           "Neogobius melanostomus"), "Gobius",taxa),
) %>%
distinct()

# add taxonomy to data
survey <- left_join(survey, clean_datras_taxa, by=c("AphiaID" = "query")) %>%
filter(!is.na(AphiaID))

#####
##### RE-CALCULATE WEIGHTS
#####
detach(package:worms)
detach(package:plyr)

# select only certain gears
# 1. summary of gears per survey
gears <- data.frame(survey) %>%
group_by(Survey, Gear) %>%
summarise(hauls = length(unique(HaulID)), years = length(unique(Year))) %>%

```

```

select(Survey, Gear, hauls, years)

# 2. only select certain gears per survey (GOV and/or most dominant in cases without GOV)
survey <- survey %>%
  filter(!(Survey=="NS-IBTS" & Gear %in% c('ABD', 'BOT', 'DHT', 'FOT', 'GRT',
                                             'H18', 'HOB', 'HT', 'KAB', 'VIN'))),
  !(Survey=="BITS" & Gear %in% c('CAM', 'CHP', 'DT', 'EGY', 'ESB', 'EXP',
                                   'FOT', 'GRT', 'H20', 'HAK', 'LBT', 'SON')),
  !(Survey=="PT-IBTS" & Gear=='CAR'),
  !(Survey=="Can-Mar" & Gear=='Y36'))

# 3. associate an LME to each haul and make final list of species
### Prepare list for estimating length-weight parameters
list.taxa <- survey %>%
  mutate(species = str_split(taxa, pattern = " ", simplify = TRUE)[,2]) %>%
  select(HaulID, Survey, ShootLat, ShootLong, family, genus, species) %>%
  distinct()

# get LME
library(rgdal)
shape1 <- readOGR(dsn = "length.weight/LME66", layer="LMEs66")
coords <- list.taxa %>%
  dplyr::select(ShootLat, ShootLong, Survey) %>%
  distinct()
str(coords)

coordinates(coords) <- ~ ShootLong + ShootLat
proj4string(coords) <- proj4string(shape1)
lme <- over(coords, shape1)

coords <- list.taxa %>%
  dplyr::select(ShootLat, ShootLong, Survey) %>%
  distinct()
coords <- cbind(coords, lme$LME_NUMBER)
setnames(coords, old='lme$LME_NUMBER', new='lme')

coords$lme <- as.factor(coords$lme)
#Select from each LME 50 long and lat
ind <- c()
for (i in 1:nlevels(coords$lme)){
  ind <- c(ind, sample(which(coords$lme==levels(coords$lme)[i]), 50,
                        replace = FALSE))
}
long50 <- coords$ShootLong[ind]
lat50 <- coords$ShootLat[ind]
lme50 <- rep(levels(coords$lme), each=50)
#For each haul without LME find a close LME that has an LME number already
nlme <- subset(coords, is.na(lme)) # many hauls without LME 710
nlme$ShootLat <- as.numeric(as.vector(nlme$ShootLat))
nlme$ShootLong <- as.numeric(as.vector(nlme$ShootLong))
long50 <- as.numeric(as.vector(long50))
lat50 <- as.numeric(as.vector(lat50))

```

```

dilme <- c()
for (i in 1:length(lme50)){
  dilme <- cbind(dilme, (nlme$ShootLat-lat50[i])**2 + (nlme$ShootLong-long50[i])**2)
}
mindi <- apply(dilme, 1, which.min)
coords$lme[is.na(coords$lme)] <- lme50[mindi]
# assign the closest LME number to each haul without LME

#Check
coords$ShootLat <- as.numeric(as.vector(coords$ShootLat))
coords$ShootLong <- as.numeric(as.vector(coords$ShootLong))

# rockall not assigned to Faroe plateau but to celtic sea LME
coords$lme <- as.character(coords$lme)
coords <- coords %>%
  mutate(lme = replace(lme, Survey == 'ROCKALL', '60')) %>%
  as.data.frame()

survey <- left_join(survey, coords, by=c('ShootLat', 'ShootLong', 'Survey'))

if(need_get_lw_rel == TRUE){
  list.taxa <- survey %>%
    select(taxa, family, genus, taxa, lme, rank, Survey) %>%
    filter(!is.na(taxa),
           rank!="Suborder") %>%
    rename(survey = Survey) %>%
    distinct()

  write.csv(data.frame(list.taxa), file="length.weight/taxa.DATRAS.FB.tofill.csv",
            row.names=FALSE)
  save.image("length.weight/DATRAS_before_lw_4AUG2021.RData")

  ### run length-weight relationships open R 32bit
  list.taxa <- read.csv("length.weight/taxa.DATRAS.FB.tofill.csv")
  get_coeffs(list.taxa, survey="DATRAS", save=TRUE)
}

# 4. re-calculate weights with length-weight relationships
#load("length.weight/DATRAS_before_lw_4AUG2021.RData")
datalw <- read.csv('length.weight/length.weight_DATRAS.csv') %>%
  rename(Survey = survey) %>%
  select(-X) %>%
  mutate(lme = as.character(lme))

survey <- survey %>%
  filter(rank!="Suborder")

# summarize abundance/weight at the haul level
survey.num <- left_join(survey, datalw, by=c("taxa", "family", "genus", "lme",
                                              "rank", "Survey")) %>%
  select(Survey, HaulID, StatRec, Year, Month, Quarter, Season, ShootLat, ShootLong,

```

```

HaulDur,Area.swept,Gear,Depth,SBT,SST,family,genus,taxa,AphiaID,worms_id,
SpecCode,kingdom, class, order,phylum,rank,
CatIdentifier,numcpue,numh,num) %>%
distinct() %>%
group_by(Survey,HaulID,StatRec,Year,Month,Quarter,Season,ShootLat,ShootLong,
HaulDur,Area.swept,Gear,Depth,SBT,SST,family,genus,taxa,AphiaID,
worms_id, SpecCode,kingdom, class, order,phylum, rank) %>%
summarize_at(.vars=c('numcpue', 'numh', 'num'), .funs = function(x) sum(x)) %>%
ungroup()

survey.wgt <- left_join(survey, datalw, by=c("taxa","family","genus","lme",
                                              "rank","Survey")) %>%
select(Survey,HaulID,StatRec,Year,Month,Quarter,Season,ShootLat,ShootLong,HaulDur,
       Area.swept,Gear,Depth,SBT,SST,family,genus,taxa,AphiaID,worms_id,SpecCode,
       kingdom, class, order,phylum,rank,
       CatIdentifier,wtcpue,wgth,wgt) %>%
distinct() %>%
group_by(Survey,HaulID,StatRec,Year,Month,Quarter,Season,ShootLat,ShootLong,
HaulDur,Area.swept,Gear,Depth,SBT,SST,family,genus,taxa,AphiaID,worms_id,
SpecCode,kingdom, class, order, phylum, rank) %>%
summarize_at(.vars=c('wtcpue', 'wgth', 'wgt'), .funs = function(x) sum(x)) %>%
ungroup()

survey1 <- full_join(survey.num, survey.wgt,
                      by=c('Survey','HaulID','StatRec','Year','Month','Quarter',
                           'Season','ShootLat','ShootLong','HaulDur','Area.swept',
                           'Gear','Depth','SBT','SST','family','genus','taxa','AphiaID',
                           'worms_id','SpecCode',
                           'kingdom','phylum','class','order','rank'))

# summarize abundance/weight from length data
survey2 <- left_join(survey, datalw, by=c("taxa","family","genus","lme",
                                             "rank","Survey")) %>%
mutate(wgtlencpue = numlencpue*a*Length^b/1000, # divide by 1000 to get kg/km2
      wgtlenh = numlenh*a*Length^b/1000) %>% # divide by 1000 to get kg/h
group_by(Survey,HaulID,StatRec,Year,Month,Quarter,Season,ShootLat,ShootLong,HaulDur,
         Area.swept,Gear,Depth,SBT,SST,family,genus,taxa, AphiaID,worms_id,SpecCode,a, b,
         kingdom, class, order,phylum, rank) %>%
summarize_at(.vars=c('numlencpue','numlenh','wgtlencpue','wgtlenh'),
             .funs=function(x) sum(x)) %>%
ungroup()

# merge both and compare
nrow(survey1)==nrow(survey2)
survey3 <- full_join(survey1, survey2, by=c('Survey','HaulID','StatRec','Year','Month',
                                              'Quarter','Season','ShootLat','ShootLong',
                                              'HaulDur','Area.swept','Gear','Depth',
                                              'SBT','SST','family','genus','taxa',
                                              'AphiaID','worms_id','SpecCode',
                                              'kingdom','phylum','class','order','rank'))

library(ggplot2)

```

```

# correlation between abundances to check calculations are right
cor(x = survey3$numh, y = survey3$numlenh, method = 'pearson')
xx <- subset(survey3, !is.na(numcpue))
cor(x = xx$numcpue, y = xx$numlenh, method = 'pearson')

# check weights
xx <- subset(survey3, wtcpue >0 & wgtlencpue>0)
cor(x = xx$wtcpue , y = xx$wgtlencpue, method = 'pearson')

xx <- subset(survey3, wgth>0 & wgtlenth>0)
cor(x = xx$wgth, y = xx$wgtlenth, method = 'pearson')

### cor = 0.92 and 0.90 so something does not work.

#####
# CHECK PER SURVEY
#####

# no zeros
xx <- subset(survey3, wgth>0 & wgtlenth>0)

# rockall looks OK
ggplot(subset(xx, Survey=='ROCKALL'), aes(x=wgth, y=wgtlenth)) + geom_point() +
  geom_abline(intercept = 0, slope = 1, color="red",
              linetype="dashed", size=0.5) + scale_x_log10() + scale_y_log10()

# IE-IGFS looks OK
ggplot(subset(xx, Survey=='IE-IGFS'), aes(x=wgth, y=wgtlenth)) + geom_point() +
  geom_abline(intercept = 0, slope = 1, color="red",
              linetype="dashed", size=0.5) + scale_x_log10() + scale_y_log10()

# NIGFS looks OK
ggplot(subset(xx, Survey=='NIGFS'), aes(x=wgth, y=wgtlenth)) + geom_point() +
  geom_abline(intercept = 0, slope = 1, color="red",
              linetype="dashed", size=0.5) + scale_x_log10() + scale_y_log10()

# PT-IBTS looks OK
ggplot(subset(xx, Survey=='PT-IBTS'), aes(x=wgth, y=wgtlenth)) + geom_point() +
  geom_abline(intercept = 0, slope = 1, color="red",
              linetype="dashed", size=0.5) + scale_x_log10() + scale_y_log10()

# FR-CGFS looks OK
ggplot(subset(xx, Survey=='FR-CGFS'), aes(x=wgth, y=wgtlenth)) + geom_point() +
  geom_abline(intercept = 0, slope = 1, color="red",
              linetype="dashed", size=0.5) + scale_x_log10() + scale_y_log10()

# SWC-IBTS issue
ggplot(subset(xx, Survey=='SWC-IBTS'), aes(x=wgth, y=wgtlenth)) + geom_point() +
  geom_abline(intercept = 0, slope = 1, color="red",
              linetype="dashed", size=0.5) + scale_x_log10() + scale_y_log10()

comp <- subset(xx, Survey=='SWC-IBTS') %>%
  select(HaulID,wgtlenth,wgth) %>%

```

```

distinct() %>%
group_by(HaulID) %>%
summarize_at(.vars=c('wgtlenh', 'wgth'), .funs = function(x) sum(x)) %>%
ungroup() %>%
as.data.frame()

ggplot(comp, aes(x=wgth, y=wgtlenh)) + geom_point() +
geom_abline(intercept = 0, slope = 1, color="red",
            linetype="dashed", size=0.5) + scale_x_log10() + scale_y_log10()

comp$factor <- comp$wgtlenh / comp$wgth
plot(comp$factor)
resc <- comp$HaulID[comp$factor > 40]

# after check with original haul length data (HL) for some resc haulid, weight
# is clearly wrong factor 100
survey3 <- survey3 %>%
  mutate(wtcpue = if_else(HaulID %in% resc, wtcpue*100,wtcpue),
         wgth = if_else(HaulID %in% resc , wgth*100,wgth),
         wgt = if_else(HaulID %in% resc , wgt*100,wgt))

# BITS issue
ggplot(subset(xx, Survey=='BITS'), aes(x=wgth, y=wgtlenh)) + geom_point() +
geom_abline(intercept = 0, slope = 1, color="red",
            linetype="dashed", size=0.5) + scale_x_log10() + scale_y_log10()

comp <- subset(xx, Survey=='BITS') %>%
  select(HaulID,wgtlenh,wgth) %>%
  distinct() %>%
group_by(HaulID) %>%
summarize_at(.vars=c('wgtlenh', 'wgth'), .funs = function(x) sum(x)) %>%
ungroup() %>%
as.data.frame()

ggplot(comp, aes(x=wgth, y=wgtlenh)) + geom_point() +
geom_abline(intercept = 0, slope = 1, color="red",
            linetype="dashed", size=0.5) + scale_x_log10() + scale_y_log10()

comp$factor <- comp$wgtlenh / comp$wgth
plot(comp$factor)
resc <- comp$HaulID[comp$factor > 40]

# after check with original haul length data (HL) for some resc haulid, weight
# is clearly wrong factor 100
survey3 <- survey3 %>%
  mutate(wtcpue = if_else(HaulID %in% resc, wtcpue*100,wtcpue),
         wgth = if_else(HaulID %in% resc , wgth*100,wgth),
         wgt = if_else(HaulID %in% resc , wgt*100,wgt))

# EVHOE may have an issue, no changes as not very clear
ggplot(subset(xx, Survey=='EVHOE'), aes(x=wgth, y=wgtlenh)) + geom_point() +
geom_abline(intercept = 0, slope = 1, color="red",
            linetype="dashed", size=0.5) + scale_x_log10() + scale_y_log10()

```

```

comp <- subset(xx, Survey=='EVHOE') %>%
  select(HaulID,wgtlenh,wgth) %>%
  distinct() %>%
  group_by(HaulID) %>%
  summarize_at(.vars=c('wgtlenh', 'wgth'), .funs = function(x) sum(x)) %>%
  ungroup() %>%
  as.data.frame()

ggplot(comp, aes(x=wgth, y=wgtlenh)) + geom_point() +
  geom_abline(intercept = 0, slope = 1, color="red",
              linetype="dashed", size=0.5) + scale_x_log10() + scale_y_log10()

comp$factor <- comp$wgtlenh / comp$wgth
plot(comp$factor)

# NS - IBTS issue
ggplot(subset(xx, Survey=='NS-IBTS'), aes(x=wgth, y=wgtlenh)) + geom_point() +
  geom_abline(intercept = 0, slope = 1, color="red",
              linetype="dashed", size=0.5) + scale_x_log10() + scale_y_log10()

comp <- subset(xx, Survey=='NS-IBTS') %>%
  select(HaulID,wgtlenh,wgth) %>%
  distinct() %>%
  group_by(HaulID) %>%
  summarize_at(.vars=c('wgtlenh', 'wgth'), .funs = function(x) sum(x)) %>%
  ungroup() %>%
  as.data.frame()

ggplot(comp, aes(x=wgth, y=wgtlenh)) + geom_point() +
  geom_abline(intercept = 0, slope = 1, color="red",
              linetype="dashed", size=0.5) + scale_x_log10() + scale_y_log10()

comp$factor <- comp$wgtlenh / comp$wgth
comp$uni <- c(1:nrow(comp))
plot(comp$factor-comp$uni,ylim=c(0,120))
points(comp$factor[comp$factor > 20]~comp$uni[comp$factor > 20],col="red")
points(comp$factor[comp$factor > 8 & comp$factor <20]~
       comp$uni[comp$factor > 8 & comp$factor <20],col="blue")

# two issues - one estimate 100 times higher based on length, the other 10 times
resc <- comp$HaulID[comp$factor > 20]
resc2 <- comp$HaulID[comp$factor > 8 & comp$factor <20]

# after check with original haul length data (HL) for some resc haulid, weight
# is clearly wrong factor 100
# and also a cluster of factor 10
survey3 <- survey3 %>%
  mutate(wtcpue = if_else(HaulID %in% resc, wtcpue*100,wtcpue),
         wgth = if_else(HaulID %in% resc , wgth*100,wgth),
         wgt = if_else(HaulID %in% resc , wgt*100,wgt))

survey3 <- survey3 %>%
  mutate(wtcpue = if_else(HaulID %in% resc2, wtcpue*10,wtcpue),

```

```

wgth = if_else(HaulID %in% resc2 , wgth*10,wgth),
wgt = if_else(HaulID %in% resc2 , wgt*10,wgt))

# check again correlations
xx <- subset(survey3, wtcpue>0 & wgtlencpue>0)
cor(x = xx$wtcpue , y = xx$wgtlencpue, method = 'pearson') # looks better

xx <- subset(survey3, wgth>0 & wgtlenth>0)
cor(x = xx$wgth, y = xx$wgtlenth, method = 'pearson') # looks better

# now check per haul without zeros, NAs
xx <- subset(survey3, wtcpue>0 & wgtlencpue>0)

comp <- xx %>%
  select(HaulID,wgtlencpue,wtcpue) %>%
  distinct() %>%
  group_by(HaulID) %>%
  summarize_at(.vars=c('wgtlencpue', 'wtcpue'), .funs = function(x) sum(x)) %>%
  ungroup() %>%
  as.data.frame()

ggplot(comp, aes(x=wtcpue, y=wgtlencpue)) + geom_point() +
  geom_abline(intercept = 0, slope = 1, color="red",
              linetype="dashed", size=0.5) + scale_x_log10() + scale_y_log10()

cor(x = xx$wtcpue , y = xx$wgtlencpue, method = 'pearson')
# [1] 0.9635742

#####
#### Fishglob format
#####

survey4 <- survey3 %>%
  rename(survey = Survey,
         haul_id = HaulID,
         stat_rec = StatRec,
         year = Year,
         month = Month,
         quarter = Quarter,
         season = Season,
         latitude = ShootLat,
         longitude = ShootLong,
         haul_dur = HaulDur,
         area_swept = Area.swept,
         gear = Gear,
         depth = Depth,
         sbt = SBT,
         sst = SST,
         verbatim_aphia_id = AphiaID,
         aphia_id = worms_id,
         accepted_name = taxa,
         ) %>%

```

```

mutate(day = NA_integer_,
       verbatim_name = NA_character_,
       station = NA_character_,
       stratum = NA_character_,
       sub_area = NA_character_,
       continent = "europe",
       country = case_when(survey=="PT-IBTS" ~ "portugal",
                            survey=="EVHOE" ~ "france",
                            survey=="IE-IGFS" ~ "ireland",
                            survey %in% c("ROCKALL","SWC-IBTS","NIGFS") ~ "uk",
                            survey=="FR-CGFS" ~ "france",
                            survey %in% c("NS-IBTS","BITS") ~ "multi-countries"),
       num = numlencpue*area_swept,
       num_h = numlenh,
       num_cpue = numlencpue,
       wgt = wgtlencpue*area_swept,
       wgt_h = wgtlenh,
       wgt_cpue = wgtlencpue,
       haul_dur = haul_dur/60,
       source = "DATRAS ICES",
       timestamp = "2021-07") %>%
# Final format
select(fishglob_data_columns$`Column name fishglob`)

#####
# Save database in Google drive
#####

# Just run this routine should be good for all
write_clean_data(data = survey4, survey = "DATRAS_v3", overwrite = T)

```

1. Overview of the survey data table

survey	source	timestamp	haul_id	country	sub_area	continent	stat
BITS	DATRAS ICES	2021-07	BITS 1992 1 SE 77AR GOV 50 1	multi-countries	NA	europe	41G
BITS	DATRAS ICES	2021-07	BITS 1992 1 SE 77AR GOV 50 1	multi-countries	NA	europe	41G
BITS	DATRAS ICES	2021-07	BITS 1992 1 SE 77AR GOV 50 1	multi-countries	NA	europe	41G
BITS	DATRAS ICES	2021-07	BITS 1992 1 SE 77AR GOV 50 1	multi-countries	NA	europe	41G
BITS	DATRAS ICES	2021-07	BITS 1992 1 SE 77AR GOV 50 1	multi-countries	NA	europe	41G

station	stratum	year	month	day	quarter	season
NA	NA	1992	3	NA	1	NA
NA	NA	1992	3	NA	1	NA
NA	NA	1992	3	NA	1	NA
NA	NA	1992	3	NA	1	NA
NA	NA	1992	3	NA	1	NA

latitude	longitude	haul_dur	area_swept	gear	depth
56.3867	12.2567	1.05	0.1897792	GOV	34
56.3867	12.2567	1.05	0.1897792	GOV	34
56.3867	12.2567	1.05	0.1897792	GOV	34
56.3867	12.2567	1.05	0.1897792	GOV	34
56.3867	12.2567	1.05	0.1897792	GOV	34

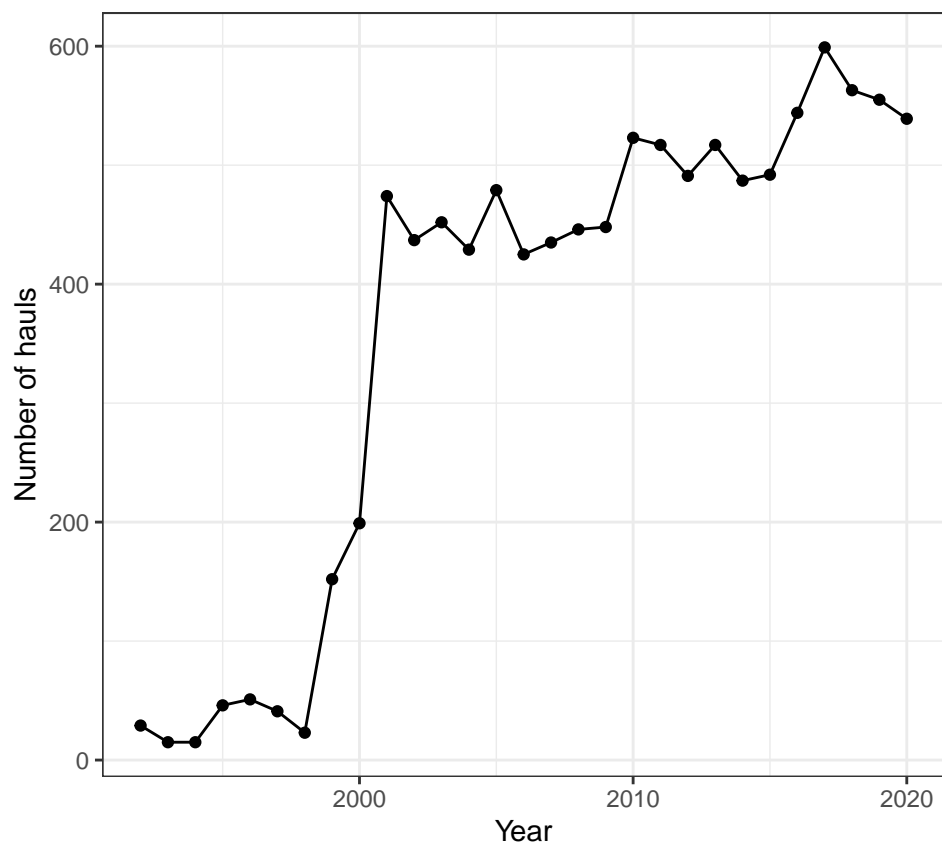
sbt	sst	num	num_h	num_cpue	wgt
NA	NA	516.60	492	2722.110865	8.7658799
NA	NA	1.05	1	5.532746	0.0483676
NA	NA	2189.25	2085	11535.774701	81.9598026
NA	NA	1770.30	1686	9328.209183	13.7967411
NA	NA	8.40	8	44.261965	7.9753360

wgt_h	wgt_cpue	verbatim_name	verbatim_aphia_id	accepted_name
8.3484571	46.1898896	NA	126793	Callionymus
0.0460644	0.2548628	NA	126792	Callionymus lyra
78.0569548	431.8692781	NA	126417	Clupea harengus
13.1397534	72.6989136	NA	126425	Sprattus sprattus
7.5955581	42.0242912	NA	126436	Gadus morhua

aphia_id	SpecCode	kingdom	phylum	class	order	family
125930	NA	Animalia	Chordata	Actinopteri	Callionymiformes	Callionymidae
126792	23	Animalia	Chordata	Actinopteri	Callionymiformes	Callionymidae
126417	24	Animalia	Chordata	Actinopteri	Clupeiformes	Clupeidae
126425	1357	Animalia	Chordata	Actinopteri	Clupeiformes	Clupeidae
126436	69	Animalia	Chordata	Actinopteri	Gadiformes	Gadidae

2. Summary of sampling intensity

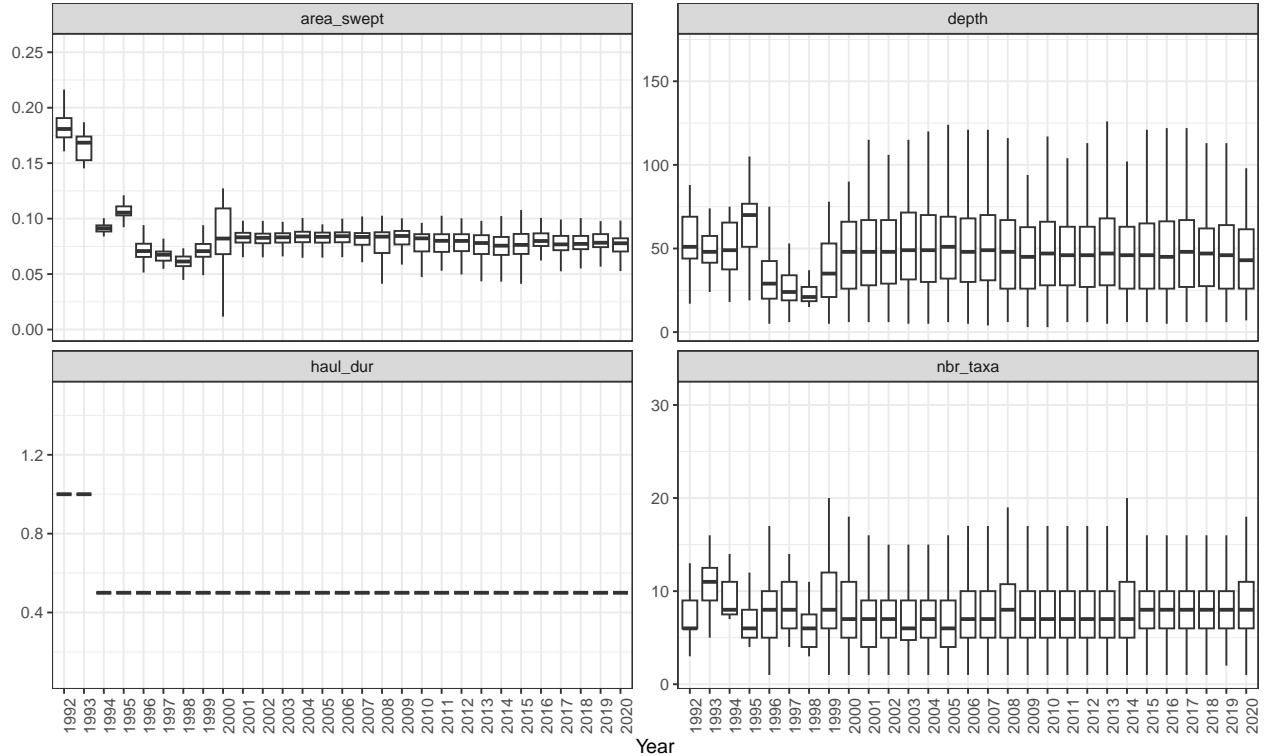
Number of hauls per year performed during the survey after data processing.



3. Summary of sampling variables from the survey

Here we show the yearly total and average of the following variables reported in the survey data:

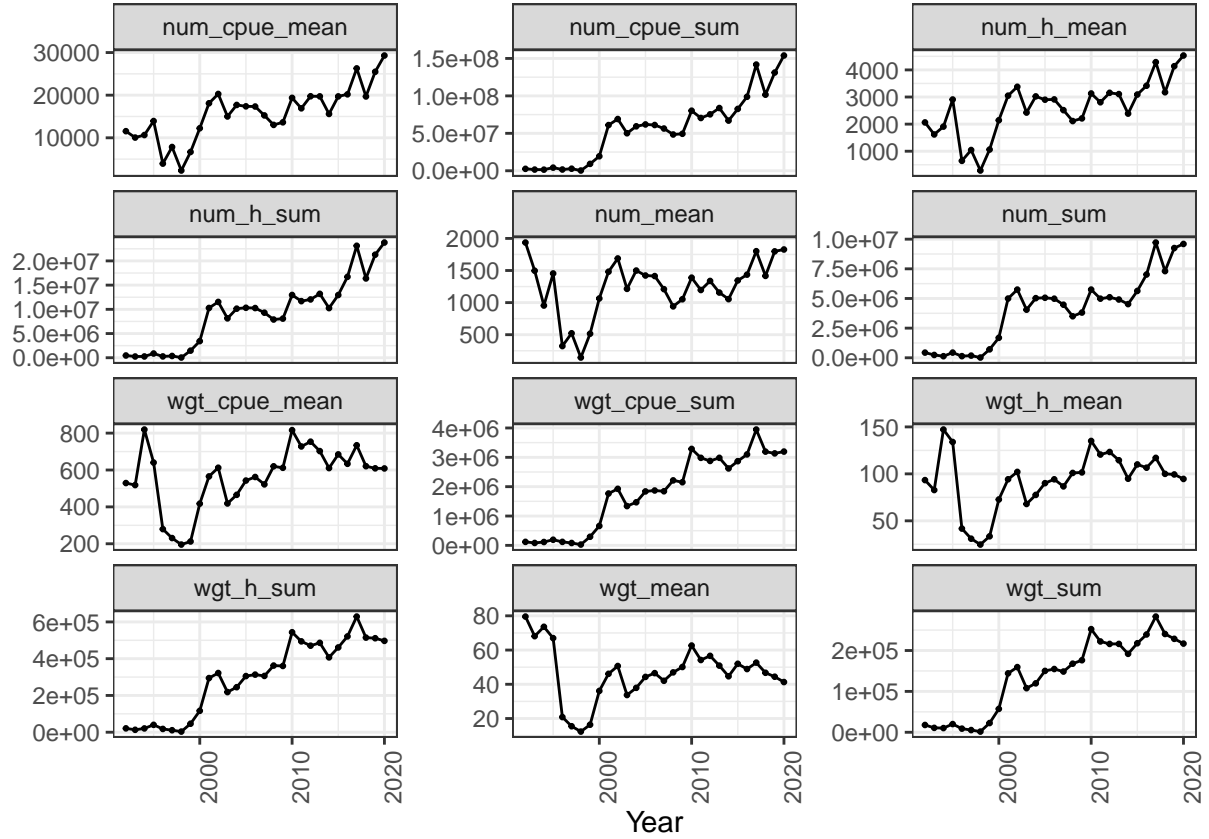
- *area_swept*, swept area by the bottom trawl gear km^2
- *depth*, sampling depth in m
- *haul_dur*, haul sampling duration *hours*
- *number of marine fish taxa*, taxa were cleaned following the last version of taxonomy from the World Register of Marine Species (<https://www.marinespecies.org/>, October 2021)



4. Summary of biological variables

Here we display the yearly total and average across hauls of the following variables recorded in the data:

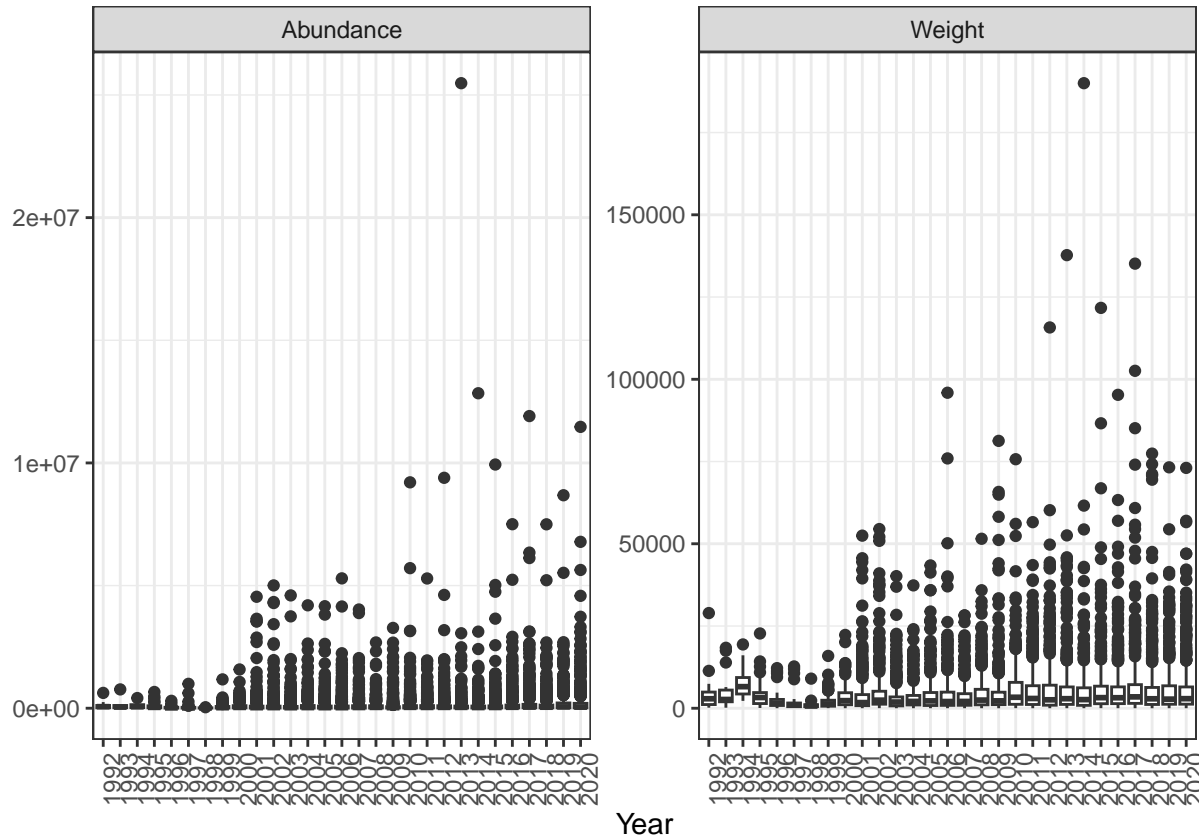
- num_cpue , number of individuals (abundance) in $\frac{individuals}{km^2}$
- num_h , number of individuals (abundance) in $\frac{individuals}{h}$
- num , number of individuals (abundance)
- wgt_cpue , weight in $\frac{kg}{km^2}$
- wgt_h , weight in $\frac{kg}{h}$
- wgt , weight in kg



5. Extreme values

Here we show a yearly total distribution of the biomass data to visualize outliers:

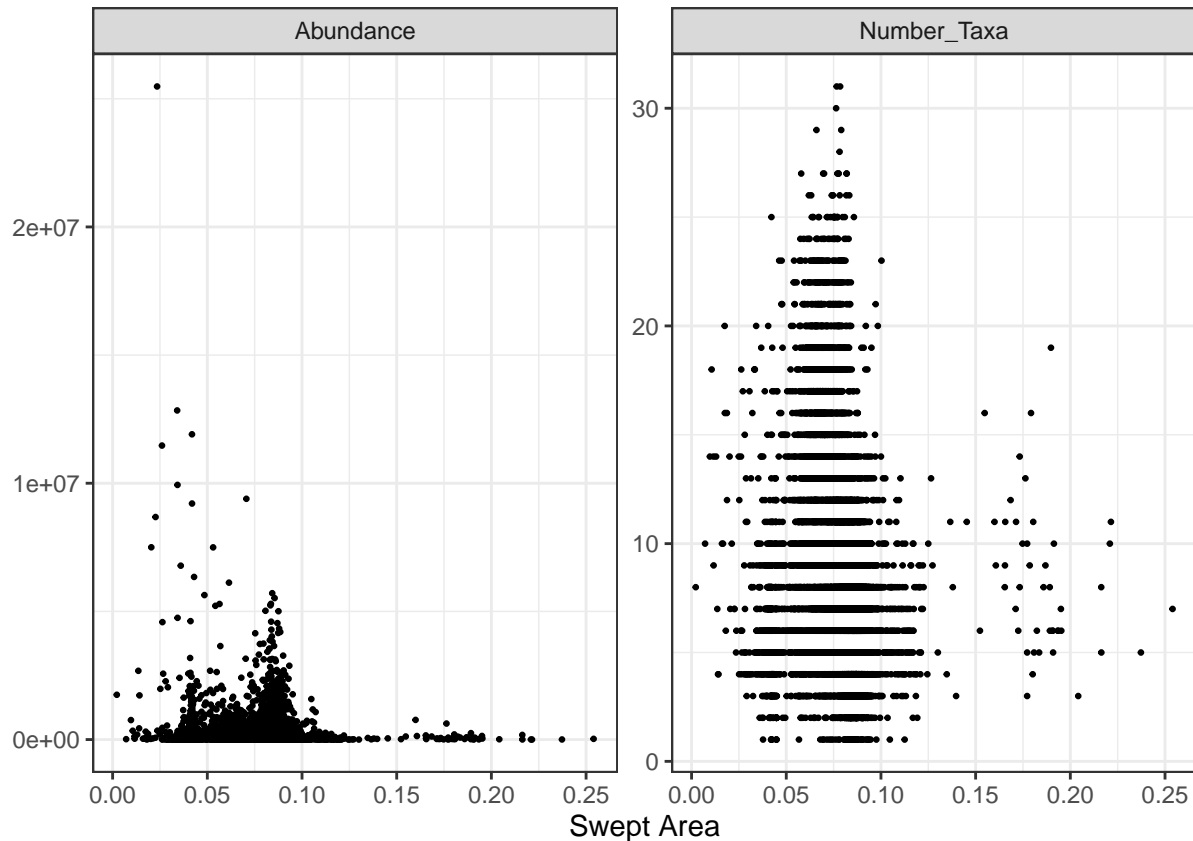
- num_cpue , number of individuals (abundance) in $\frac{individuals}{km^2}$
- wgt_cpue , weight in $\frac{kg}{km^2}$



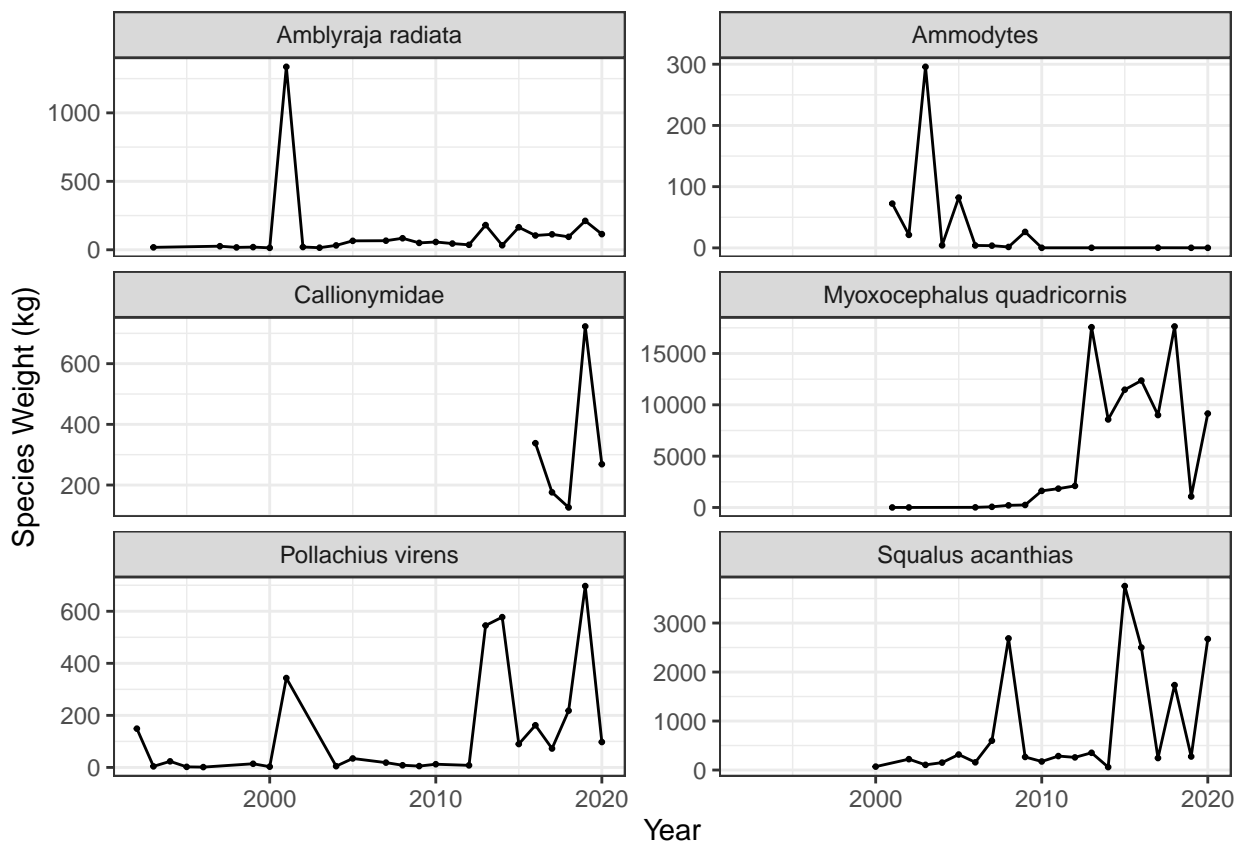
6. Summary of variables against swept area

Here we show the total abundance and number of taxa relationships with the area swept:

- nbr_taxa , number of marine fish taxa after taxonomic data cleaning
- num_cpue , number of individuals (abundance) in $\frac{individuals}{km^2}$
- wgt_cpue , weight in $\frac{kg}{km^2}$

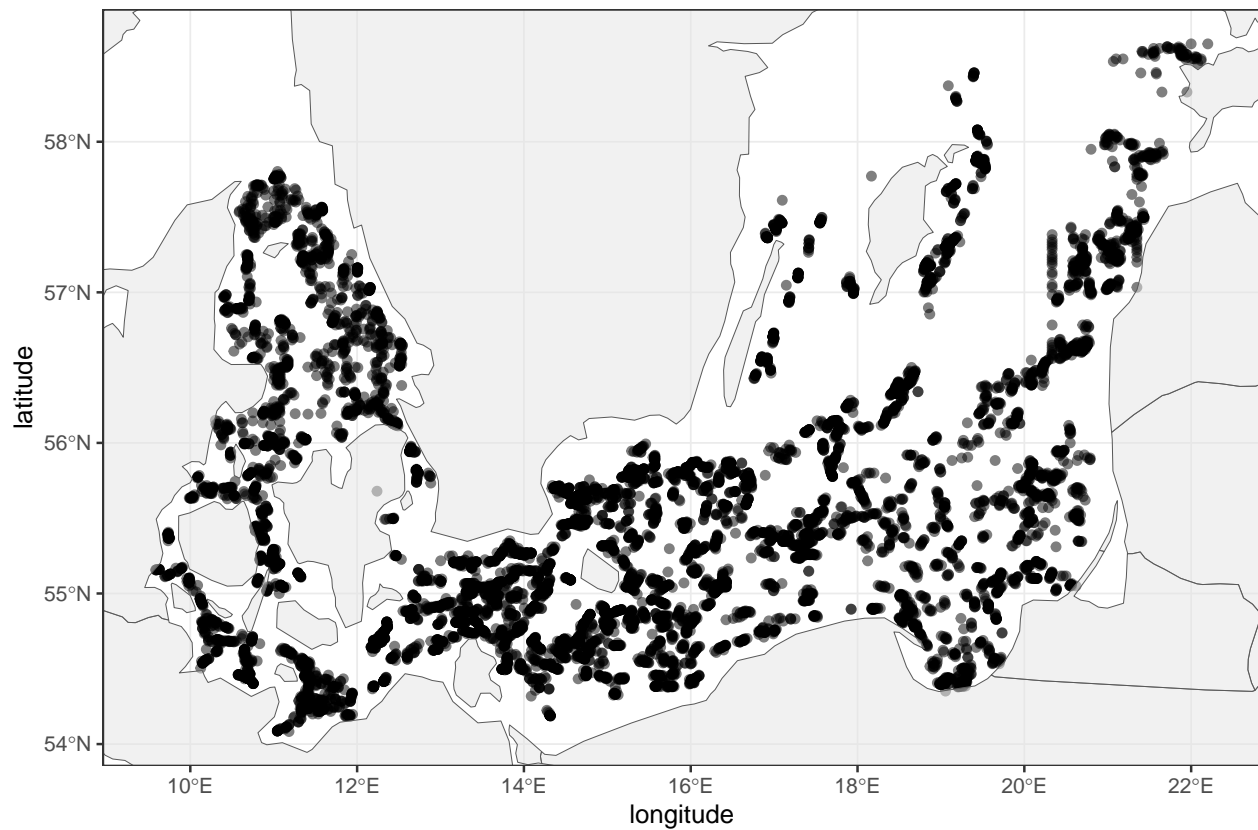


7. Abundance or Weight trends of the six most abundant species



8. Distribution mapping

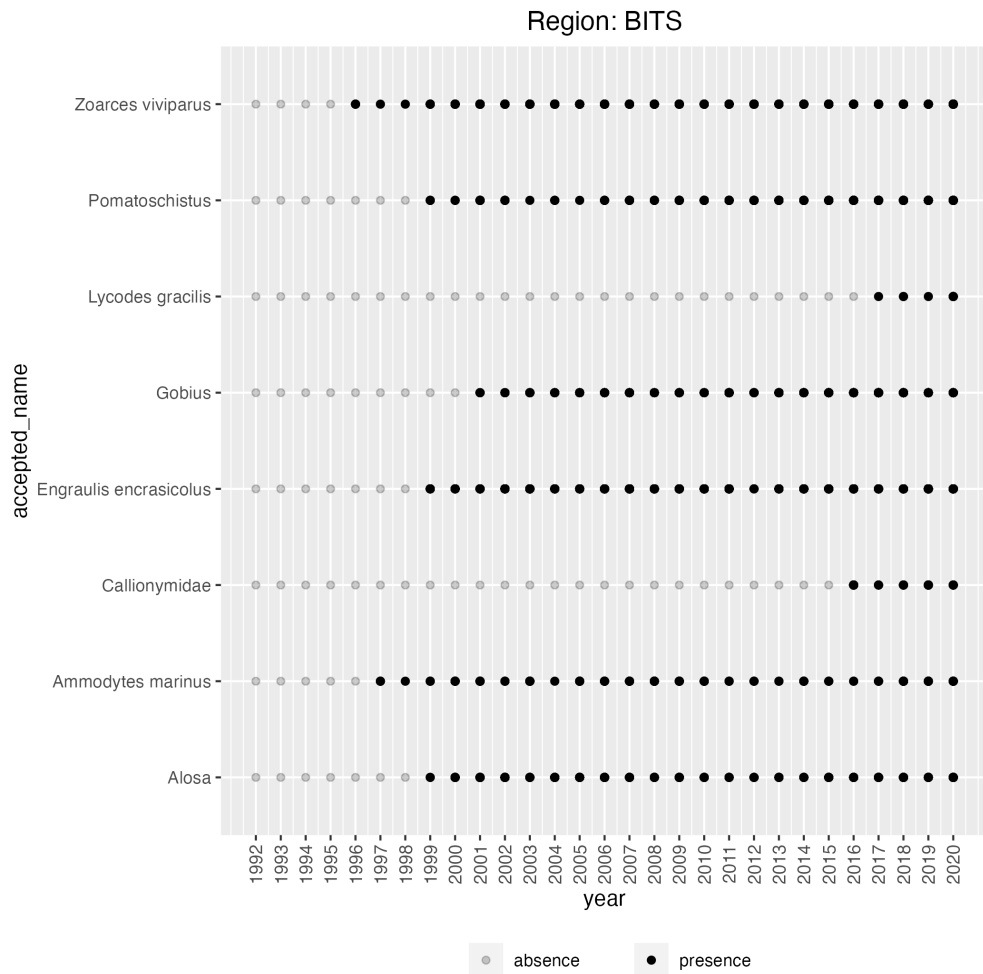
Map of the sampling distribution in space. Note that we only show one year per coordinate.



9. Taxonomic flagging

This species flagging method was adapted from <https://github.com/pinskylab/OceanAdapt/blob/master/R/add-spp-to-taxonomy.Rmd#L33>

Visualization of flagged taxa



Statistics related to the taxonomic flagging outputs

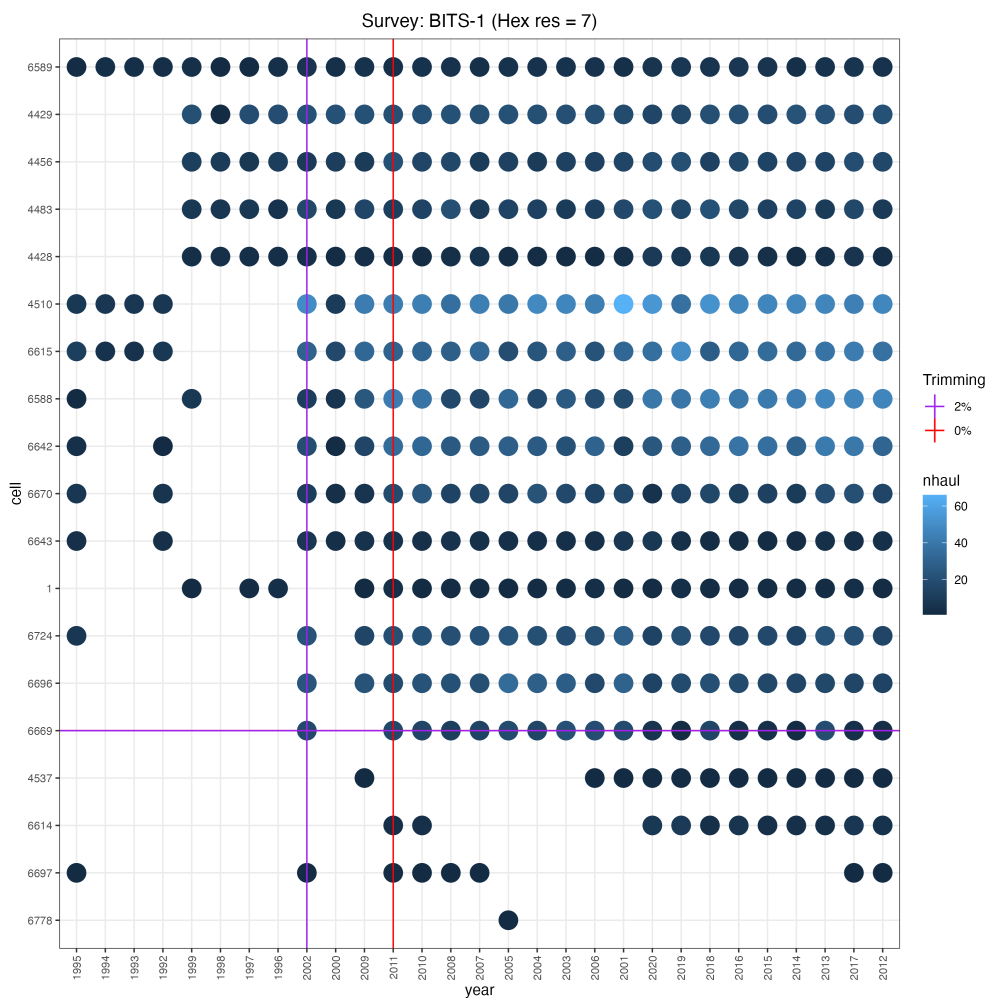
Total number of species	126.0
Percentage of species flagged	6.3

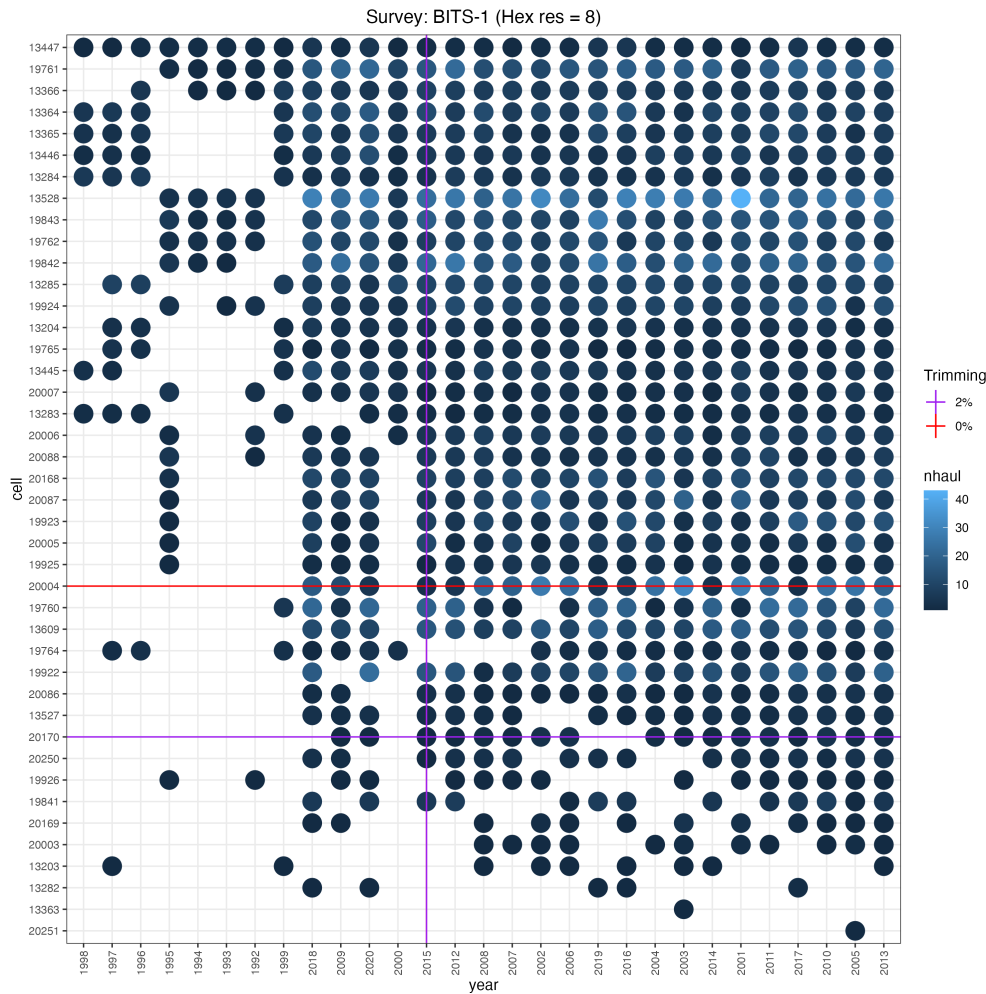
10. Spatio-temporal standardization: BITS-1

a. Standardization method 1

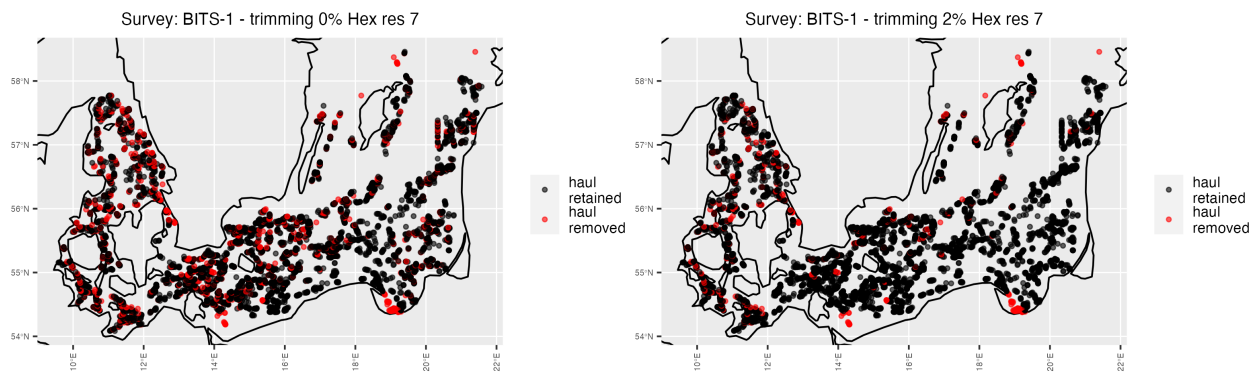
This standardization method was adapted from https://github.com/zoekitchel/trawl_spatial_turnover/blob/master/data_prep_code/species/explore_NorthSea_trimming.Rmd
It was run for hex resolution 7 and 8.

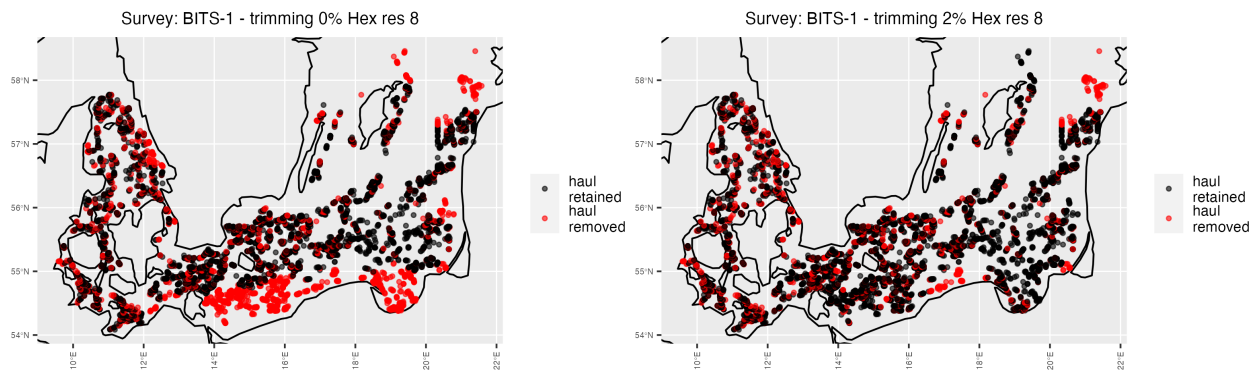
Plot of number of cells x years with overlaid flagging options



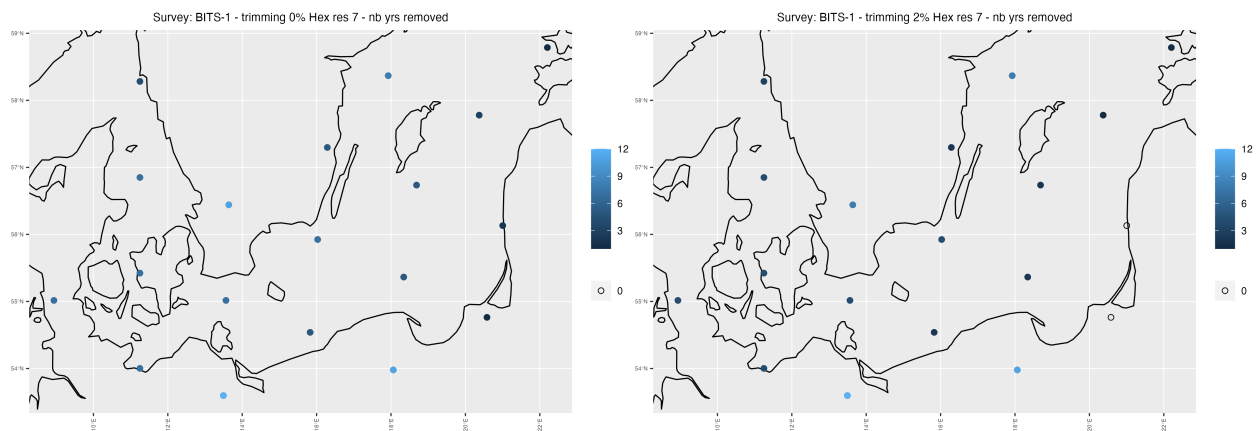


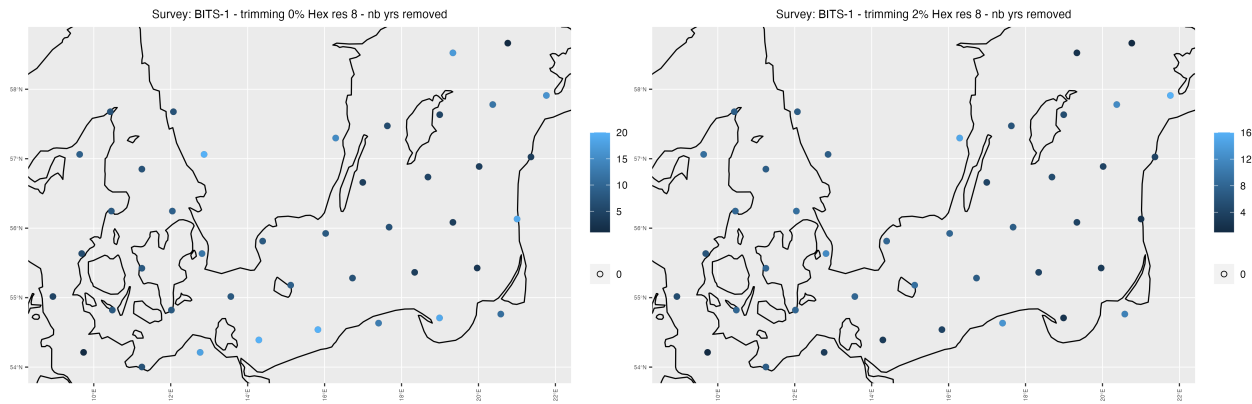
Map of hauls retained and removed per flagging method and threshold





Map of numbers of years removed per grid cell and flagging method/threshold

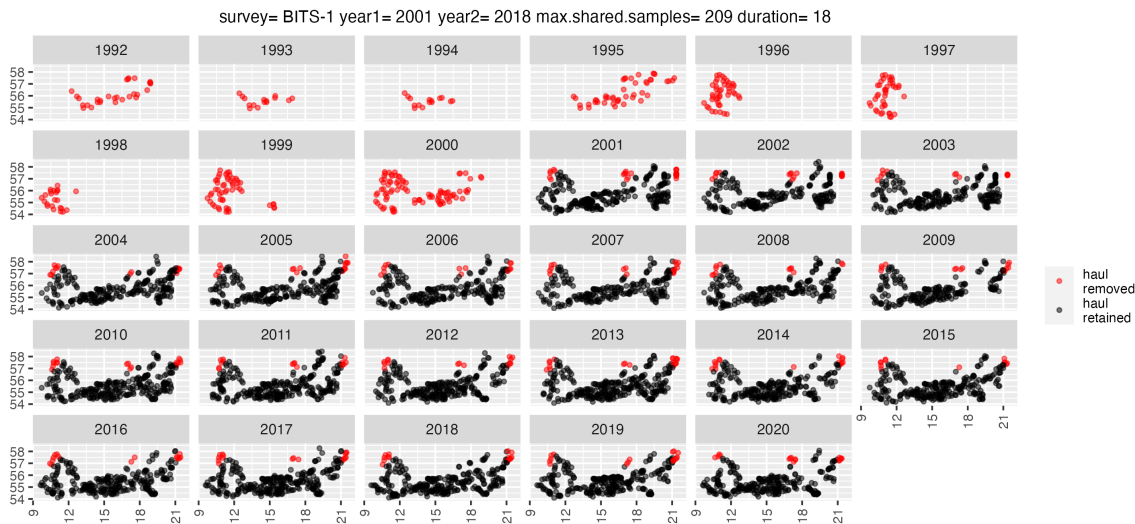




b. Standardization method 2

This standardization method was adapted from BioTIME code from https://github.com/Wubing-Xu/Range_size_winners_losers

Map of hauls retained and removed



c. Standardization summary

Statistics of hauls removed for each standardization method

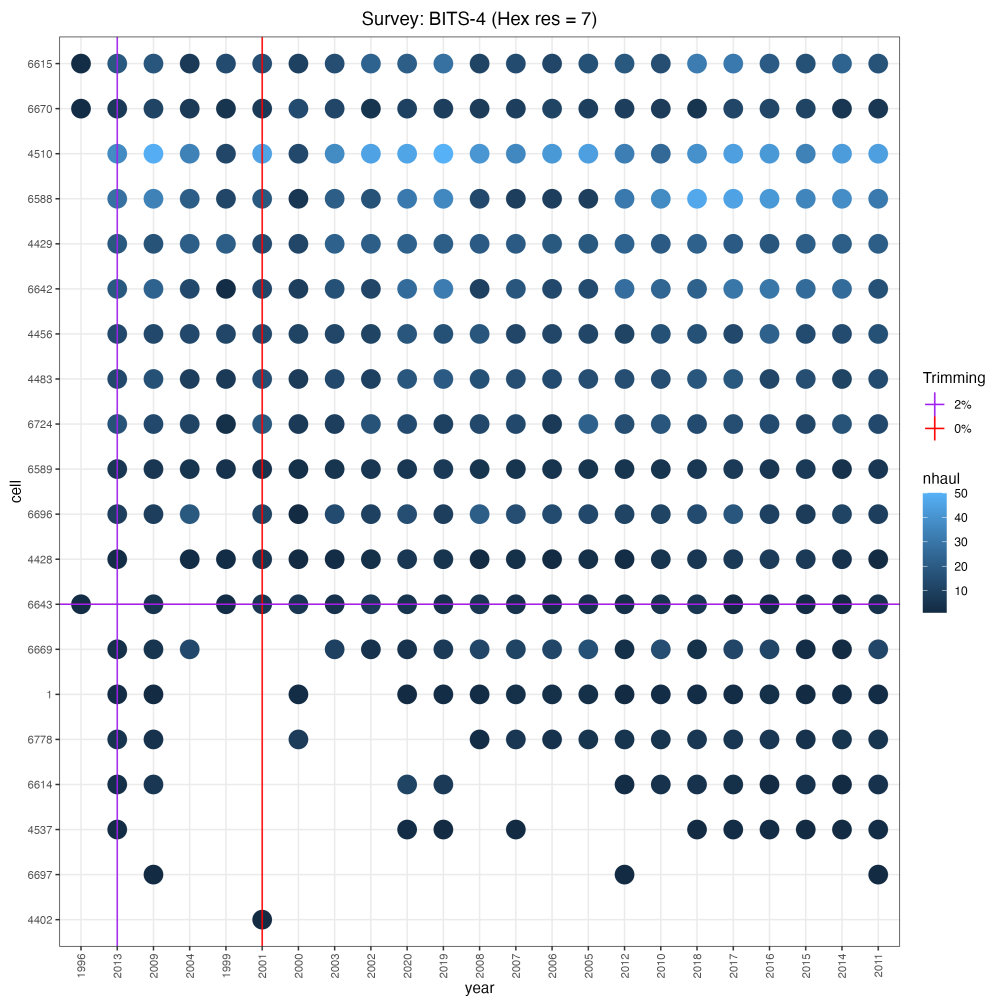
summary	grid cell 7, 0% threshold	grid cell 7, 2% threshold	grid cell 8, 0% threshold	grid cell 8, 2% threshold	method 2 (biotime)
number of hauls removed	900.0	351.0	2058.0	1309.0	5982.0
percentage of hauls removed	15.5	6.1	35.5	22.6	13.3

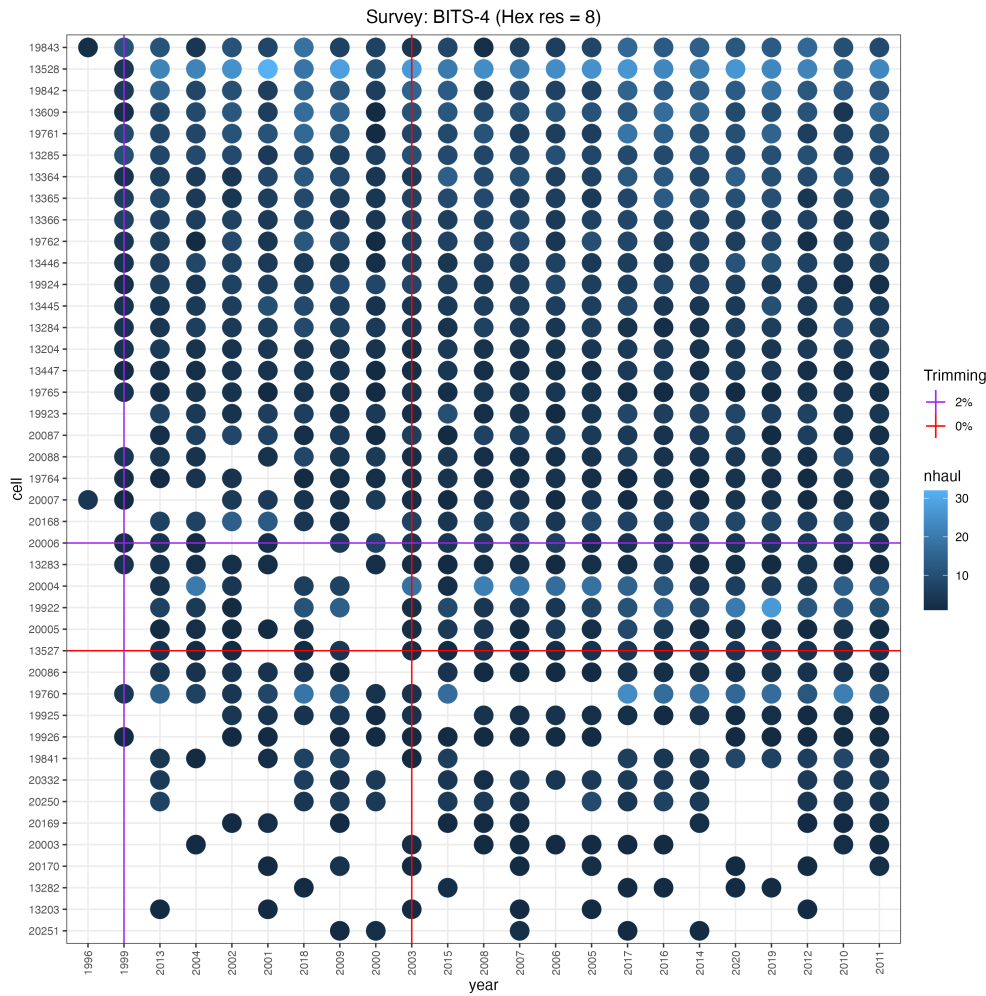
11. Spatio-temporal standardization: BITS-4

a. Standardization method 1

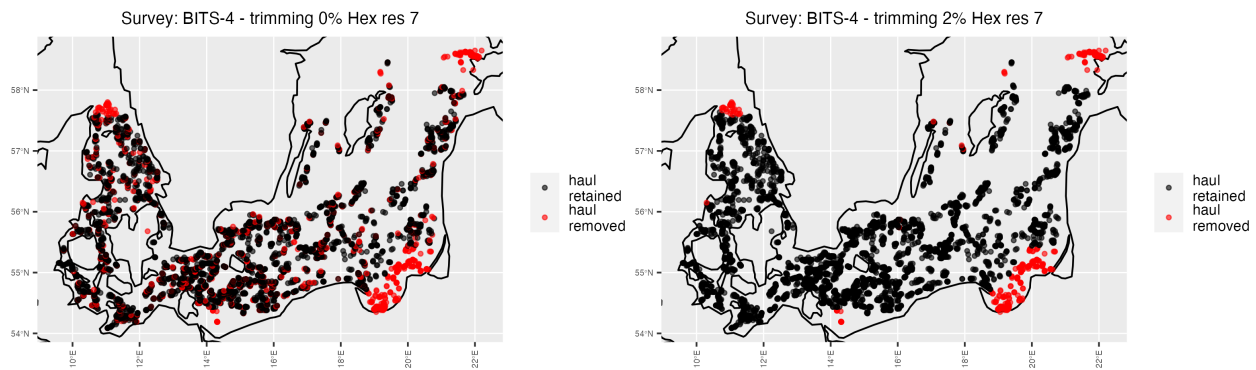
This standardization method was adapted from https://github.com/zoekitchel/trawl_spatial_turnover/blob/master/data_prep_code/species/explore_NorthSea_trimming.Rmd
It was run for hex resolution 7 and 8.

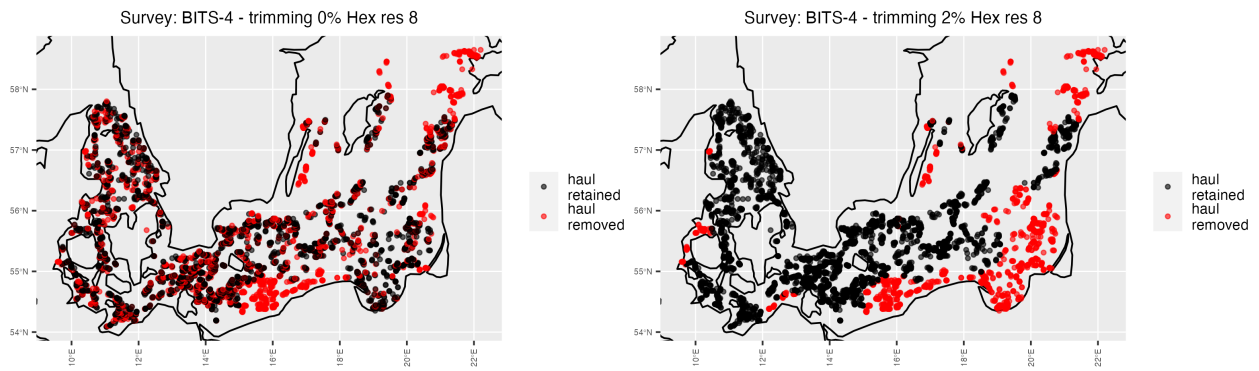
Plot of number of cells x years with overlaid flagging options



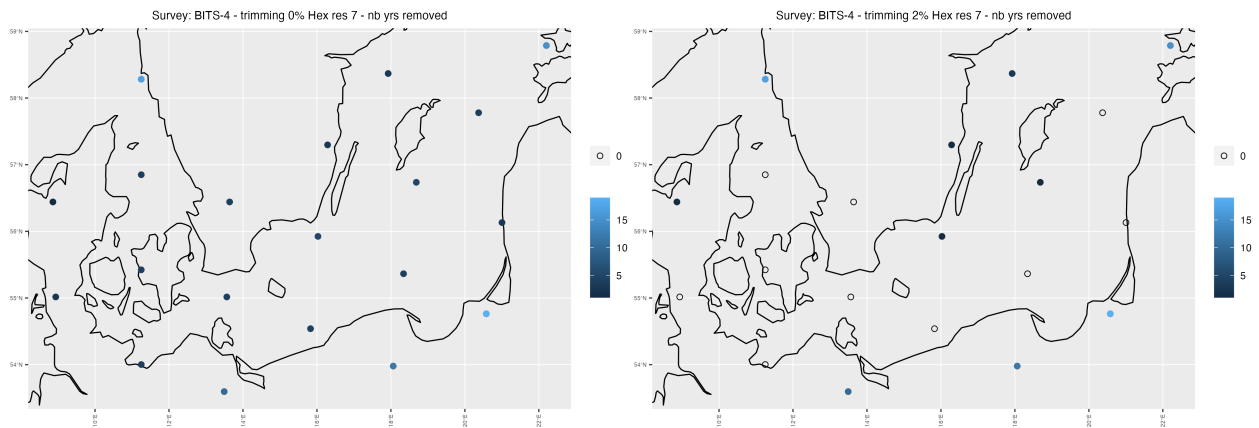


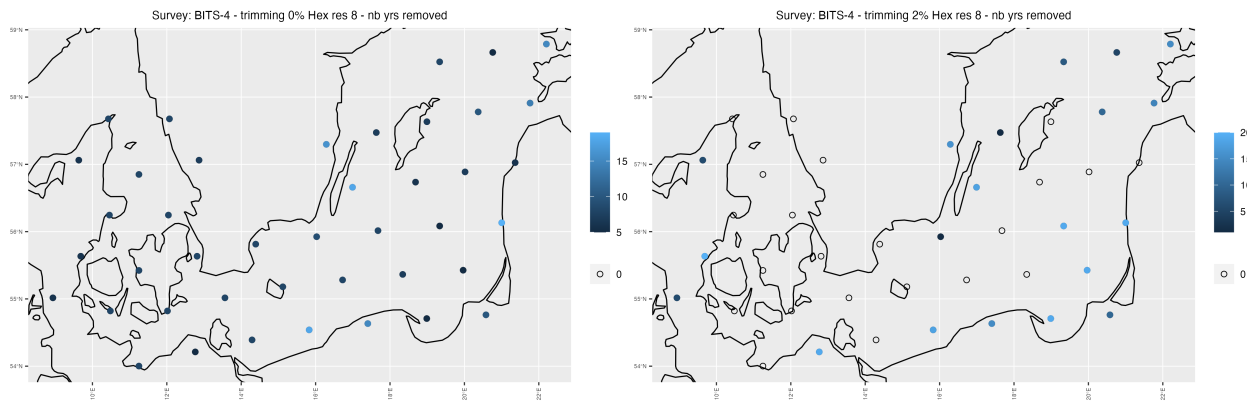
Map of hauls retained and removed per flagging method and threshold





Map of numbers of years removed per grid cell and flagging method/threshold

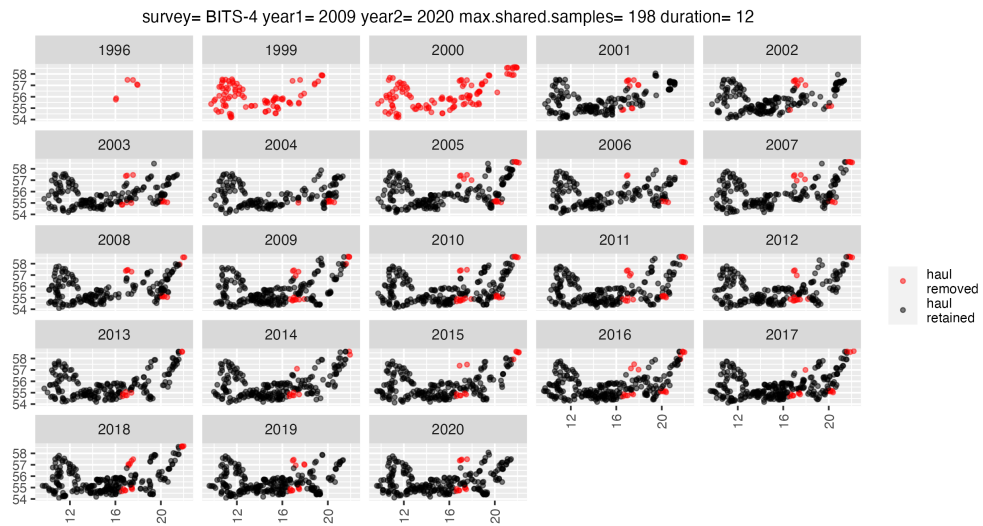




b. Standardization method 2

This standardization method was adapted from BioTIME code from https://github.com/Wubing-Xu/Range_size_winners_losers

Map of hauls retained and removed



c. Standardization summary

Statistics of hauls removed for each standardization method

summary	grid cell 7, 0% threshold	grid cell 7, 2% threshold	grid cell 8, 0% threshold	grid cell 8, 2% threshold	method 2 (biotime)
number of hauls removed	1003.0	342.0	1878.0	1140.0	3277.0
percentage of hauls removed	21.7	7.4	40.6	24.6	7.8