WCTRI: West Coast US Triennial survey data processing summary

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General info

This document presents the cleaning code and summary of the West Coast US Triennial bottom trawl survey provided by Aimee Keller and John Buchanan. It contains data from 1977 and up to 2004.

Data cleaning in R

```
#### R code to clean trawl survey West Coast US Triennial Survey (WCTRI)
#### Public data Ocean Adapt
#### Contacts: Aimee Keller
                        smartt@dnr.sc.gov, Fisheries Research Surveys Supervisor,
####
            NOAA, NMFS, NWFSC, FRAM
#####
          John Buchanan john.buchanan@noaa.gov Fisheries Biologist,
             Groundfish Ecology Program, Northwest Fisheries Science Center
#### Coding: Michelle Stuart, Dan Forrest, Zoë Kitchel November 2021
#### LOAD LIBRARIES AND FUNCTIONS ####
library(rfishbase) #needs R 4.0 or more recent
library(tidyverse)
library(lubridate)
```

```
library(googledrive)
library(taxize) # for getting correct species names
library(magrittr) # for names wrangling
source("functions/clean taxa.R")
source("functions/write_clean_data.R")
#Data for the West Coast US can be best accessed using the public Pinsky
#Lab Ocean Adapt Git Hub Repository.
#Contact malin.pinsky@rutgers.edu for questions or help accessing
#### PULL IN AND EDIT RAW DATA FILES ####
wctri_catch <- read_csv(</pre>
  "https://github.com/pinskylab/OceanAdapt/raw/master/data_raw/wctri_catch.csv",
  col_types = cols(
 CRUISEJOIN = col_integer(),
 HAULJOIN = col_integer(),
 CATCHJOIN = col_integer(),
 REGION = col_character(),
 VESSEL = col_integer(),
 CRUISE = col_integer(),
 HAUL = col_integer(),
  SPECIES_CODE = col_integer(),
 WEIGHT = col_double(),
 NUMBER_FISH = col_integer(),
 SUBSAMPLE_CODE = col_character(),
 VOUCHER = col_character(),
 AUDITJOIN = col_integer()
))
wctri_haul <- read_csv(</pre>
  "https://raw.githubusercontent.com/pinskylab/OceanAdapt/master/data_raw/wctri_haul.csv",
  col_types =
                         cols(
                           CRUISEJOIN = col_integer(),
                           HAULJOIN = col_integer(),
                           REGION = col_character(),
                           VESSEL = col_integer(),
                           CRUISE = col_integer(),
                           HAUL = col_integer(),
                           HAUL_TYPE = col_integer(),
                           PERFORMANCE = col_double(),
                           START_TIME = col_character(),
                           DURATION = col_double(),
                           DISTANCE_FISHED = col_double(),
                           NET_WIDTH = col_double(),
    #Net widths ranged from 9.8 to 17.6 m, with a standard deviation of 0.96 m.
                           NET_MEASURED = col_character(),
                           NET_HEIGHT = col_double(),
```

```
STRATUM = col_integer(),
                           START_LATITUDE = col_double(),
                           END_LATITUDE = col_double(),
                           START_LONGITUDE = col_double(),
                           END_LONGITUDE = col_double(),
                           STATIONID = col_character(),
                           GEAR_DEPTH = col_integer(),
                           BOTTOM_DEPTH = col_integer(),
                           BOTTOM_TYPE = col_integer(),
                           SURFACE_TEMPERATURE = col_double(),
                           GEAR_TEMPERATURE = col_double(),
                           WIRE_LENGTH = col_integer(),
                           GEAR = col_integer(),
                           ACCESSORIES = col_integer(),
                           SUBSAMPLE = col_integer(),
                           AUDITJOIN = col_integer()
                         ))
wctri_species <- read_csv(</pre>
"https://raw.githubusercontent.com/pinskylab/OceanAdapt/master/data_raw/wctri_species.csv",
 col_types = cols(
 SPECIES_CODE = col_integer(),
 SPECIES_NAME = col_character(),
 COMMON_NAME = col_character(),
 REVISION = col_character(),
 BS = col_character(),
 GOA = col_character(),
 WC = col_character(),
 AUDITJOIN = col_integer()
))
#### REFORMAT AND MERGE DATA FILES ####
# Add haul info to catch data
wctri <- left_join(wctri_catch, wctri_haul, by = c(</pre>
                          "CRUISEJOIN", "HAULJOIN", "VESSEL", "CRUISE", "HAUL"))
# add species names
wctri <- left_join(wctri, wctri_species, by = "SPECIES_CODE")</pre>
wctri <- wctri %>%
  # trim to standard hauls and good performance (applicable to fishglob too)
  filter(HAUL_TYPE == 3 & PERFORMANCE == 0) %>%
  # Create a unique haul_id
 mutate(
   haul_id = paste(formatC(VESSEL, width=3, flag=0), CRUISE,
                    formatC(HAUL, width=3, flag=0), START_LONGITUDE,
                    START_LATITUDE, sep=''),
    # Extract year where needed
```

```
year = substr(CRUISE, 1, 4),
   month = substr(CRUISE, 5,6),
   day = NA,
   quarter = case_when(month %in% c(1,2,3) ~ 1,
                        month %in% c(4,5,6) \sim 2,
                        month %in% c(7,8,9) \sim 3,
                        month %in% c(10,11,12) ~ 4),
   season = NA character ,
    # Add "strata" (define by lat, lon and depth bands) where needed # degree bins
    # 100 m bins # no need to use lon grids on west coast (so narrow)
   stratum = paste(floor(START_LATITUDE)+0.5, floor(BOTTOM_DEPTH/100)*100 + 50, sep= "-"),
   area_swept = DISTANCE_FISHED*(NET_WIDTH/1000), #distanced_fished in km *
    #net_width in meters / 1000 m/km
    # adjust for tow area # weight per km (1000 m2)
   wgt_cpue = WEIGHT/area_swept, #kq/km^2
   wgt_h = WEIGHT/DURATION, #kq/hour
   num_h = NUMBER_FISH/DURATION, # ind/hour
   num_cpue = NUMBER_FISH/area_swept #ind/km2
  )
wctri <- wctri %>%
  rename(
   haul dur = DURATION, #DURATION is in hours
   svvessel = VESSEL,
   latitude = START LATITUDE,
   longitude = START LONGITUDE,
   depth = BOTTOM DEPTH,
   spp = SPECIES_NAME,
   sst = SURFACE_TEMPERATURE,
   num = NUMBER_FISH,
   gear = GEAR,
   station = STATIONID,
   verbatim_name = SPECIES_NAME,
   wgt = WEIGHT
 ) %>%
  filter(
    verbatim_name != "" &
      !grepl("egg", verbatim_name)
  ) %>%
  # adjust spp names
  mutate(verbatim_name = ifelse(grepl("Lepidopsetta", verbatim_name),
                                "Lepidopsetta sp.", verbatim_name),
         verbatim_name = ifelse(grepl("Bathyraja", verbatim_name),
                                 'Bathyraja sp.', verbatim_name),
         verbatim_name = ifelse(grepl("Squalus", verbatim_name),
                                 'Squalus suckleyi', verbatim_name),
         sbt = NA,) \%>\%
  # add survey column
  mutate(survey = "WCTRI",
         source = "NOAA",
         timestamp = mdy("02/06/2019"),
         country = "United States",
         continent = "n_america",
```

```
sub_area = NA,
         stat_rec = NA) %>%
  select(survey, haul_id, source, timestamp,
         country, sub_area, continent, stat_rec, station,
         stratum, year, month, day, quarter, season, latitude, longitude,
         haul_dur, area_swept, gear, depth, sbt, sst,
         num, num_h, num_cpue, wgt, wgt_h, wgt_cpue, verbatim_name)
#sum duplicates
wctri <- wctri %>%
  group_by(survey,
           source, timestamp,
           haul_id, country, sub_area, continent, stat_rec, station, stratum,
           year, month, day, quarter, season, latitude, longitude, haul_dur, area_swept,
           gear, depth, sbt, sst,verbatim_name) %>%
  summarise(num = sum(num, na.rm = T),
           num_h = sum(num_h, na.rm = T),
           num_cpue = sum(num_cpue, na.rm = T),
           wgt = sum(wgt, na.rm = T),
            wgt_h = sum(wgt_h, na.rm = T),
            wgt_cpue = sum(wgt_cpue, na.rm = T)) %>% ungroup()
#check for duplicates, should not be any with more than 1 obs
#check for duplicates
count_wctri <- wctri %>%
  group_by(haul_id, verbatim_name) %>%
 mutate(count = n())
#none!
#which ones are duplicated?
unique_name_match <- count_wctri %>%
  group_by(verbatim_name) %>%
 filter(count>1) %>%
 distinct(verbatim_name)
unique_name_match
#empty
#the following verbatim names are duplicated within haul_id without fix above
#Bathyraja sp.
#Actinauge verrilli
#Rossia pacifica
#Sebastes alutus
#### INTEGRATE CLEAN TAXA FROM TAXA ANALYSIS ####
# Get WoRM's id for sourcing
wrm <- gnr_datasources() %>%
 filter(title == "World Register of Marine Species") %>%
```

```
pull(id)
### Automatic cleaning
# Set Survey code
wctri_survey_code <- "WCTRI"</pre>
wctri <- wctri %>%
 mutate(
   taxa2 = str_squish(verbatim_name),
   taxa2 = str_remove_all(taxa2," spp. | sp. | spp | sp | NO "),
   taxa2 = str_to_sentence(str_to_lower(taxa2)))
# Get clean taxa (setting save = T means we will get an output of missing taxa)
clean_auto <- clean_taxa(unique(wctri$taxa2), input_survey = wctri_survey_code)</pre>
# takes 20 mins!
#This cuts out the following species which are all inverts
#Cheiraster dawsoni
#Crangon communis
#Cancer gracilis
#Cancer anthonyi
#### INTEGRATE CLEAN TAXA in WCTRI survey data ####
clean_taxa <- clean_auto %>%
  select(-survey)
clean_wctri <- left_join(wctri, clean_taxa, by=c("taxa2"="query")) %>%
  filter(!is.na(taxa)) %>% # query does not indicate taxa entry that were removed in the cleaning proce
  # so all NA taxa have to be removed from the surveys because: non-existing, non marine or non fish
  rename(accepted_name = taxa,
         aphia_id = worms_id) %>%
  mutate(verbatim_aphia_id = NA) %>%
  select(survey, haul_id, source, timestamp,
         country, sub_area, continent, stat_rec, station, stratum,
         year, month, day, quarter, season, latitude, longitude,
         haul_dur, area_swept, gear, depth, sbt, sst, num, num_h, num_cpue, wgt, wgt_h, wgt_cpue,
         verbatim_name, verbatim_aphia_id, accepted_name, aphia_id, SpecCode,
         kingdom, phylum, class, order, family, genus, rank)
#check for duplicates
count_clean_wctri <- clean_wctri %>%
  group_by(haul_id, accepted_name) %>%
 mutate(count = n())
#none!
#which ones are duplicated?
unique_name_match <- count_clean_wctri %>%
  group_by(verbatim_name, accepted_name) %>%
```

1. Overview of the survey data table

survey	haul_id	source	timestamp	country	sub_area	continent
WCTRI	004197701002-119.3334.09	NOAA	2019-02-06	United States	NA	n_america
WCTRI	004197701002 - 119.3334.09	NOAA	2019-02-06	United States	NA	n _america
WCTRI	004197701002-119.3334.09	NOAA	2019-02-06	United States	NA	n _america
WCTRI	004197701002 - 119.3334.09	NOAA	2019-02-06	United States	NA	$n_america$
WCTRI	004197701002-119.3334.09	NOAA	2019-02-06	United States	NA	$n_america$

stat_rec	station	stratum	year	month	day	quarter	season
NA	NA	34.5 - 250	1977	1	NA	NA	NA
NA	NA	34.5 - 250	1977	1	NA	NA	NA
NA	NA	34.5 - 250	1977	1	NA	NA	NA
NA	NA	34.5 - 250	1977	1	NA	NA	NA
NA	NA	34.5 - 250	1977	1	NA	NA	NA

latitude	longitude	haul_dur	area_swept	gear	depth
34.09	-119.33	0.5	0.032256	160	254
34.09	-119.33	0.5	0.032256	160	254
34.09	-119.33	0.5	0.032256	160	254
34.09	-119.33	0.5	0.032256	160	254
34.09	-119.33	0.5	0.032256	160	254

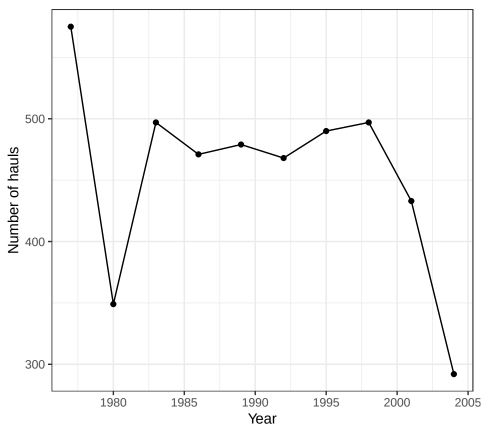
sbt	sst	num	num_h	num_cpue	wgt
NA	16.2	2	4	62.00397	1.81
NA	16.2	1	2	31.00198	0.45
NA	16.2	2	4	62.00397	0.13
NA	16.2	9	18	279.01786	2.72
NA	16.2	1	2	31.00198	0.45

wgt_h	wgt_cpue	verbatim_name	verbatim_aphia_id	accepted_name
3.62	56.113591	Eopsetta jordani	NA	Eopsetta jordani
0.90	13.950893	Hippoglossina stomata	NA	Hippoglossina stomata
0.26	4.030258	Microstomus pacificus	NA	Microstomus pacificus
5.44	84.325397	Parophrys vetulus	NA	Parophrys vetulus
0.90	13.950893	Pleuronichthys verticalis	NA	Pleuronichthys verticalis

aphia_id	${\bf SpecCode}$	kingdom	phylum	class	order	family
280690	4237	Animalia	Chordata	Actinopteri	Pleuronectiformes	Pleuronectidae
275827	4225	Animalia	Chordata	Actinopteri	Pleuronectiformes	Paralichthyidae
274294	4247	Animalia	Chordata	Actinopteri	Pleuronectiformes	Pleuronectidae
254393	4248	Animalia	Chordata	Actinopteri	Pleuronectiformes	Pleuronectidae
282295	4254	Animalia	Chordata	Actinopteri	Pleuronectiformes	Pleuronectidae

2. Summary of sampling intensity

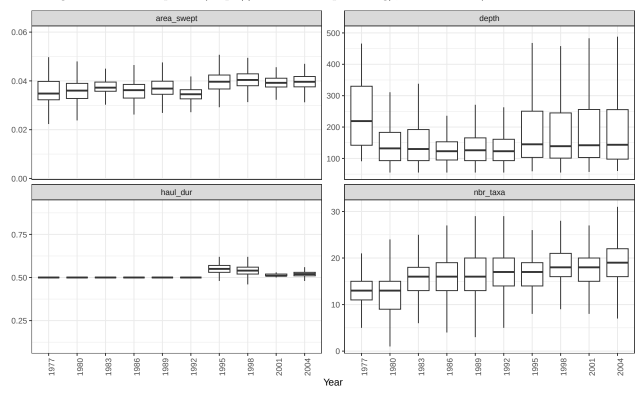
Number of hauls per year performed during the survey after data processing.



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:

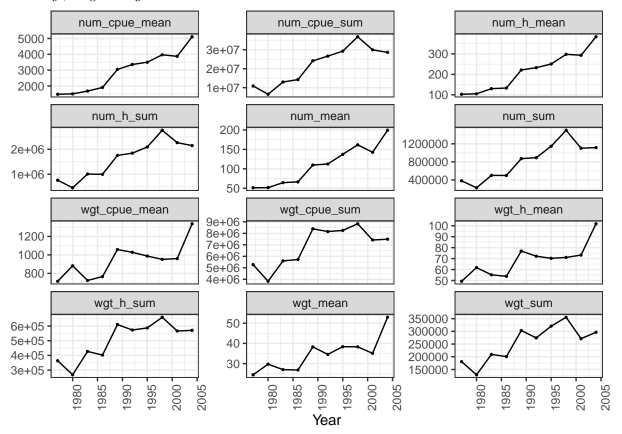
- $area_swept$, swept area by the bottom trawl gear km^2
- depth, sampling depth in m
- haul_dur, haul sampling duration hour
- number of marine fish taxa, taxa were cleaned following the last version of taxonomy from the World Register of Marine Species (https://www.marinespecies.org/, October 2021)



4. Summary of biological variables

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

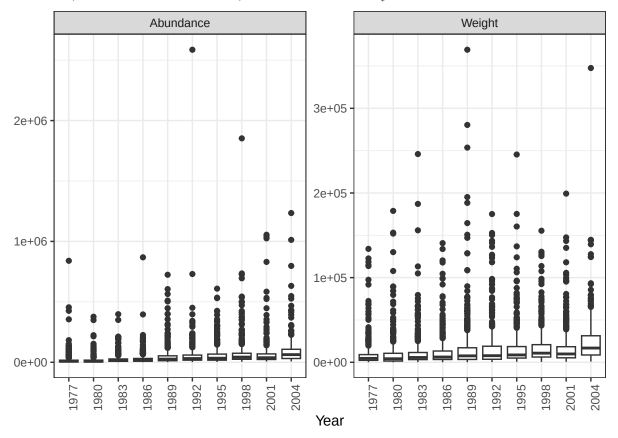
- num_cpue , number of individuals (abundance) in $\frac{individuals}{km^2}$
- num_h , number of individuals (abundance) in $\frac{individuals}{h}$
- num, number of individuals (abundance)
- wgt_cpue , weight in $\frac{kg}{km^2}$
- wgt_h , weight in $\frac{kg}{h}$
- wgt, weight in kg



5. Extreme values

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

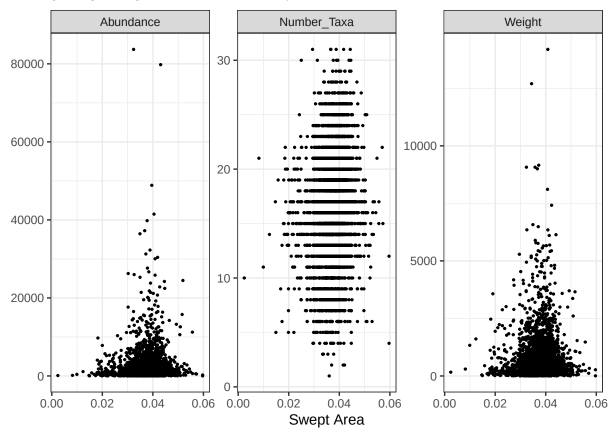
- wgt, total weight in kg per haul and year per haul and year, if available in the survey data
- num, total number of individuals, if available in the survey data



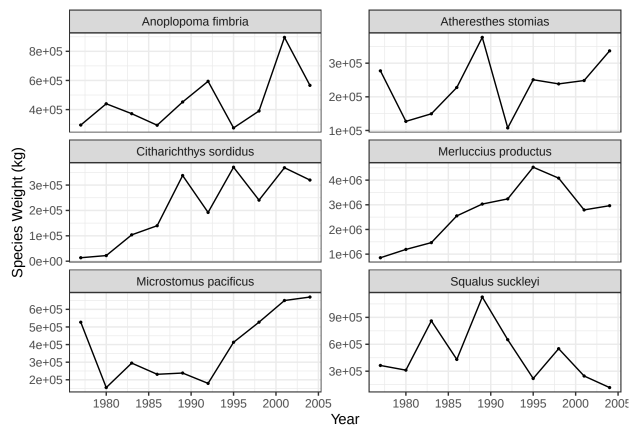
6. Summary of variables against swept area

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, number of individuals, if available in the survey data
- wgt, weight in kg, if available in the survey data

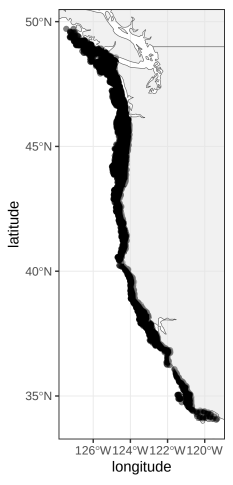


7. Abundance or Weight trends of the six most abundant species



8. Distribution mapping

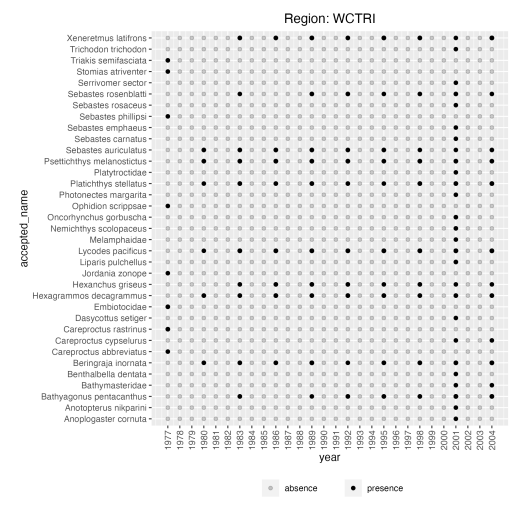
Map of the sampling distribution in space. Note that we only show one year per coordinate.



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

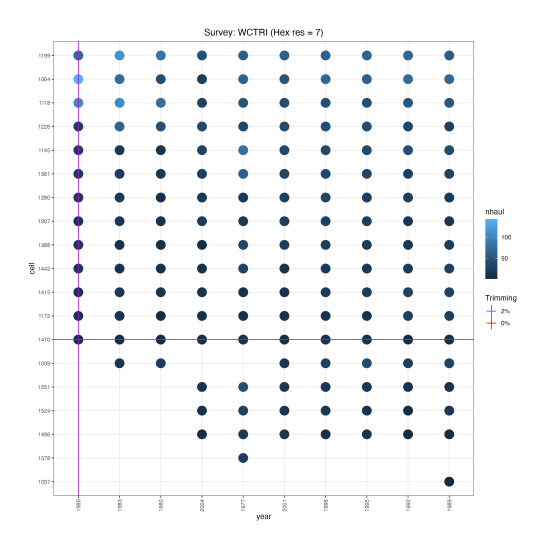
Total number of species	302.0
Percentage of species flagged	11.6

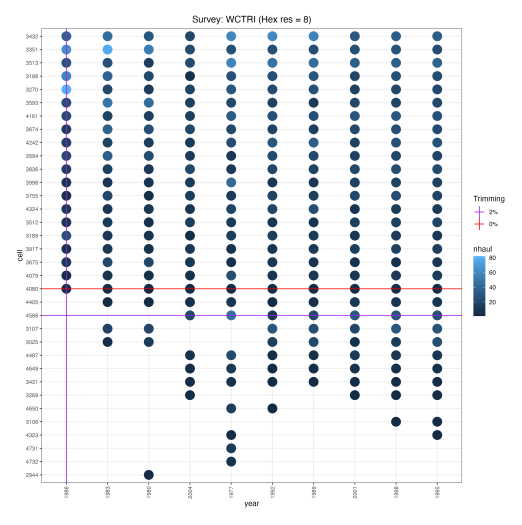
10. Spatio-temporal standardization

a. Standardization method 1

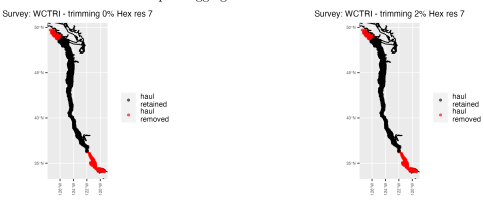
This standardization method was adapted from https://github.com/zoekitchel/trawl_spatial_turnover/blob/master/data_prep_code/species/explore_NorthSea_trimming.Rmd It was run for hex resolution 7 and 8.

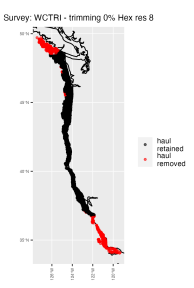
Plot of number of cells x years with overlaid flagging options

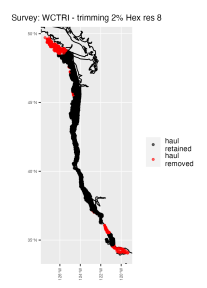




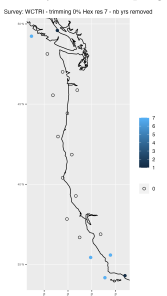
Map of hauls retained and removed per flagging method and threshold

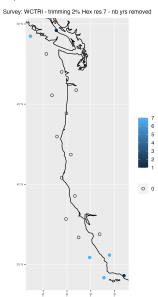


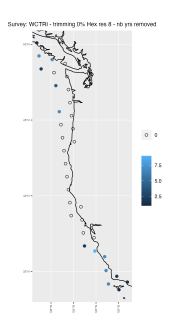


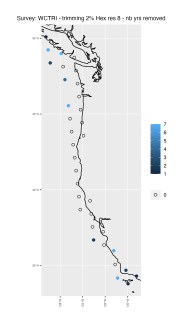


Map of numbers of years removed per grid cell and flagging method/threshold







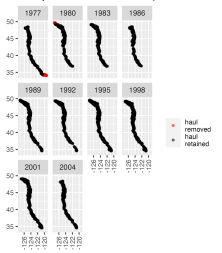


b. Standardization method 2

This standardization method was adapted from BioTIME code from https://github.com/Wubing-Xu/Range $_$ size $_$ winners $_$ losers

Map of hauls retained and removed

survey= WCTRI year1= 1977 year2= 1998 max.shared.samples= 401 duration= 22



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	490.0	490.0	651.0	426.0	383.0
percentage of hauls removed	10.8	10.8	14.3	9.4	0.5