Final Report

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Introduction

Animal pollinators, particularly bees, play a crucial role in the producing global food crops (Klein et al., 2007) and maintaining ecosystem health and biodiversity by facilitating plant reproduction (Biesmeijer et al., 2006). Understanding the dynamics of bee-plant interactions is essential for conservation efforts, agricultural productivity, sustainable development and ecosystem resilience (Patel et al., 2021, Huang et al., 2021).

In this study, we utilize the Oregon Bee Atlas (OBA) dataset which is a citizen science project in which trained volunteers catch bee species around Oregon which is then verified by experts before logging its information and preserving the specimen (Best et al., 2021, 2022). This dataset covers the state of Oregon in the Pacific Northwest of the US, spanning two years (2018-2019) and includes information on bee species, the plants they were foraging on, date caught, and sex. Combined with other open datasets, OBA can provide a valuable opportunity to explore various aspects of bee-plant interactions and their implications for ecosystem functioning.

Background and Research Questions

Part 1: Effect of Sampling Protocols and Urbanization on Bee Sex Ratio

Background: Sampling Protocol and Taxon level biases

Different types of sampling protocols used to collect bees in the OBA are aerial net, blue vane trap, pan trap and trap nest. The aerial net is considered active method while the rest are considered passive. Many studies in the literature have shown that taxon level biases exist between different sampling protocols, some of them are listed below:

- 1. Aerial Net involves using nets to capture insects or small animals while they are flying or moving through the air. Active methods, such as aerial netting, rely heavily on the researcher's observational abilities and netting skill. These techniques involve physically capturing or directly interacting with organisms in their natural environment. Success in these methods often depends on the researcher's ability to accurately observe and predict the behavior of the target organisms, as well as their skill in using the equipment effectively (Roulston et al 2007).
- 2. Blue Vane Traps are effective for capturing larger bees, particularly species like Bombus and Eucerine bees (Geroff et al., 2014; McCravy et al., 2017; Kimoto et al., 2012; Buchanan et al., 2017). These traps typically consist of a blue vane that attracts bees, leading them into a capturing mechanism.
- 3. Pan Traps are useful for catching small bees, especially Halictid bees, but they capture bigger bees less frequently (Droege et al., 2010; Richards et al., 2011; Toler et al., 2005). These traps involve shallow pans filled with soapy water, which attracts and drowns the bees.
- 4. **Trap Nests** are designed to catch cavity-building and more solitary bees, though only a small subset of potential cavity-nesting bee species actually use them (Campbell et al., 2017). These traps consist of tubes or cavities that mimic natural nesting sites for bees.

Background: Importance of Bee Sex Ratio

Bee Sex Ration is an important factor as it can influence population dynamics of bee communities. It also has strong effects on foraging behavior of the bees. Studies have shown that there are differences in floral preferences (which flowers the bees visit), floral constancy (how likely is a bee to visit the flower of the same species), pollen transfer efficiency (efficiency of pollen transfer between the flowers), and flight distance between foraging bouts among male and female bees (Ritchie et al., 2016; Ne'eman et al., 2006; Ostevik et al., 2010). These differences can have implications for the overall foraging efficiency and reproductive success of bee populations. Moreover, there is evidence to suggest that the sex ratio of bee populations can change with levels of urbanization, with urban areas showing a more male-biased sex ratio. This change is mainly driven by a decline in medium and large-bodied ground-nesting female bees in urban environments (Fitch et al., 2019). Understanding these sex-based differences and their impact on population dynamics is crucial for effective conservation strategies for bee species.

Research Questions

1A. Effect of Sampling Protocols Are different sampling protocols more likely to capture bees of one sex than another?

1B. Effect of Human population (marker for urbanization) How does human population levels affect bee distribution and sex ratio?

Part 2: Effect of Weather on Bee Populations

Background

Weather patterns play a crucial role in influencing the behavior and ecology of bees, both in their nursery and foraging activities. Studies have shown that increasing temperatures can enhance flight activity among bees, while decreasing temperatures have the opposite effect (Riessberger & Crailsheim, 1997; Burrill & Dietz, 1981). Moreover, temperature and precipitation have been identified as more important predictors of bee behavior than landscape composition, quality, or topography (Kammerer et al., 2021). Long-term precipitation patterns can also significantly impact bee communities by affecting the availability of floral resources (Thomson, 2016).

Despite the importance of weather patterns, there is a limited understanding of how short-term weather fluctuations affect wild bee populations in Oregon. Previous studies have primarily focused on long-term trends and the effects of climate change on bees. Understanding the short-term impacts of weather on bee populations is crucial for developing effective conservation strategies, especially in the face of climate change and increasing weather variability (Burrill & Dietz, 1981; Kammerer et al., 2021).

Research Questions

How does the number of foraging bees change with temperature and precipitation?

Additional Data Sources Used

- 1. Oregon County level Population from Oregon Demographics by Cubit (https://www.oregon-demographics.com)
- 2. Oregon Climate Data from National Oceanic and Atmospheric Administration (https://graphical.weather.gov/sectors/oregon.php)

Code

Import OBA datasets

```
getwd()
## [1] "/home/piyush/Dropbox/Academics/BI510L Data Sci in R/BI510L Data Sci in R_code/bi510l_project"
setwd("/home/piyush/Dropbox/Academics/BI510L Data Sci in R/BI510L Data Sci in R_code/bi510l_project")
oba_2018 <- read.csv("data/2018_occurrence.csv")
oba_2019 <- read.csv("data/2019_occurrence.csv")
cat("The colnames are identical:", identical(colnames(oba_2018), colnames(oba_2019)))
## The colnames are identical: TRUE
oba_2018_2019 <- rbind(oba_2018, oba_2019)
# head(oba_2018_2019, 3)</pre>
```

Clean dataset using regex

[10] ""

I will clean and update the associatedTaxa column which is supposed to have the plant name

1. Only keep the plant names in the string

```
sample(oba_2018_2019$associatedTaxa, size = 10)
   [1] "foraging on : \"Mentha\""
   [2] "foraging on : \"Sedum\""
   [3] "foraging on : \"Rhododendron macrophyllum\""
  [4] ""
  [5] ""
##
   [6] ""
##
   [7] "\"foraging on\" : \"Anaphalis margaritacea\""
   [8] ""
## [9] "foraging on : \"Erigeron canadensis\""
## [10] ""
# using regex to select for the pattern to keep rather than the pattern to drop
# noticed that the plant name is always bounded by the second double quotes ""
# Define the regex pattern
pattern <- '(.*)"(?<plant>.*)"$'
# Use str_match() to extract the match
match <- str_match(oba_2018_2019$associatedTaxa, pattern)
plant_name_list <- match[, "plant"]</pre>
plant_name_list[is.na(plant_name_list)] <- ""</pre>
oba_2018_2019$associatedTaxa <- plant_name_list
sample(oba_2018_2019$associatedTaxa, size = 10)
  [1] "Madia elegans"
##
                                "Eryngium maritimum"
   [4] "Prunus virginiana"
                                "Prunus emarginata"
                                                         "Cirsium vulgare"
## [7] ""
                                "Potentilla glandulosa" "Rubus laciniatus"
```

2. Remove all weed names and keep only one plant name

```
cat("We have these many weeds:", sum(grepl("Weedy", oba 2018 2019 associatedTaxa)), "\n")
## We have these many weeds: 3
oba_2018_2019$associatedTaxa <- gsub("Weedy yellow comp.", "", oba_2018_2019$associatedTaxa)
oba_2018_2019$associatedTaxa <- gsub("Weedy", "", oba_2018_2019$associatedTaxa)
cat("Now we have these many weeds:", sum(grepl("Weedy", oba_2018_2019$associatedTaxa)), "\n")
## Now we have these many weeds: 0
# get rid of;
cat("We have these many ';'", sum(grepl(";", oba_2018_2019$associatedTaxa)), "\n")
## We have these many ';' 11
oba_2018_2019$associatedTaxa[grepl(";", oba_2018_2019$associatedTaxa)] <- sapply(
    strsplit(
        oba_2018_2019$associatedTaxa[grepl(";", oba_2018_2019$associatedTaxa)], ";"
   ),
    function(x) x[1]
cat("We have these many ';'", sum(grepl(";", oba_2018_2019$associatedTaxa)), "\n")
## We have these many ';' 0
```

Add more information

1. Add plant resolution

We want to make a new column called plantResolution and assign it to "Family", "Genus" or "Species" depending on the level of resolution associated taxa is resolved to.

```
oba_2018_2019$plantTaxaWordCount <- str_count(oba_2018_2019$associatedTaxa, "\\w+")
head(oba_2018_2019$associatedTaxa, 10)
    [1] "Vaccinium ovatum" ""
                                                                     "Whipplea modesta"
                                                 11 11
    [5] "Whipplea modesta" ""
##
    [9] ""
                            "Horkelia fusca"
head(oba_2018_2019$plantTaxaWordCount, 10)
   [1] 2 0 0 2 2 0 0 0 0 2
summary(oba_2018_2019$plantTaxaWordCount)
##
      Min. 1st Qu.
                    Median
                               Mean 3rd Qu.
                                                Max.
             0.000
                      1.000
                              1.084
                                       2.000
                                               4.000
plant_resolution_list <- rep(NA, nrow(oba_2018_2019))</pre>
for (i in seq_along(plant_resolution_list)) {
    word_count <- oba_2018_2019$plantTaxaWordCount[i]</pre>
    plant_name <- oba_2018_2019$associatedTaxa[i]</pre>
    if (word_count == 1) { # can be genus or family
        if (grepl("aceae$", plant_name)) { # family
            plant resolution list[i] <- "family"</pre>
```

```
} else {
            plant_resolution_list[i] <- "genus"</pre>
    } else if (word_count > 1) { # it is species/subspecies
        plant_resolution_list[i] <- "species"</pre>
    }
}
oba_2018_2019$plantResolution <- plant_resolution_list
head(oba_2018_2019$associatedTaxa, 10)
   [1] "Vaccinium ovatum" ""
                                                                   "Whipplea modesta"
    [5] "Whipplea modesta" ""
   [9] ""
                            "Horkelia fusca"
head(oba_2018_2019$plantTaxaWordCount, 10)
## [1] 2 0 0 2 2 0 0 0 0 2
head(oba_2018_2019$plantResolution, 10)
  [1] "species" NA
                                       "species" "species" NA
                                                                      NA
## [8] NA
                             "species"
```

2. Add plant genus

[8] NA

Create a new column called plantGenus that is the genus if the associated taxa was resolved to species or genus, and NA if it was resolved to family.

```
plant_genus_list <- rep(NA, nrow(oba_2018_2019))</pre>
for (i in seq_along(plant_genus_list)) {
    plant_res <- oba_2018_2019$plantResolution[i]</pre>
    plant_name <- oba_2018_2019$associatedTaxa[i]</pre>
    if (!is.na(plant_res) && (plant_res == "genus")) {
        plant_genus_list[i] <- plant_name</pre>
    } else if (!is.na(plant_res) && (plant_res == "species")) {
        plant_genus_list[i] <- strsplit(plant_name, " ")[[1]][1]</pre>
    }
}
oba_2018_2019$plantGenus <- plant_genus_list
head(oba_2018_2019$associatedTaxa, 10)
    [1] "Vaccinium ovatum" ""
                                                                     "Whipplea modesta"
                                                 11 11
##
    [5] "Whipplea modesta" ""
   [9] ""
                            "Horkelia fusca"
head(oba_2018_2019$plantTaxaWordCount, 10)
## [1] 2 0 0 2 2 0 0 0 0 2
head(oba_2018_2019$plantResolution, 10)
                                        "species" "species" NA
  [1] "species" NA
                                                                        NA
                             NA
```

"species"

```
head(oba_2018_2019$plantGenus, 10)

## [1] "Vaccinium" NA NA "Whipplea" "Whipplea" NA

## [7] NA NA NA "Horkelia"
```

Save/Read the new dataframe as csv

Drop the columns that we are not going to use

```
col_to_save <- c(</pre>
    "gbifID", "genericName", "specificEpithet", "species", "sex",
    "taxonRank", "eventDate", "year", "month", "samplingProtocol",
    "associatedTaxa", "county", "decimalLatitude", "decimalLongitude",
    "plantResolution", "plantGenus"
oba_2018_2019 <- oba_2018_2019[, col_to_save]
colnames(oba_2018_2019)
##
  [1] "gbifID"
                           "genericName"
                                              "specificEpithet" "species"
## [5] "sex"
                           "taxonRank"
                                              "eventDate"
                                                                  "year"
## [9] "month"
                                                                  "county"
                           "samplingProtocol" "associatedTaxa"
## [13] "decimalLatitude" "decimalLongitude" "plantResolution" "plantGenus"
Save
write.csv(oba_2018_2019, "data/cleaned_oba_2018_2019.csv", row.names = FALSE)
```

Read

```
oba_2018_2019 <- read.csv("data/cleaned_oba_2018_2019.csv")

# check if all values of column gbifID are unique
unique_gbifID <- length(unique(oba_2018_2019$gbifID))
total_gbifID <- nrow(oba_2018_2019)

if (unique_gbifID == total_gbifID) {
    print("All values of column 'gbifID' are unique.")
} else {
    print("Not all values of column 'gbifID' are unique.")
}</pre>
```

[1] "All values of column 'gbifID' are unique."

This means we can use gbifID as a unique identifier

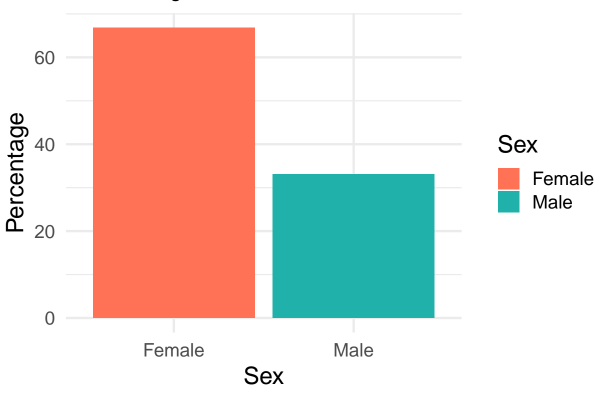
1. Bee Sex Ratio

A. Effect of Sampling Protocols

```
# check the TOTAL PERCENTAGE of MALE vs FEMALE bees caught
total_male <- sum(oba_2018_2019$sex == "MALE", na.rm = TRUE)
total_female <- sum(oba_2018_2019$sex == "FEMALE", na.rm = TRUE)
total_bees <- nrow(oba_2018_2019)</pre>
```

```
percentage_male <- (total_male / total_bees) * 100</pre>
percentage_female <- (total_female / total_bees) * 100</pre>
# Create a data frame for the plot
sex_per_df <- data.frame(</pre>
    Sex = c("Female", "Male"),
    Percentage = c(percentage_female, percentage_male)
# Create a bar plot with ggplot
p_title <- "Percentage of Male and Female Bees"</pre>
p <- ggplot(sex_per_df, aes(x = Sex, y = Percentage, fill = Sex)) +</pre>
    geom_bar(stat = "identity") +
    scale_fill_manual(values = c("coral1", "lightseagreen")) +
    labs(x = "Sex", y = "Percentage") +
    ggtitle(p_title) +
    theme_minimal(18) +
    theme(
        axis.text = element_text(size = 14),
        plot.title = element_text(hjust = 0.5, size = 16)
p
```

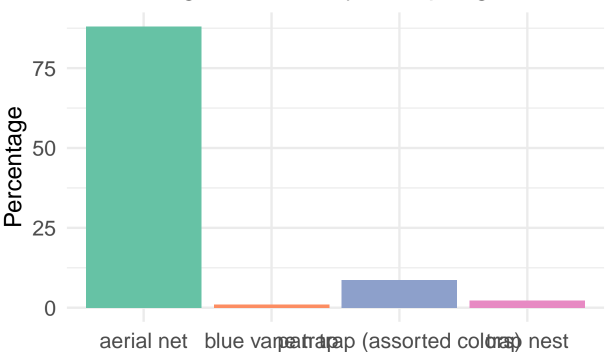
Percentage of Male and Female Bees



ggsave(glue("plots/{p_title}.png"), dpi = 300, plot = p)

```
# Calculate the percentage of bees caught using different sampling patterns
sampling_patterns <- unique(oba_2018_2019$samplingProtocol)</pre>
percentage_per_pattern <- sapply(sampling_patterns, function(pattern) {</pre>
    total_bees <- sum(oba_2018_2019$samplingProtocol == pattern, na.rm = TRUE)
    percentage <- (total_bees / nrow(oba_2018_2019)) * 100</pre>
    return(percentage)
})
sampling_per_df <- data.frame(</pre>
    patterns = sampling_patterns,
    percentage = percentage_per_pattern
# Create a stacked bar plot with percentages
p_title <- "Percentage of Bees by Sampling Protocol"</pre>
p <- ggplot(sampling_per_df, aes(x = patterns, y = percentage, fill = patterns)) +
    geom_bar(stat = "identity", position = "stack", show.legend = F) +
    labs(x = "Sex", y = "Percentage", fill = "Sampling Protocol") +
    ggtitle(p_title) +
    theme_minimal(18) +
    theme(
        axis.text = element_text(size = 16),
        plot.title = element_text(hjust = 0.5)
    scale_fill_brewer(palette = "Set2")
р
```

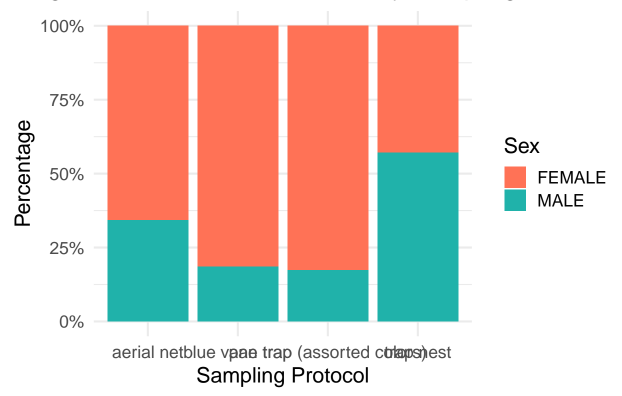
Percentage of Bees by Sampling Protocol



Sex

```
\# ggsave(glue("plots/{p_title}).png"), dpi = 300, plot = p)
p_title <- "Percentage of Male and Female Bees by Sampling Protocol"
p <- oba_2018_2019 %>%
    filter(sex != "") %>%
    group_by(sex, samplingProtocol) %>%
    summarise(n_group = n()) %>%
    ungroup() %>%
    group_by(samplingProtocol) %>%
    mutate(
       n_sampling = sum(n_group),
       percentage = n_group / n_sampling
    ggplot(aes(x = samplingProtocol, y = percentage, fill = sex)) +
    geom_bar(stat = "identity", position = "stack") +
    labs(x = "Sampling Protocol", y = "Percentage", fill = "Sex") +
    ggtitle(p_title) +
    scale_fill_manual(values = c("coral1", "lightseagreen")) +
    scale_y_continuous(labels = scales::percent_format()) +
    theme_minimal(16) +
    theme(
       axis.text = element_text(size = 13),
       plot.title = element_text(hjust = 0.5)
## `summarise()` has grouped output by 'sex'. You can override using the `.groups`
## argument.
p
```

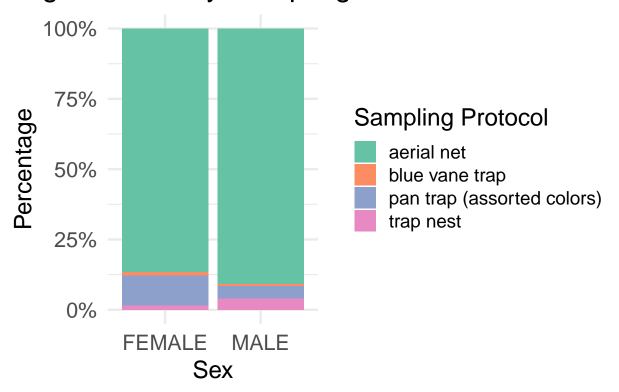
entage of Male and Female Bees by Sampling Protocol



```
# ggsave(glue("plots/{p_title}.png"), dpi = 300, plot = p)
# Generate the plot
p_title <- "Percentage of Bees by Sampling Protocol_2"</pre>
p <- oba_2018_2019 %>%
   filter(sex != "") %>%
    group_by(sex, samplingProtocol) %>%
    summarise(n_group = n()) %>%
   ungroup() %>%
    group_by(sex) %>%
   mutate(
        n_sampling = sum(n_group),
        percentage = n_group / n_sampling
    ggplot(aes(x = sex, y = percentage, fill = samplingProtocol)) +
   geom_bar(stat = "identity", position = "stack") +
   labs(x = "Sex", y = "Percentage", fill = "Sampling Protocol") +
   ggtitle(p title) +
   scale_fill_brewer(palette = "Set2") +
    scale_y_continuous(labels = scales::percent_format()) +
   theme_minimal(18) +
    theme(
        axis.text = element_text(size = 16),
        plot.title = element_text(hjust = 0.5)
```

```
## `summarise()` has grouped output by 'sex'. You can override using the `.groups`
## argument.
p
```

ntage of Bees by Sampling Protocol_2



```
# ggsave(glue("plots/{p_title}.png"), dpi = 300, plot = p)
```

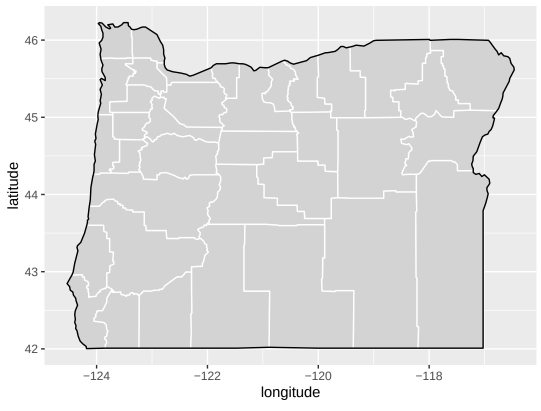
B. Effect of Human Population

Download basemaps into R using the map_data function in ggplot

```
# get Oregon map
or_map_base_df <- map_data("state") %>%
 filter(region == "oregon") %>%
 select(longitude = long, latitude = lat, group_id = group, state_name = region)
str(or_map_base_df)
## 'data.frame':
                   236 obs. of 4 variables:
## $ longitude : num -117 -117 -117 -117 ...
   $ latitude : num 46 46 45.9 45.9 45.9 ...
   $ group_id : num 44 44 44 44 44 44 44 44 44 ...
  $ state_name: chr "oregon" "oregon" "oregon" "oregon" ...
or_map_county_df <- map_data("county", "oregon") %>%
 select(longitude = long, latitude = lat, group_id = group, county_name = subregion)
str(or_map_county_df)
## 'data.frame':
                   1633 obs. of 4 variables:
## $ longitude : num -117 -117 -118 -118 ...
```

```
: num 44.3 44.3 44.4 44.4 44.4 ...
## $ group_id : num 1 1 1 1 1 1 1 1 1 1 ...
## $ county_name: chr "baker" "baker" "baker" "baker" ...
# plot Oregon map with bee location overlaid
plot_or_w_county <- ggplot() +</pre>
  geom_polygon(
    data = or_map_county_df, fill = "lightgray", color = "white",
    aes(x = longitude, y = latitude, group = county_name)
  ) +
  geom_polygon(
    data = or_map_base_df, fill = NA, color = "black",
    aes(x = longitude, y = latitude, group = state_name)
  ) +
  theme(plot.title = element_text(hjust = 0.5))
plot_or_w_county +
  ggtitle(glue("Map of Oregon with County Boundaries")) +
  coord_quickmap()
```

Map of Oregon with County Boundaries



```
# data taken from https://www.oregon-demographics.com/counties_by_population

# make a data frame
or_dem <- readxl::read_xlsx("./data/oregon_demographics_by_county.xlsx", skip = 4, col_names = TRUE)
head(or_dem)

## # A tibble: 6 x 67

## name state county city most_current_pop_year population pop_dens_sq_mi mhhi</pre>
```

```
## 2 Bake~ Oreg~ NA
                                                2022
                                                           16685
                                                                             5.4 51657
## 3 Bent~ Oreg~ NA
                                                2022
                                                                                 72882
                         NΑ
                                                           95615
                                                                           142.
## 4 Clac~ Oreg~ NA
                                                 2022
                                                          420925
                                                                           225
                                                                                 95740
                                                2022
## 5 Clat~ Oreg~ NA
                                                           41190
                                                                            NA
                                                                                    ΝA
## 6 Colu~ Oreg~ NA
                         NA
                                                           52865
                                                                                    NΑ
## # i 59 more variables: hhi_total_hh <dbl>, hhi_hh_w_lt_25k <dbl>,
       \label{lem:lem:hhi_hh_w_25k_49k <dbl>, hhi_hh_w_50k_74k <dbl>, hhi_hh_w_75k_99k <dbl>,
## #
       hhi_hh_w_100k_149k <dbl>, hhi_hh_w_150k_199k <dbl>,
       hhi_hh_w_200k_plus <dbl>, race_ethnicity_total <dbl>,
## #
       race_ethnicity_white <dbl>, race_ethnicity_black <dbl>,
## #
       race_ethnicity_native <dbl>, race_ethnicity_asian <dbl>,
       race_ethnicity_islander <dbl>, race_ethnicity_other <dbl>, ...
colnames (or_dem)
##
   [1] "name"
                                              "state"
##
    [3] "county"
                                             "city"
##
   [5] "most_current_pop_year"
                                              "population"
  [7] "pop_dens_sq_mi"
## [9] "hhi_total_hh"
                                              \verb|"hhi_hh_w_lt_25k||
## [11] "hhi_hh_w_25k_49k"
                                              "hhi_hh_w_50k_74k"
## [13] "hhi_hh_w_75k_99k"
                                             "hhi_hh_w_100k_149k"
## [15] "hhi_hh_w_150k_199k"
                                             "hhi_hh_w_200k_plus"
                                             "race_ethnicity_white"
## [17] "race_ethnicity_total"
## [19] "race_ethnicity_black"
                                             "race_ethnicity_native"
## [21] "race_ethnicity_asian"
                                             "race_ethnicity_islander"
## [23] "race_ethnicity_other"
                                             "race_ethnicity_two"
                                              "age_total"
## [25] "race_ethnicity_hispanic"
## [27] "age_f_0_9"
                                             "age_f_10_19"
## [29] "age_f_20_29"
                                             "age_f_30_39"
## [31] "age_f_40_49"
                                             "age_f_50_59"
## [33] "age_f_60_69"
                                              "age_f_70_plus"
## [35] "age_m_0_9"
                                             "age_m_10_19"
                                             "age_m_30_39"
## [37] "age_m_20_29"
## [39] "age_m_40_49"
                                             "age_m_50_59"
## [41] "age_m_60_69"
                                              "age_m_70_plus"
## [43] "median_age"
                                             "edu_att_pop_25_plus"
## [45] "edu_att_no_diploma"
                                             "edu_att_high_school"
## [47] "edu_att_some_college"
                                             "edu_att_bachelors"
## [49] "edu_att_graduate"
                                             "family_hh_total"
## [51] "family_poverty_pct"
                                             "emp_status_civ_labor_force"
## [53] "unemployment_pct"
                                             "housing_units"
## [55] "occupied_units"
                                              "owner_occupied"
## [57] "renter_occupied"
                                              "median_value_owner_occupied_units"
## [59] "households"
                                             "hh families"
## [61] "hh_mc_families"
                                             "hh_mc_with_own_children_under_18"
## [63] "hh_sp_families"
                                             "hh_sp_with_own_children_under_18"
## [65] "hh_non_families"
                                             "aland_sq_m"
## [67] "geoid"
# select only two columns, remove oregon total population, add 'county' col, convert everything to lowe
or_pop_by_county_df <- or_dem %>%
  select(name, population) %>%
```

<dbl>

2022

<dbl>

4229374

<dbl> <dbl> 44.1 76632

<chr> <chr> <lgl>

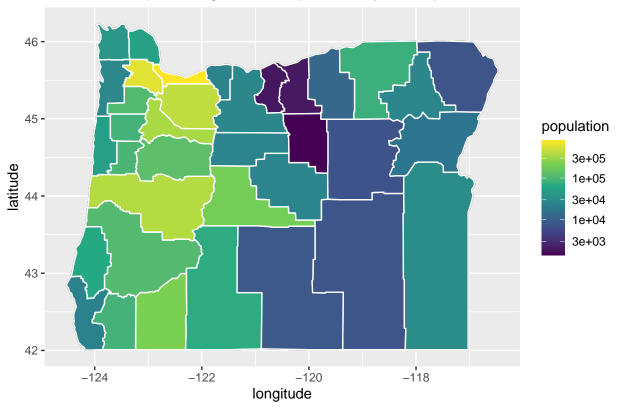
1 Oreg~ Oreg~ NA

<lgl>

NA

```
filter(name != "Oregon") %>%
  mutate(county_name = tolower(str_replace(name, " County$", ""))) %>%
  select(-name)
head(or_pop_by_county_df)
## # A tibble: 6 x 2
    population county_name
          <dbl> <chr>
          16685 baker
## 1
## 2
          95615 benton
## 3
       420925 clackamas
## 4
         41190 clatsop
## 5
         52865 columbia
## 6
         64908 coos
head(or_map_county_df)
     longitude latitude group_id county_name
## 1 -117.2042 44.30683
                               1
                                       baker
## 2 -117.4907 44.30683
                               1
                                       baker
## 3 -117.4907 44.38704
                               1
                                       baker
## 4 -117.5366 44.42142
                                       baker
                               1
## 5 -117.5709 44.42142
                               1
                                       baker
## 6 -117.5996 44.43861
                               1
                                       baker
# now the county_name in or_pop_by_county_df corresponds to or_map_county_df, join them in a single df
or_map_with_population_df <- left_join(or_map_county_df, or_pop_by_county_df, by = "county_name")
head(or_map_with_population_df)
     longitude latitude group_id county_name population
## 1 -117.2042 44.30683
                              1
                                       baker
                                                  16685
## 2 -117.4907 44.30683
                                       baker
                                                  16685
## 3 -117.4907 44.38704
                                                  16685
                                       baker
                               1
## 4 -117.5366 44.42142
                               1
                                       baker
                                                  16685
## 5 -117.5709 44.42142
                               1
                                       baker
                                                  16685
## 6 -117.5996 44.43861
                                       baker
                                                  16685
plot_or_w_county +
  ggtitle(glue("Map of Oregon with Population by County")) +
  geom_polygon(data = or_map_with_population_df,
  aes(x = longitude, y = latitude, fill = population, group = county_name), color = "white") +
  scale_fill_continuous(trans = "log10", type = "viridis")
```

Map of Oregon with Population by County

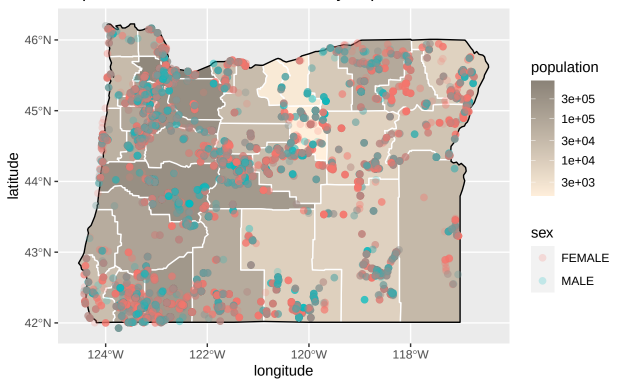


coord_sf(datum = sf::st_crs(4326))

```
<ggproto object: Class CoordSf, CoordCartesian, Coord, gg>
##
       aspect: function
       backtransform_range: function
##
       clip: on
##
       crs: NULL
##
##
       datum: crs
       default: FALSE
##
##
       default_crs: NULL
##
       determine_crs: function
       distance: function
##
       expand: TRUE
##
##
       fixup_graticule_labels: function
       get_default_crs: function
##
##
       is_free: function
##
       is_linear: function
##
       label_axes: list
##
       label_graticule:
##
       labels: function
##
       limits: list
##
       lims_method: cross
       modify_scales: function
##
##
       ndiscr: 100
##
       params: list
##
       range: function
```

```
##
       record bbox: function
##
       render_axis_h: function
##
       render axis v: function
##
       render_bg: function
##
       render_fg: function
##
       setup data: function
##
       setup layout: function
##
       setup_panel_guides: function
##
       setup_panel_params: function
##
       setup_params: function
##
       train_panel_guides: function
##
       transform: function
       super: <ggproto object: Class CoordSf, CoordCartesian, Coord, gg>
summary(oba_2018_2019$decimalLatitude)
##
      Min. 1st Qu. Median
                              Mean 3rd Qu.
                                               Max.
                                                       NA's
     41.93
             43.17
                     44.60
                             44.23
                                     45.17
                                              46.99
                                                         12
summary(oba_2018_2019$decimalLongitude)
      Min. 1st Qu. Median
                                                       NA's
                              Mean 3rd Qu.
                                               Max.
## -124.42 -123.23 -122.89 -121.97 -121.27
                                              43.37
                                                         12
all_species_data <- oba_2018_2019 %>%
  filter((decimalLongitude < -116) & (decimalLatitude < 46.5) & (sex != "")) %>%
  select(x = decimalLongitude, y = decimalLatitude, sex)
all_species_spatial <- st_as_sf(all_species_data, coords = c("x", "y"), crs = "WGS84")
p title <- "Map of All Bee Locations with County Population info"
p <- ggplot() +</pre>
  geom_polygon(
   data = or_map_county_df, fill = "lightgray", color = "white",
    aes(x = longitude, y = latitude, group = county_name)
  ) +
  geom_polygon(
   data = or_map_with_population_df,
   aes(x = longitude, y = latitude, fill = population, group = county_name), color = "white"
  ) +
  scale_fill_gradient(trans = "log10", low = "antiquewhite1", high = "antiquewhite4") +
  geom_polygon(
   data = or_map_base_df, fill = NA, color = "black",
   aes(x = longitude, y = latitude, group = state_name)
  ) +
  # qeom sf(data = all species spatial, color = "darkorange", alpha = 0.1, size = 2) +
  geom_sf(data = all_species_spatial, aes(color = sex), alpha = 0.2, size = 2) +
  # geom_segment(aes(x = -118.4, y = 42.3, xend = -117.21, yend = 42.3), linewidth = 4) +
  # annotate("text", x = -117.8, y = 42.43, label = "100 km", size = 4) +
  # theme(plot.title = element_text(hjust = 0.5)) +
  ggtitle(p_title)
p
```

Map of All Bee Locations with County Population info



```
\# ggsave(glue("plots/{p_title}_by_sex.png"), dpi = 300, plot = p)
```

2. Weather and Foraging Patterns

Lane county

```
# get climate data for lane county
# Read csv files
avg_temp <- read.csv("data/lane_county_avg_temp_2018_2019.csv", skip = 3, header = 4)
avg_ppt <- read.csv("data/lane_county_precipitation_2018_2019.csv", skip = 3, header = 4)</pre>
# rename columns
colnames(avg_temp)[colnames(avg_temp) == "Date"] <- "date"</pre>
colnames(avg_temp)[colnames(avg_temp) == "Value"] <- "temp_f"</pre>
colnames(avg_ppt) [colnames(avg_ppt) == "Date"] <- "date"</pre>
colnames(avg_ppt) [colnames(avg_ppt) == "Value"] <- "ppt_inch"</pre>
# add column 'year' and 'month' from the first four string character from the 'Date' column
avg_temp$year <- substr(avg_temp$date, 1, 4)</pre>
avg_temp$month <- substr(avg_temp$date, 5, 6)</pre>
avg_temp$year <- as.numeric(avg_temp$year)</pre>
avg_temp$month <- as.numeric(avg_temp$month)</pre>
avg_temp$temp_c <- (avg_temp$temp_f - 32) * (5 / 9) # add celsius</pre>
head(avg_temp)
```

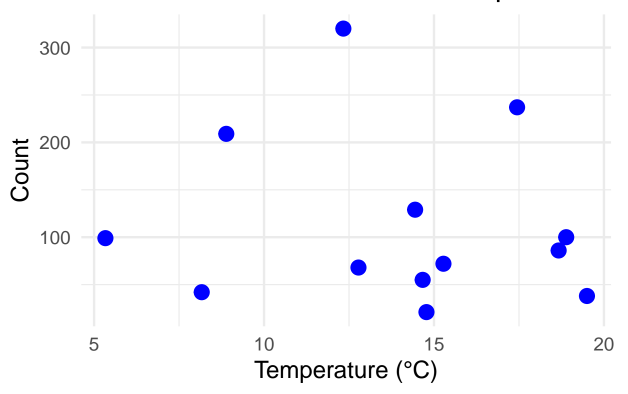
```
date temp_f year month
                             temp c
## 1 201801 42.0 2018 1 5.555556
## 2 201802 38.2 2018
                         2 3.444444
## 3 201803 41.3 2018
                        3 5.166667
                       4 8.166667
## 4 201804 46.7 2018
                       5 12.777778
## 5 201805 55.0 2018
## 6 201806 58.4 2018
                       6 14.666667
avg ppt$year <- substr(avg ppt$date, 1, 4)</pre>
avg_ppt$month <- substr(avg_ppt$date, 5, 6)</pre>
avg_ppt$year <- as.numeric(avg_ppt$year)</pre>
avg_ppt$month <- as.numeric(avg_ppt$month)</pre>
avg_ppt$ppt_cm <- avg_ppt$ppt_inch * 2.54 # add cm ppt</pre>
head(avg ppt)
##
      date ppt_inch year month ppt_cm
## 1 201801
             8.41 2018
                         1 21.3614
## 2 201802
              4.73 2018
                           2 12.0142
## 3 201803
              5.84 2018
                         3 14.8336
## 4 201804
             6.55 2018
                        4 16.6370
## 5 201805
             0.83 2018
                         5 2.1082
6 4.7244
lane_oba_2018_2019 <- oba_2018_2019 %>%
   filter(county == "Lane")
# find the count of total number of 'gbifID' by year and month in lane_oba_2018_2019
net_count_per_month <- lane_oba_2018_2019 %>%
   group_by(year, month) %>%
   summarise(count = n distinct(gbifID))
## `summarise()` has grouped output by 'year'. You can override using the
## `.groups` argument.
species_count_per_month <- lane_oba_2018_2019 %>%
   filter(species != "") %>%
   group_by(year, month) %>%
   summarise(count = n_distinct(species))
## `summarise()` has grouped output by 'year'. You can override using the
## `.groups` argument.
# qet and add ppt_cm information from avq_ppt df by year and month
net_count_per_month <- net_count_per_month %>%
   left_join(avg_ppt, by = c("year", "month")) %>%
   left_join(avg_temp, by = c("year", "month"))
species_count_per_month <- species_count_per_month %>%
   left_join(avg_ppt, by = c("year", "month")) %>%
   left_join(avg_temp, by = c("year", "month"))
head(net_count_per_month)
## # A tibble: 6 x 9
## # Groups:
             year [1]
     <dbl> <dbl> <int> <int> <dbl>
                                     <dbl> <int> <dbl> <dbl>
##
```

```
## 1 2018
                 42 201804
                             6.55 16.6
                                        201804
                                                46.7
                                                      8.17
## 2 2018
                 68 201805
                             0.83 2.11
            5
                                        201805
                                                55
                                                     12.8
                                                58.4 14.7
## 3 2018
            6 55 201806
                             1.86 4.72
                                        201806
## 4 2018
            7
                 38 201807
                             0.01 0.0254 201807
                                                67.1 19.5
                             0.13 0.330 201808
                                                65.6 18.7
## 5 2018
            8
                 86 201808
## 6 2018
            9
                 21 201809
                             0.41 1.04
                                        201809
                                                58.6 14.8
head(species_count_per_month)
## # A tibble: 6 x 9
## # Groups:
            year [1]
##
     <dbl> <dbl> <int> <int>
                            <dbl>
                                  <dbl> <int> <dbl> <dbl>
## 1 2018
            4
                 8 201804
                             6.55 16.6
                                        201804
                                                46.7
                                                      8.17
## 2 2018
            5
                 14 201805
                             0.83 2.11
                                                     12.8
                                        201805
                                                55
## 3 2018
            6
                 15 201806
                             1.86 4.72
                                        201806
                                                58.4 14.7
## 4 2018
            7
                 8 201807
                             0.01 0.0254 201807
                                                67.1 19.5
                                                65.6 18.7
## 5 2018
            8
                 21 201808
                             0.13 0.330 201808
## 6 2018
            9
                 10 201809
                             0.41 1.04
                                        201809
                                                58.6 14.8
```

Temperature effect

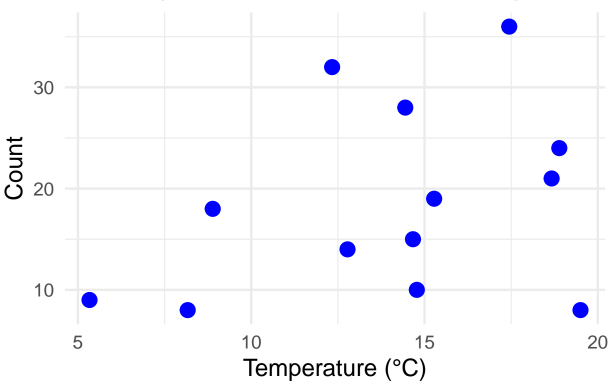
```
# Scatter plot
ggplot(net_count_per_month) +
    geom_point(aes(x = temp_c, y = count), size = 5, color = "blue") +
    labs(x = "Temperature (°C)", y = "Count") +
    ggtitle("Net Count vs Mean Temp") +
    theme_minimal(18) +
    theme(plot.title = element_text(hjust = 0.5))
```

Net Count vs Mean Temp



```
ggplot(species_count_per_month) +
   geom_point(aes(x = temp_c, y = count), size = 5, color = "blue") +
   labs(x = "Temperature (°C)", y = "Count") +
   ggtitle("Species Count vs Mean Temp") +
   theme_minimal(18) +
   theme(plot.title = element_text(hjust = 0.5))
```

Species Count vs Mean Temp



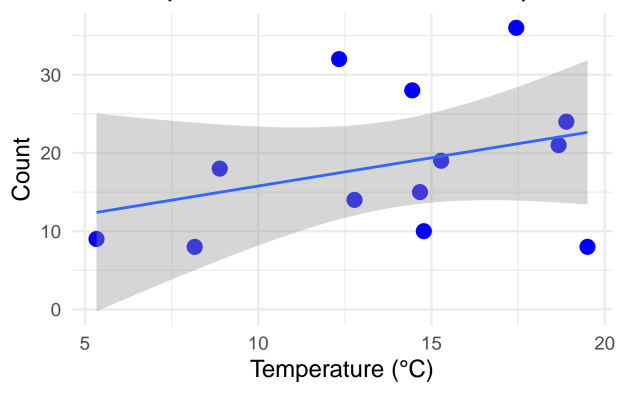
```
lin_reg <- lm(species_count_per_month$count ~ species_count_per_month$temp_c)</pre>
lin_reg_slope <- summary(lin_reg)$coefficients[2]</pre>
lin_reg_pval <- summary(lin_reg)$coefficients[7]</pre>
summary(lin_reg)
##
## Call:
## lm(formula = species_count_per_month$count ~ species_count_per_month$temp_c)
##
## Residuals:
##
       Min
                1Q Median
                                 3Q
                                        Max
## -14.634 -4.143 -1.032
                              3.030 14.851
##
## Coefficients:
##
                                   Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                     8.5498
                                                8.7324
                                                          0.979
                                                                   0.349
                                     0.7223
## species_count_per_month$temp_c
                                                0.5999
                                                          1.204
                                                                   0.254
## Residual standard error: 9.1 on 11 degrees of freedom
## Multiple R-squared: 0.1164, Adjusted R-squared: 0.03613
## F-statistic: 1.45 on 1 and 11 DF, p-value: 0.2538
print(lin_reg_slope)
## [1] 0.7222794
print(lin_reg_pval)
```

[1] 0.3485819

```
ggplot(species_count_per_month, aes(x = temp_c, y = count)) +
    geom_point(size = 5, color = "blue") +
    labs(x = "Temperature (°C)", y = "Count") +
    ggtitle("Species Count vs Mean Temp") +
    theme_minimal(18) +
    geom_smooth(method = "lm", se = TRUE) +
    theme(plot.title = element_text(hjust = 0.5))
```

`geom_smooth()` using formula = 'y ~ x'

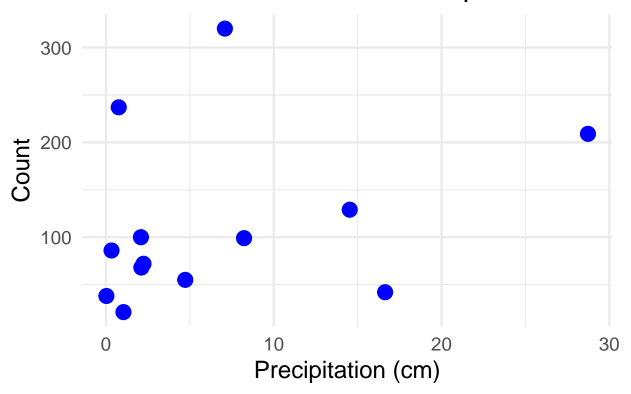
Species Count vs Mean Temp



Precipitation effect

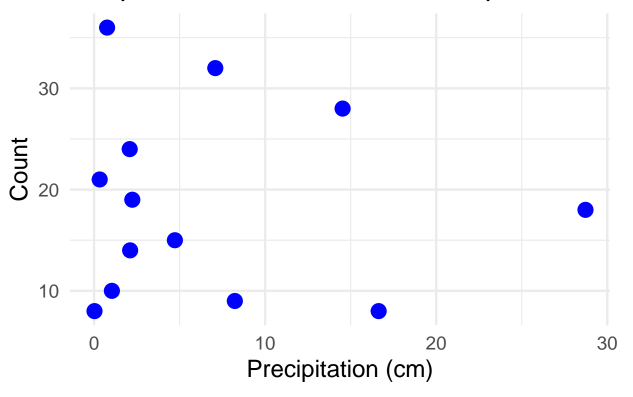
```
# Scatter plot
ggplot(net_count_per_month) +
    geom_point(aes(x = ppt_cm, y = count), size = 5, color = "blue") +
    labs(x = "Precipitation (cm)", y = "Count") +
    ggtitle("Net Count vs Mean Precipitation") +
    theme_minimal(18)+
    theme(plot.title = element_text(hjust = 0.5))
```

Net Count vs Mean Precipitation



```
ggplot(species_count_per_month) +
   geom_point(aes(x = ppt_cm, y = count), size = 5, color = "blue") +
   labs(x = "Precipitation (cm)", y = "Count") +
   ggtitle("Species Count vs Mean Precipitation") +
   theme_minimal(18) +
   theme(plot.title = element_text(hjust = 0.5))
```

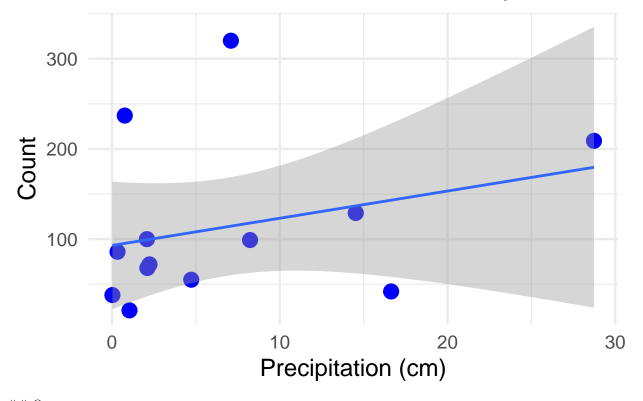
Species Count vs Mean Precipitation



```
lin_reg <- lm(net_count_per_month$count ~ species_count_per_month$ppt_cm)</pre>
lin_reg_slope <- summary(lin_reg)$coefficients[2]</pre>
lin_reg_pval <- summary(lin_reg)$coefficients[7]</pre>
summary(lin_reg)
##
## Call:
## lm(formula = net_count_per_month$count ~ species_count_per_month$ppt_cm)
##
## Residuals:
       Min
                  1Q
                       Median
                                    3Q
## -101.189 -52.249 -18.824
                                 0.721 205.624
## Coefficients:
##
                                  Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                    92.995
                                                32.139
                                                         2.893
                                                                 0.0146 *
## species_count_per_month$ppt_cm
                                     3.017
                                                 3.023
                                                         0.998
                                                                 0.3398
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 88.99 on 11 degrees of freedom
## Multiple R-squared: 0.08301,
                                  Adjusted R-squared: -0.0003575
## F-statistic: 0.9957 on 1 and 11 DF, p-value: 0.3398
print(lin_reg_slope)
```

`geom_smooth()` using formula = 'y ~ x'

Net Count vs Mean Temp



```
## Oregon

# get climate data for Oregon

# Read csv files
avg_temp <- read.csv("data/oregon_avg_temp_2018_2019.csv", skip = 3, header = 4)
avg_ppt <- read.csv("data/oregon_precipitation_2018_2019.csv", skip = 3, header = 4)

# rename columns
colnames(avg_temp)[colnames(avg_temp) == "Date"] <- "date"
colnames(avg_temp)[colnames(avg_temp) == "Value"] <- "temp_f"
colnames(avg_ppt)[colnames(avg_ppt) == "Date"] <- "date"
colnames(avg_ppt)[colnames(avg_ppt) == "Date"] <- "date"
colnames(avg_ppt)[colnames(avg_ppt) == "Value"] <- "ppt_inch"

# add column 'year' and 'month' from the first four string character from the 'Date' column</pre>
```

```
avg_temp$year <- substr(avg_temp$date, 1, 4)</pre>
avg_temp$month <- substr(avg_temp$date, 5, 6)</pre>
avg_temp$year <- as.numeric(avg_temp$year)</pre>
avg_temp$month <- as.numeric(avg_temp$month)</pre>
avg_temp$temp_c <- (avg_temp$temp_f - 32) * (5 / 9) # add celsius</pre>
head(avg_temp)
##
      date temp_f year month
                                temp_c
## 1 201801 37.5 2018 1 3.0555556
## 2 201802 33.2 2018
                         2 0.6666667
## 4 201804 45.0 2018 4 7.2222222
## 5 201805 55.4 2018 5 13.0000000
                        6 14.7222222
## 6 201806 58.5 2018
avg_ppt$year <- substr(avg_ppt$date, 1, 4)</pre>
avg_ppt$month <- substr(avg_ppt$date, 5, 6)</pre>
avg ppt$year <- as.numeric(avg ppt$year)</pre>
avg_ppt$month <- as.numeric(avg_ppt$month)</pre>
avg_ppt$ppt_cm <- avg_ppt$ppt_inch * 2.54 # add cm ppt</pre>
head(avg_ppt)
##
      date ppt_inch year month ppt_cm
## 1 201801
              4.14 2018
                         1 10.5156
## 2 201802
              2.29 2018 2 5.8166
## 3 201803
             3.42 2018 3 8.6868
## 4 201804
              3.26 2018 4 8.2804
             1.27 2018
                           5 3.2258
## 5 201805
## 6 201806
              1.13 2018 6 2.8702
# find the count of total number of 'gbifID' by year and month in oba_2018_2019
net_count_per_month <- oba_2018_2019 %>%
   group_by(year, month) %>%
   summarise(count = n distinct(gbifID))
## `summarise()` has grouped output by 'year'. You can override using the
## `.groups` argument.
species_count_per_month <- oba_2018_2019 %>%
   filter(species != "") %>%
   group_by(year, month) %>%
   summarise(count = n_distinct(species))
## `summarise()` has grouped output by 'year'. You can override using the
## `.groups` argument.
# qet and add ppt_cm information from avq_ppt df by year and month
net_count_per_month <- net_count_per_month %>%
   left_join(avg_ppt, by = c("year", "month")) %>%
   left_join(avg_temp, by = c("year", "month"))
species_count_per_month <- species_count_per_month %>%
   left join(avg ppt, by = c("year", "month")) %>%
   left_join(avg_temp, by = c("year", "month"))
head(net_count_per_month)
```

```
## # A tibble: 6 x 9
## # Groups:
               year [1]
      year month count date.x ppt_inch ppt_cm date.y temp_f temp_c
                                         <dbl>
                                                <int>
                                                        <dbl>
##
     <dbl> <dbl> <int>
                        <int>
                                  <dbl>
                                                               <dbl>
## 1 2017
               3
                     1
                            NA
                                     NA
                                            NA
                                                           NA
                                                                  NA
## 2
     2017
               5
                            NA
                                     NA
                                            NA
                                                    NA
                                                           MΔ
                                                                  NA
                    31
## 3
     2017
               6
                    14
                            NA
                                     NA
                                            NΑ
                                                    NΑ
                                                                  NΑ
## 4
     2017
               7
                    70
                            NA
                                     NA
                                            NA
                                                    NA
                                                           NA
                                                                  NA
## 5
      2017
               8
                    70
                            NA
                                     NA
                                            NA
                                                    NA
                                                           NA
                                                                  NA
## 6 2017
               9
                                     NA
                    43
                            NA
                                            NA
                                                    NA
                                                           NA
                                                                  NA
head(species_count_per_month)
## # A tibble: 6 x 9
## # Groups:
               year [1]
##
      year month count date.x ppt_inch ppt_cm date.y temp_f temp_c
     <dbl> <dbl> <int>
                        <int>
                                  <dbl>
                                         <dbl>
                                                <int>
                                                        <dbl>
## 1 2017
               5
                     4
                            NA
                                     NA
                                            NA
                                                    NA
                                                           NA
                                                                  NA
## 2
     2017
               6
                     7
                            NA
                                     NA
                                            NA
                                                    NA
                                                           NA
                                                                  NA
               7
## 3 2017
                    15
                            NA
                                     NA
                                            NA
                                                    NA
                                                           NA
                                                                  NA
     2017
               8
                    20
                            NA
                                     NA
                                            NA
                                                    NA
                                                           NA
                                                                  NA
## 5
     2017
               9
                    12
                            NA
                                     NA
                                            NA
                                                    NA
                                                           NA
                                                                  NA
## 6 2017
                      2
                            NΑ
                                     NA
                                            NA
                                                                  NΑ
Temperature Effect
# Scatter plot
lin_reg <- lm(net_count_per_month$count ~ net_count_per_month$temp_c)</pre>
# lin_reg_slope <- summary(lin_reg)$coefficients[2]</pre>
# lin_reg_pval <- summary(lin_reg)$coefficients[7]</pre>
summary(lin_reg)
##
## lm(formula = net_count_per_month$count ~ net_count_per_month$temp_c)
##
## Residuals:
       Min
##
                1Q Median
                                 3Q
                                        Max
## -1516.5 -975.3 -422.3
                              278.2 3616.4
##
## Coefficients:
##
                               Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                -264.18
                                            724.20 -0.365 0.71977
                                             58.28
                                                     3.364 0.00368 **
## net_count_per_month$temp_c
                                 196.07
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1528 on 17 degrees of freedom
     (8 observations deleted due to missingness)
## Multiple R-squared: 0.3997, Adjusted R-squared: 0.3644
## F-statistic: 11.32 on 1 and 17 DF, p-value: 0.003681
p_title = "Oregon - Net Count vs Mean Monthly Temperature"
p <- ggplot(net_count_per_month, aes(x = temp_c, y = count)) +</pre>
```

geom_point(size = 5, color = "#ff0000") +

```
labs(x = "Temperature (°C)", y = "Total #Bees") +
    ggtitle(p_title) +
    theme_minimal(18) +
    geom_smooth(method = "lm", se = TRUE, color="#ff0000") +
    theme(plot.title = element_text(hjust = 0.5))

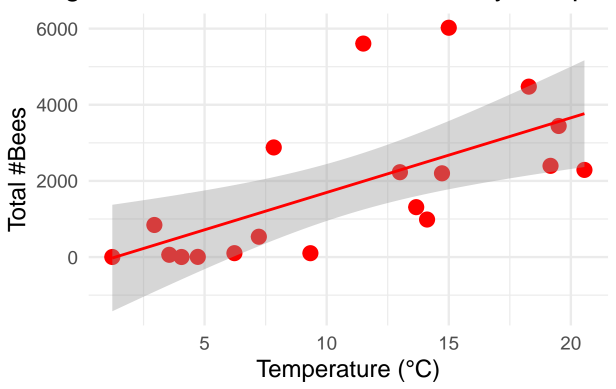
p

## `geom_smooth()` using formula = 'y ~ x'

## Warning: Removed 8 rows containing non-finite values (`stat_smooth()`).

## Warning: Removed 8 rows containing missing values (`geom_point()`).
```

Oregon - Net Count vs Mean Monthly Tempera



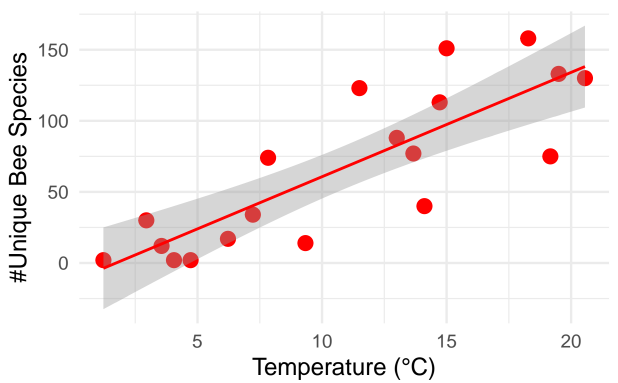
```
# ggsave(glue("plots/{p_title}.png"), dpi = 300, plot = p)

lin_reg <- lm(species_count_per_month$count ~ species_count_per_month$temp_c)
summary(lin_reg)

## Call:
## lm(formula = species_count_per_month$count ~ species_count_per_month$temp_c)
##
## Residuals:
## Min 1Q Median 3Q Max
## -52.952 -15.521 -1.403 19.370 53.621
###</pre>
```

```
## Coefficients:
##
                                  Estimate Std. Error t value Pr(>|t|)
                                               14.883
## (Intercept)
                                   -12.687
                                                      -0.852
                                     7.338
## species_count_per_month$temp_c
                                                1.198
                                                        6.126 1.12e-05 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 31.4 on 17 degrees of freedom
     (7 observations deleted due to missingness)
## Multiple R-squared: 0.6883, Adjusted R-squared: 0.6699
## F-statistic: 37.53 on 1 and 17 DF, p-value: 1.119e-05
p_title <- "Oregon - Species Count vs Mean Monthly Temperature"</pre>
p <- ggplot(species_count_per_month, aes(x = temp_c, y = count)) +</pre>
    geom_point(size = 5, color = "#ff0000") +
    labs(x = "Temperature (°C)", y = "#Unique Bee Species") +
    ggtitle(p_title) +
    theme_minimal(18) +
    geom_smooth(method = "lm", se = TRUE, color = "#ff0000") +
    theme(plot.title = element_text(hjust = 0.5))
p
## 'geom_smooth()' using formula = 'y ~ x'
## Warning: Removed 7 rows containing non-finite values (`stat_smooth()`).
## Warning: Removed 7 rows containing missing values (`geom_point()`).
```

Pregon – Species Count vs Mean Monthly Tempe

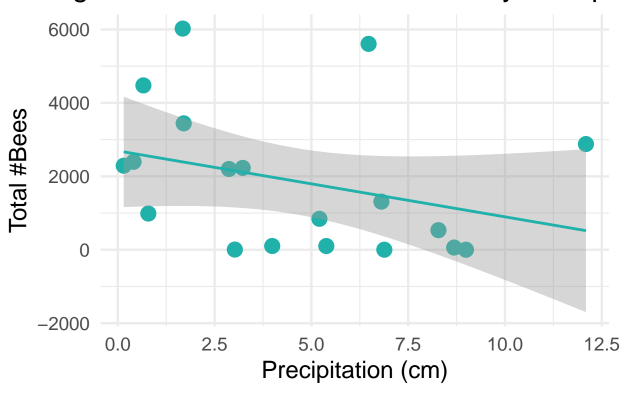


```
# ggsave(glue("plots/{p_title}.png"), dpi = 300, plot = p)
```

Precipitation effect

```
lin_reg <- lm(net_count_per_month$count ~ net_count_per_month$ppt_cm)</pre>
# lin_reg_slope <- summary(lin_reg)$coefficients[2]</pre>
# lin_reg_pval <- summary(lin_reg)$coefficients[7]</pre>
summary(lin_reg)
##
## Call:
## lm(formula = net_count_per_month$count ~ net_count_per_month$ppt_cm)
##
## Residuals:
##
       Min
                1Q Median
                                3Q
                                       Max
## -2145.4 -1265.1 -377.3 585.1 4077.4
##
## Coefficients:
##
                              Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                2693.7
                                            727.9
                                                   3.700 0.00178 **
                                            128.1 -1.403 0.17865
## net_count_per_month$ppt_cm
                                -179.7
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1867 on 17 degrees of freedom
     (8 observations deleted due to missingness)
## Multiple R-squared: 0.1038, Adjusted R-squared: 0.05104
## F-statistic: 1.968 on 1 and 17 DF, p-value: 0.1786
p_title <- "Oregon - Net Count vs Mean Monthly Precipitation"</pre>
p <- ggplot(net_count_per_month, aes(x = ppt_cm, y = count)) +</pre>
    geom_point(size = 5, color = "lightseagreen") +
   labs(x = "Precipitation (cm)", y = "Total #Bees") +
   ggtitle(p_title) +
   theme minimal(18) +
    geom_smooth(method = "lm", se = TRUE, color = "lightseagreen") +
    theme(plot.title = element text(hjust = 0.5))
p
## `geom_smooth()` using formula = 'y ~ x'
## Warning: Removed 8 rows containing non-finite values (`stat_smooth()`).
## Warning: Removed 8 rows containing missing values (`geom_point()`).
```

Oregon - Net Count vs Mean Monthly Precipit



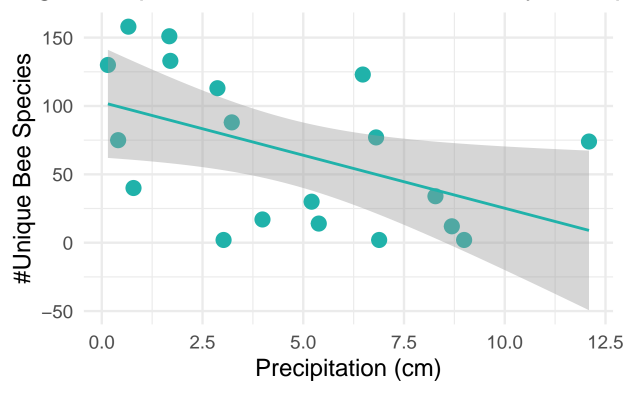
```
# ggsave(glue("plots/{p_title}.png"), dpi = 300, plot = p)
lin_reg <- lm(species_count_per_month$count ~ species_count_per_month$ppt_cm)</pre>
summary(lin_reg)
##
## Call:
## lm(formula = species_count_per_month$count ~ species_count_per_month$ppt_cm)
##
## Residuals:
                1Q Median
                                3Q
  -77.292 -39.670 -4.536 37.999 70.485
##
## Coefficients:
##
                                  Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                   102.721
                                                19.151
                                                         5.364 5.16e-05 ***
## species_count_per_month$ppt_cm
                                    -7.751
                                                3.370 -2.300
                                                                 0.0344 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 49.11 on 17 degrees of freedom
     (7 observations deleted due to missingness)
## Multiple R-squared: 0.2373, Adjusted R-squared: 0.1924
## F-statistic: 5.29 on 1 and 17 DF, p-value: 0.03439
p_title <- "Oregon - Species Count vs Mean Monthly Precipitation"</pre>
p <- ggplot(species_count_per_month, aes(x = ppt_cm, y = count)) +</pre>
```

```
geom_point(size = 5, color = "lightseagreen") +
labs(x = "Precipitation (cm)", y = "#Unique Bee Species") +
ggtitle(p_title) +
theme_minimal(18) +
geom_smooth(method = "lm", se = TRUE, color = "lightseagreen") +
theme(plot.title = element_text(hjust = 0.5))
p
## `geom_smooth()` using formula = 'y ~ x'
```

Warning: Removed 7 rows containing non-finite values (`stat_smooth()`).

Warning: Removed 7 rows containing missing values (`geom_point()`).

Dregon – Species Count vs Mean Monthly Precip



ggsave(glue("plots/{p_title}.png"), dpi = 300, plot = p)

Results

- 1. A. Effect of Sampling Protocols: Are different sampling protocols more likely to capture bees of one sex than another?
- Yes! Trap nests capture a higher percentage of male bees. For example, in a colony, the male-to-female ratio is typically 1:100, while for solitary bees, it's closer to 1:1 (Willmer, 2011).
- 1. B. Effect of Human Population (Marker for Urbanization): How does the human population level affect bee distribution and sex ratio?
- There is a higher occurrence of bees near population centers. However, this observation comes with a

- caveat, as it can be influenced by sampling bias. My studies have not shown a clear difference in the sex ratio of bees in urbanized areas compared to other habitats.
- 2. Effect of Local Weather on Bee Populations: How does the number of foraging bees change with temperature and precipitation?
- In my study, I found that the number of unique bee species significantly increases with increasing average monthly temperature. Conversely, the number of foraging bees tends to significantly decrease with increasing average monthly precipitation. These findings highlight the intricate relationship between local weather patterns and bee populations, emphasizing the importance of considering climatic factors in bee conservation efforts.