Bird Diversity Analysis

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Summary

This project studies landscape and bird diversity change in a mountainous area (the Willmore Wilderness Park in Alberta, Canada) over the last century.

We have data on past and current land cover composition (historic and repeat photos & their land cover classifications). The 10 land cover categories are: Coniferous forest, Broadleaf forest, Mixedwood forest, Wetland, Shrub, Herbaceous, Rock, Water, Regenerating area, Snow/Ice.

We have species distribution models (made from auditory survey data and Landsat-based land cover map). The 15 bird species are: American pipit, American robin, Chipping sparrow, Dark-eyed junco, Golden-crowned kinglet, Golden-crowned sparrow, Gray jay, Hermit thrush, Pine siskin, Ruby-crowned kinglet, Savannah sparrow, Swainson's thrush, Varied thrush, Wilson's warbler, Yellow-rumped warbler.

```
# Load packages
library(here)
library(readxl)
library(tidyverse)
library(vegan)
```

Load data

- Raw data consists of Mountain Legacy Project (MLP) photos (Photographs.zip)
- Pre-processed data consists of image classifications (LandCoverClassifications.zip)
- Processed data is in ImageAnalysis.xlsx
 - Each tab is an MLP station (i.e. photograph pair)
 - Classified photos were uploaded to the MLP's Image Analysis Toolkit
 - Land cover % outputs from IAT were copied back into ImageAnalysis.xlsx
 - Model backtransformations are done to get probability of occurrence in both historical and repeat photos for each station
 - Then there is a tab for land cover summary across all photograph pairs
 - And a tab aggregating outputs for all historical images
 - And a tab aggregating outputs for all repeat images

```
historicdata <- read_xlsx(here("ProcessedData","ImageAnalysis.xlsx"), sheet="Historic") %>%
   mutate(TIME = "Historic")

repeatdata <- read_xlsx(here("ProcessedData","ImageAnalysis.xlsx"), sheet="Repeat") %>%
   mutate(TIME = "Repeat")

data <- rbind(historicdata, repeatdata) %>%
   pivot_longer(cols=CF:SN, names_to="LANDCOVER", values_to="LANDCOVER_PERCENT") %>%
   pivot_longer(cols=GRAJ:VATH, names_to="SPECIES", values_to="SPECIES_OCCURRENCE") %>%
   mutate(across(c(STATION, TIME, LANDCOVER, SPECIES), as.factor))

landcovertypes <- read.csv(here("ProcessedData", "LandcoverLookup.csv"))
```

```
species <- read.csv(here("ProcessedData", "SpeciesLookup.csv")) %>%
mutate(NAMESEP = gsub(" ", " \n ", NAME)) # for plotting later
```

Calculate diversity

Calculate diversity indices for both land cover and species.

```
data <- data %>%
  group_by(STATION, TIME, SPECIES) %>%
  mutate(LANDCOVER_DIV = diversity(LANDCOVER_PERCENT, index="shannon")) %>%
  ungroup() %>%
  group_by(STATION, TIME, LANDCOVER) %>%
  mutate(SPECIES_DIV = diversity(SPECIES_OCCURRENCE, index="shannon")) %>%
  ungroup()
```

Examine land cover change

Run paired t-tests for each land cover type and for land cover diversity. We chose paired t-tests because the samples (photographs) are not independent - it's the same landscape at two different times.

T-tests are relatively robust to deviations from assumptions, but assumptions are:

- Continuous data
- Random samples

Paired t-test

- Homogeneity of variance
- Distribution approximately normal

```
# Prep data
landcoverdata <- data %>%
  select(STATION, TIME, LANDCOVER, LANDCOVER_PERCENT) %>%
  unique()
# Print t-test results for each land cover type
for (lc in landcovertypes$CODE) {
  print(landcovertypes$NAME[which(landcovertypes$CODE==lc)])
  print(t.test(landcoverdata %>% filter(LANDCOVER==1c, TIME=="Historic") %>% pull(LANDCOVER_PERCENT),
         landcoverdata %>% filter(LANDCOVER==lc, TIME=="Repeat") %>% pull(LANDCOVER_PERCENT),
         paired=T))
}
## [1] "Coniferous Forest"
##
##
  Paired t-test
##
## data: landcoverdata %>% filter(LANDCOVER == lc, TIME == "Historic") %>% pull(LANDCOVER_PERCENT) and
## t = -4.5633, df = 45, p-value = 3.876e-05
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -17.913748 -6.942774
## sample estimates:
## mean difference
##
         -12.42826
##
## [1] "Broadleaf Forest"
##
```

```
##
## data: landcoverdata %>% filter(LANDCOVER == 1c, TIME == "Historic") %>% pull(LANDCOVER_PERCENT) and
## t = 0.47923, df = 45, p-value = 0.6341
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.05570012 0.09048273
## sample estimates:
## mean difference
##
         0.0173913
##
## [1] "Mixedwood Forest"
##
## Paired t-test
##
## data: landcoverdata %>% filter(LANDCOVER == lc, TIME == "Historic") %>% pull(LANDCOVER_PERCENT) and
## t = -0.8365, df = 45, p-value = 0.4073
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -2.896609 1.196609
## sample estimates:
## mean difference
             -0.85
##
## [1] "Wetlands"
##
## Paired t-test
##
## data: landcoverdata %>% filter(LANDCOVER == lc, TIME == "Historic") %>% pull(LANDCOVER_PERCENT) and
## t = 2.8607, df = 45, p-value = 0.006389
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.1955858 1.1261533
## sample estimates:
## mean difference
##
         0.6608696
##
## [1] "Shrub Cover"
##
## Paired t-test
##
## data: landcoverdata %>% filter(LANDCOVER == 1c, TIME == "Historic") %>% pull(LANDCOVER_PERCENT) and
## t = 1.706, df = 45, p-value = 0.0949
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.2716922 3.2803879
## sample estimates:
## mean difference
##
          1.504348
## [1] "Herbaceous Cover"
##
## Paired t-test
##
## data: landcoverdata %>% filter(LANDCOVER == 1c, TIME == "Historic") %>% pull(LANDCOVER_PERCENT) and
```

```
## t = 6.083, df = 45, p-value = 2.351e-07
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 4.337647 8.631918
## sample estimates:
## mean difference
          6.484783
##
##
## [1] "Rock"
##
## Paired t-test
##
## data: landcoverdata %>% filter(LANDCOVER == lc, TIME == "Historic") %>% pull(LANDCOVER_PERCENT) and
## t = 1.8056, df = 45, p-value = 0.07767
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.08457856 1.54979595
## sample estimates:
## mean difference
##
         0.7326087
##
## [1] "Water"
##
## Paired t-test
##
## data: landcoverdata %>% filter(LANDCOVER == 1c, TIME == "Historic") %>% pull(LANDCOVER_PERCENT) and
## t = 3.6465, df = 45, p-value = 0.000687
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.04379322 0.15185896
## sample estimates:
## mean difference
##
        0.09782609
##
## [1] "Regenerating Area"
##
## Paired t-test
##
## data: landcoverdata %>% filter(LANDCOVER == lc, TIME == "Historic") %>% pull(LANDCOVER_PERCENT) and
## t = 1.2113, df = 45, p-value = 0.2321
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -1.994084 8.011475
## sample estimates:
## mean difference
          3.008696
##
##
## [1] "Snow/Ice"
##
## Paired t-test
##
## data: landcoverdata %>% filter(LANDCOVER == 1c, TIME == "Historic") %>% pull(LANDCOVER_PERCENT) and
## t = 1.996, df = 45, p-value = 0.052
## alternative hypothesis: true mean difference is not equal to 0
```

```
## 95 percent confidence interval:
## -0.006732943 1.493689464
## sample estimates:
## mean difference
        0.7434783
# Overall landscape diversity
print(t.test(data %>% select(STATION, TIME, LANDCOVER_DIV) %>% unique() %>% filter(TIME=="Historic") %>
             data %>% select(STATION, TIME, LANDCOVER_DIV) %>% unique() %>% filter(TIME=="Repeat") %>%
             paired=T))
##
##
   Paired t-test
## data: data %>% select(STATION, TIME, LANDCOVER_DIV) %>% unique() %>% filter(TIME == "Historic") %>%
## t = 5.1271, df = 45, p-value = 6.018e-06
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.1228375 0.2817877
## sample estimates:
## mean difference
##
         0.2023126
# Historical landscape diversity
data %>% select(STATION, TIME, LANDCOVER_DIV) %>% unique() %>% filter(TIME=="Historic") %>% pull(LANDCO
## [1] 1.199805
data %>% select(STATION, TIME, LANDCOVER_DIV) %>% unique() %>% filter(TIME=="Historic") %>% pull(LANDCO
## [1] 0.03935819
# Repeat landscape diversity
data %>% select(STATION, TIME, LANDCOVER_DIV) %>% unique() %>% filter(TIME=="Repeat") %>% pull(LANDCOVE
## [1] 0.9974924
data %>% select(STATION, TIME, LANDCOVER_DIV) %>% unique() %>% filter(TIME=="Repeat") %>% pull(LANDCOVE
## [1] 0.04447595
```

Given these outputs, we can see which land cover types changed. We can also be confident that the landscape diversity declined between the historical photos and the repeat photos.

Examine bird diversity change

Run paired t-tests for each bird species and for bird diversity. We chose paired t-tests because the samples (photographs) are not independent - it's the same landscape at two different times.

```
## [1] "Gray Jay"
##
##
  Paired t-test
##
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and bird
## t = 3.8312, df = 45, p-value = 0.0003928
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.02374718 0.07639195
## sample estimates:
## mean difference
##
       0.05006956
##
## [1] "Wilson's Warbler"
##
## Paired t-test
##
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and bird
## t = 0.40219, df = 45, p-value = 0.6894
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.01739597 0.02607694
## sample estimates:
## mean difference
      0.004340486
##
## [1] "Savannah Sparrow"
##
## Paired t-test
##
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and bird
## t = -3.514, df = 45, p-value = 0.001019
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.18669617 -0.05065448
## sample estimates:
## mean difference
##
        -0.1186753
##
## [1] "Golden-crowned Kinglet"
## Paired t-test
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and bird
## t = 4.5144, df = 45, p-value = 4.543e-05
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.01242143 0.03243330
## sample estimates:
## mean difference
##
        0.02242736
##
## [1] "Ruby-crowned Kinglet"
##
```

```
## Paired t-test
##
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and bird
## t = 3.0555, df = 45, p-value = 0.00377
\#\# alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.004442516 0.021626548
## sample estimates:
## mean difference
##
                  0.01303453
## [1] "Dark-eyed Junco"
## Paired t-test
##
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and bird
## t = 5.8957, df = 45, p-value = 4.459e-07
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.01882403 0.03835902
## sample estimates:
## mean difference
                  0.02859153
##
##
## [1] "American Robin"
## Paired t-test
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and birddiscrete birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and birddiscrete 
## t = -3.907, df = 45, p-value = 0.0003112
\mbox{\tt \#\#} alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.04997135 -0.01597497
## sample estimates:
## mean difference
                -0.03297316
##
##
## [1] "Hermit Thrush"
##
## Paired t-test
##
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and bird
## t = 2.8944, df = 45, p-value = 0.00584
\mbox{\tt \#\#} alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.007576746 0.042247916
## sample estimates:
## mean difference
##
                  0.02491233
##
## [1] "Pine Siskin"
##
## Paired t-test
##
```

```
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and bird
## t = 4.1807, df = 45, p-value = 0.0001323
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.02617540 0.07484153
## sample estimates:
## mean difference
##
       0.05050846
##
## [1] "American Pipit"
##
  Paired t-test
##
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and bird
## t = -2.0487, df = 45, p-value = 0.04635
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.0741438345 -0.0006321223
## sample estimates:
## mean difference
##
      -0.03738798
##
## [1] "Golden-crowned Sparrow"
##
## Paired t-test
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and bird
## t = -1.9743, df = 45, p-value = 0.05451
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.0692928547 0.0006916874
## sample estimates:
## mean difference
       -0.03430058
##
## [1] "Swainson's Thrush"
##
## Paired t-test
##
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and bird
## t = 0.46603, df = 45, p-value = 0.6434
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.008776938 0.014061380
## sample estimates:
## mean difference
##
      0.002642221
##
## [1] "Yellow-rumped Warbler"
##
## Paired t-test
##
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and bird
## t = 6.7136, df = 45, p-value = 2.713e-08
```

```
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.1167426 0.2168094
## sample estimates:
## mean difference
##
                      0.166776
## [1] "Chipping Sparrow"
##
##
      Paired t-test
##
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and birddiscrete birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and birddiscrete 
## t = -2.8055, df = 45, p-value = 0.007394
\mbox{\tt \#\#} alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.054107230 -0.008884972
## sample estimates:
## mean difference
                  -0.0314961
##
##
## [1] "Varied Thrush"
##
     Paired t-test
##
## data: birddata %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE) and bird
## t = -0.8566, df = 45, p-value = 0.3962
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.03186980 0.01285023
## sample estimates:
## mean difference
##
             -0.009509786
# Overall bird diversity
print(t.test(data %>% select(STATION, TIME, SPECIES_DIV) %>% unique() %>% filter(TIME=="Historic") %>% ;
                             data %>% select(STATION, TIME, SPECIES_DIV) %>% unique() %>% filter(TIME=="Repeat") %>% pu
                             paired=T))
##
##
      Paired t-test
##
## data: data %>% select(STATION, TIME, SPECIES_DIV) %>% unique() %>% filter(TIME == "Historic") %>% p
## t = -0.1605, df = 45, p-value = 0.8732
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.006930295 0.005907297
## sample estimates:
## mean difference
             -0.000511499
At the species level, many species were affected and changed in probability of occurrence between the two
```

At the species level, many species were affected and changed in probability of occurrence between the two time points. However, we cannot say that the species diversity changed significantly between the two time periods.

Plot bird diversity changes

Create a data frame with the t-test outputs to make it easier to plot.

Link to external data on each species:

- Breeding habitat
- Non-breeding habitat
- Conservation concern

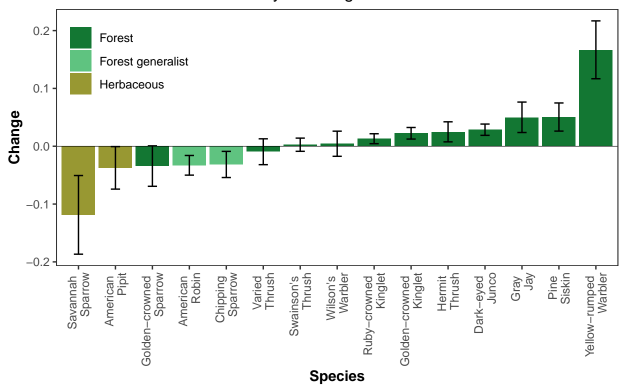
Source: State of North America's Birds 2016 Report

```
sonab <- read_xlsx(here("ProcessedData", "assessment-table-full.xlsx"), sheet="SONAB database for distri"</pre>
  filter(`English Name` %in% species$NAME) %>%
  select(`English Name`, `Breeding Habitats`, `Non-breeding Habitats`, CCSmax) %>%
  rename(NAME = `English Name`)
# Regroup habitats based on classes that match the categories we used in our analysis
sonab <- sonab %>%
  rename(BREEDING = `Breeding Habitats`,
         NONBREEDING = `Non-breeding Habitats`,
         CONSERVATION = CCSmax) %>%
  mutate(BREEDING = gsub("Arctic tundra", "", BREEDING), # remove classes that don't exist in our study
         BREEDING = gsub("Temperate western forest", "", BREEDING),
         BREEDING = gsub("Temperate eastern forest", "", BREEDING),
         BREEDING = gsub("Mexican highland forest", "", BREEDING),
         BREEDING = gsub("Mexican pine and oak forest", "", BREEDING),
         BREEDING = gsub("Temperate grassland", "", BREEDING),
         BREEDING = gsub("Agricultural", "", BREEDING),
         BREEDING = gsub("Alpine tundra", "Herbaceous", BREEDING), # rename to match our class names
         BREEDING = gsub("Boreal forest", "Forest", BREEDING),
         BREEDING = gsub(";", "", BREEDING),
         BREEDING = trimws(BREEDING)) %>%
  mutate(NONBREEDING = gsub("Beach and estuary", "", NONBREEDING), # remove classes that don't exist in
         NONBREEDING = gsub("Agricultural", "", NONBREEDING),
         NONBREEDING = gsub("Coastal saltmarsh", "", NONBREEDING),
         NONBREEDING = gsub("Tropical highland forest", "", NONBREEDING),
         NONBREEDING = gsub("Temperate grassland", "Herbaceous", NONBREEDING), # rename to match our cl
         NONBREEDING = gsub("Temperate western forest", "Forest", NONBREEDING),
         NONBREEDING = gsub("Tropical evergreen forest", "Herbaceous", NONBREEDING),
         NONBREEDING = gsub("Boreal forest", "Forest", NONBREEDING),
         NONBREEDING = gsub(";", "", NONBREEDING),
```

NONBREEDING = trimws(NONBREEDING)) %>%

```
mutate(CONSERVATION = replace(CONSERVATION, which(CONSERVATION %in% as.character(5:8)), "Low"), # cut
         CONSERVATION = replace(CONSERVATION, which(CONSERVATION %in% as.character(9:13)), "Moderate"))
  left_join(species, by="NAME")
df <- left_join(df, sonab, by="CODE")</pre>
df$ESTIMATE <- as.numeric(df$ESTIMATE)</pre>
df$CI LOW <- as.numeric(df$CI LOW)</pre>
df$CI HIGH <- as.numeric(df$CI HIGH)</pre>
df$NAME <- factor(df$NAME)</pre>
df$BREEDING <- factor(df$BREEDING)</pre>
df$NONBREEDING <- factor(df$NONBREEDING)</pre>
df$CONSERVATION <- factor(df$CONSERVATION)</pre>
df$NAMESEP <- factor(df$NAMESEP)</pre>
speciesorder <- levels(reorder(df$NAMESEP, df$ESTIMATE)) # we want to use this specific order for all p
df$NAMESEP <- factor(df$NAMESEP, speciesorder)</pre>
Make plots used in manuscript.
plot1 <- ggplot(data=df, aes(x=NAMESEP, y=ESTIMATE, fill=BREEDING)) +</pre>
  theme_bw() +
  geom_bar(stat="identity",position=position_dodge(0.9)) +
  geom_errorbar(aes(ymax=CI_HIGH, ymin=CI_LOW) ,position=position_dodge(0.9),width=0.25) +
  scale_fill_manual("Legend", values = c("Forest" = rgb(19/256,119/256,51/256),
                                           "Herbaceous" = rgb(153/256, 153/256, 50/256),
                                           "Forest generalist" = rgb(95/256,196/256,128/256))) +
  labs(x="Species",y="Change") +
  ggtitle("Change in species occurrence \nby breeding habitat") +
  theme(plot.title = element_text(hjust = 0.5),
        axis.text = element_text(),
        axis.text.x = element_text(angle=90, vjust=0.5, hjust=1),
        axis.title = element_text(face="bold"),
                   = element text(),
        \# panel.background = element_rect(fill="transparent", colour=NA), \# transparent background for
        # plot.background = element_rect(fill="transparent", colour=NA),
        panel.grid.minor = element_blank(),
        panel.grid.major = element_blank(),
        legend.title = element_blank(),
        legend.position = c(0.01, 0.99),
        legend.background = element_blank(),
        legend.justification = c("left","top"))+
  geom_hline(yintercept = 0, linewidth=0.2)
plot1
```

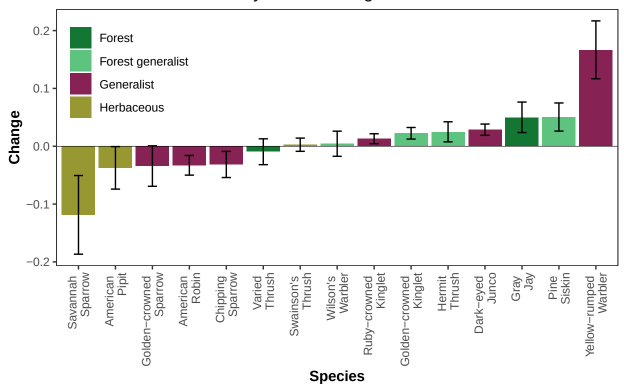
Change in species occurrence by breeding habitat



As we can see from this plot, species whose main breeding habitat is not forest have declined more than species that breed in forest.

```
plot2 <- ggplot(data=df, aes(x=NAMESEP, y=ESTIMATE, fill=NONBREEDING)) +</pre>
  theme_bw() +
  geom_bar(stat="identity", position=position_dodge(0.9)) +
  geom_errorbar(aes(ymax=CI_HIGH, ymin=CI_LOW) ,position=position_dodge(0.9),width=0.25) +
  scale_fill_manual("Legend", values = c("Forest" = rgb(19/256,119/256,51/256),
                                          "Herbaceous" = rgb(153/256, 153/256, 50/256),
                                          "Forest generalist" = rgb(95/256,196/256,128/256),
                                          "Generalist" = rgb(136/256,35/256,84/256))) +
  labs(x="Species",y="Change") +
  ggtitle("Change in species occurrence \nby non-breeding habitat") +
  theme(plot.title = element_text(hjust = 0.5),
        axis.text = element_text(),
        axis.text.x = element_text(angle=90, vjust=0.5, hjust=1),
        axis.title = element_text(face="bold"),
                   = element text(),
        panel.grid.minor = element_blank(),
        panel.grid.major = element_blank(),
        legend.title = element_blank(),
        legend.position = c(0.01,0.99),
        legend.background = element_blank(),
        legend.justification = c("left","top"))+
  geom_hline(yintercept = 0, linewidth=0.2)
plot2
```

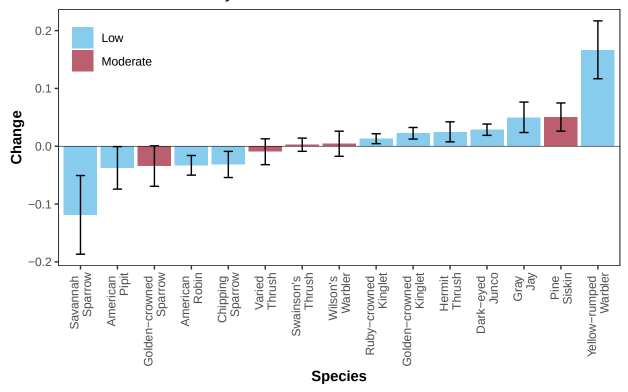
Change in species occurrence by non-breeding habitat



As we can see from this plot, species' non-breeding habitat does not show a clear correlation with direction of change. Generally, forest-dwelling species are positively affected and herbaceous-dwelling species are negatively affected.

```
plot3 <- ggplot(data=df, aes(x=NAMESEP, y=ESTIMATE, fill=CONSERVATION)) +</pre>
  theme_bw() +
  geom_bar(stat="identity",position=position_dodge(0.9)) +
  geom_errorbar(aes(ymax=CI_HIGH, ymin=CI_LOW) ,position=position_dodge(0.9),width=0.25) +
  scale_fill_manual("Legend", values = c("Low" = rgb(136/256,205/256,238/256),
                                         "Moderate" = rgb(188/256,94/256,109/256))) +
  labs(x="Species",y="Change") +
  ggtitle("Change in species occurrence \nby level of conservation concern") +
  theme(plot.title = element_text(hjust = 0.5),
        axis.text = element_text(),
        axis.text.x = element_text(angle=90, vjust=0.5, hjust=1),
        axis.title = element_text(face="bold"),
                   = element_text(),
        panel.grid.minor = element_blank(),
        panel.grid.major = element_blank(),
        legend.title = element blank(),
        legend.position = c(0.01,0.99),
        legend.background = element_blank(),
        legend.justification = c("left","top"))+
  geom_hline(yintercept = 0, linewidth=0.2)
plot3
```

Change in species occurrence by level of conservation concern



As we can see from this plot, the level of conservation concern of the species does not show a clear correlation with direction of change.

Sensitivity Analysis

Our species distribution models report coefficients as well as standard errors (SE). To ensure we are dealing with error propagation, we run a sensitivity analysis. The results we produced for the main text show the model predictions using the mean coefficients. Here we also show the predictions using coefficient-SE and coefficient+SE and evaluate whether our conclusions change.

Load the lower bound (coefficient-SE) and upper bound (coefficient+SE) data.

```
historicdata_lb <- read_xlsx(here("ProcessedData","ImageAnalysis_SensitivityAnalysis_MinusSE.xlsx"), sh
  mutate(TIME = "Historic")
repeatdata_lb <- read_xlsx(here("ProcessedData","ImageAnalysis_SensitivityAnalysis_MinusSE.xlsx"), shee
  mutate(TIME = "Repeat")
data_lb <- rbind(historicdata_lb, repeatdata_lb) %>%
  pivot_longer(cols=CF:SN, names_to="LANDCOVER", values_to="LANDCOVER_PERCENT") %>%
  pivot_longer(cols=GRAJ:VATH, names_to="SPECIES", values_to="SPECIES_OCCURRENCE_LB") %>%
  mutate(across(c(STATION, TIME, LANDCOVER, SPECIES), as.factor)) %>%
  select(STATION, TIME, SPECIES, SPECIES_OCCURRENCE_LB) %>%
  distinct()

data_lb <- data_lb %>%
  group_by(STATION, TIME) %>%
  mutate(SPECIES_DIV = diversity(SPECIES_OCCURRENCE_LB, index="shannon")) %>%
  ungroup()
```

```
# Historical bird diversity, lower bound
data_lb %>% select(STATION, TIME, SPECIES_DIV) %>% unique() %>% filter(TIME=="Historic") %>% pull(SPECI
## [1] 2.695998
# Repeat bird diversity, lower bound
data_lb %>% select(STATION, TIME, SPECIES_DIV) %>% unique() %>% filter(TIME=="Repeat") %>% pull(SPECIES
## [1] 2.699622
historicdata_ub <- read_xlsx(here("ProcessedData", "ImageAnalysis_SensitivityAnalysis_PlusSE.xlsx"), she
  mutate(TIME = "Historic")
repeatdata_ub <- read_xlsx(here("ProcessedData", "ImageAnalysis_SensitivityAnalysis_PlusSE.xlsx"), sheet
  mutate(TIME = "Repeat")
data_ub <- rbind(historicdata_ub, repeatdata_ub) %>%
  pivot_longer(cols=CF:SN, names_to="LANDCOVER", values_to="LANDCOVER_PERCENT") %>%
  pivot_longer(cols=GRAJ:VATH, names_to="SPECIES", values_to="SPECIES_OCCURRENCE_UB") %>%
  mutate(across(c(STATION, TIME, LANDCOVER, SPECIES), as.factor)) %>%
  select(STATION, TIME, SPECIES, SPECIES_OCCURRENCE_UB) %>%
  distinct()
data_ub <- data_ub %>%
  group_by(STATION, TIME) %>%
  mutate(SPECIES DIV = diversity(SPECIES OCCURRENCE UB, index="shannon")) %>%
  ungroup()
# Historical bird diversity, upper bound
data_ub %>% select(STATION, TIME, SPECIES_DIV) %>% unique() %>% filter(TIME=="Historic") %>% pull(SPECI
## [1] 2.619888
# Repeat bird diversity, upper bound
data_ub %>% select(STATION, TIME, SPECIES_DIV) %>% unique() %>% filter(TIME=="Repeat") %>% pull(SPECIES
## [1] 2.606102
Re-run the paired t-tests with the lower bound (coefficient-SE) and upper bound (coefficient+SE) on individual
species and on species diversity.
birddata_lb <- data_lb %>%
  select(STATION, TIME, SPECIES, SPECIES_OCCURRENCE_LB) %>%
  unique()
for (spp in species$CODE) {
  print(species$NAME[which(species$CODE==spp)])
  print(t.test(birddata_lb %>% filter(SPECIES==spp, TIME=="Historic") %>% pull(SPECIES_OCCURRENCE_LB),
               birddata_lb %% filter(SPECIES==spp, TIME=="Repeat") %>% pull(SPECIES_OCCURRENCE_LB),
               paired=T))
## [1] "Gray Jay"
##
## Paired t-test
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) an
## t = 4.2679, df = 45, p-value = 0.0001003
## alternative hypothesis: true mean difference is not equal to 0
```

```
## 95 percent confidence interval:
## 0.006546118 0.018245967
## sample estimates:
## mean difference
##
       0.01239604
##
## [1] "Wilson's Warbler"
##
## Paired t-test
##
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) and
## t = 2.3318, df = 45, p-value = 0.02425
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.001323701 0.018107324
## sample estimates:
## mean difference
##
       0.009715512
##
## [1] "Savannah Sparrow"
##
## Paired t-test
##
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) an
## t = -3.0426, df = 45, p-value = 0.003906
\#\# alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.1399712 -0.0284692
## sample estimates:
## mean difference
##
       -0.08422019
##
## [1] "Golden-crowned Kinglet"
##
##
  Paired t-test
##
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) an
## t = 4.6025, df = 45, p-value = 3.411e-05
\#\# alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.0009320615 0.0023825733
## sample estimates:
## mean difference
       0.001657317
##
## [1] "Ruby-crowned Kinglet"
##
## Paired t-test
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) an
## t = 3.5842, df = 45, p-value = 0.0008274
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.0004912145 0.0017514509
```

```
## sample estimates:
## mean difference
##
       0.001121333
##
## [1] "Dark-eyed Junco"
##
## Paired t-test
##
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) an
## t = 7.5834, df = 45, p-value = 1.402e-09
\#\# alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.01801018 0.03103671
## sample estimates:
## mean difference
##
        0.02452345
##
## [1] "American Robin"
##
## Paired t-test
##
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) an
## t = -3.3546, df = 45, p-value = 0.001621
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.036494291 -0.009112119
## sample estimates:
## mean difference
       -0.02280321
##
##
## [1] "Hermit Thrush"
##
## Paired t-test
##
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) and
## t = 2.4257, df = 45, p-value = 0.01935
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.001058028 0.011412449
## sample estimates:
## mean difference
##
       0.006235238
##
## [1] "Pine Siskin"
## Paired t-test
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) an
## t = 3.7199, df = 45, p-value = 0.0005508
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.01105227 0.03715161
## sample estimates:
## mean difference
```

```
##
       0.02410194
##
## [1] "American Pipit"
##
##
  Paired t-test
##
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) an
## t = -1.6827, df = 45, p-value = 0.09937
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.056933203 0.005104453
## sample estimates:
## mean difference
      -0.02591437
##
##
## [1] "Golden-crowned Sparrow"
##
## Paired t-test
##
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) an
## t = -1.6666, df = 45, p-value = 0.1025
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.048403947 0.004570439
## sample estimates:
## mean difference
##
      -0.02191675
## [1] "Swainson's Thrush"
##
## Paired t-test
##
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) an
## t = 1.7669, df = 45, p-value = 0.08403
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -1.922358e-05 2.940224e-04
## sample estimates:
## mean difference
##
     0.0001373994
## [1] "Yellow-rumped Warbler"
## Paired t-test
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) and
## t = 6.9286, df = 45, p-value = 1.3e-08
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.05164396 0.09397432
## sample estimates:
## mean difference
##
       0.07280914
##
```

```
## [1] "Chipping Sparrow"
##
##
   Paired t-test
##
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) an
## t = -1.9406, df = 45, p-value = 0.05858
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.0513965814 0.0009549555
## sample estimates:
## mean difference
       -0.02522081
##
##
## [1] "Varied Thrush"
##
## Paired t-test
##
## data: birddata_lb %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_LB) an
## t = -0.51179, df = 45, p-value = 0.6113
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.003558681 0.002116589
## sample estimates:
## mean difference
    -0.0007210462
##
# Overall bird diversity
print(t.test(data_lb %>% select(STATION, TIME, SPECIES_DIV) %>% unique() %>% filter(TIME=="Historic") %
             data_lb %>% select(STATION, TIME, SPECIES_DIV) %>% unique() %>% filter(TIME=="Repeat") %>%
##
##
   Paired t-test
##
## data: data_lb %>% select(STATION, TIME, SPECIES_DIV) %>% unique() %>% filter(TIME == "Historic") %>
## t = -1.9687, df = 45, p-value = 0.05517
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -7.331250e-03 8.357808e-05
## sample estimates:
## mean difference
      -0.003623836
birddata ub <- data ub %>%
  select(STATION, TIME, SPECIES, SPECIES_OCCURRENCE_UB) %>%
  unique()
for (spp in species$CODE) {
  print(species$NAME[which(species$CODE==spp)])
  print(t.test(birddata_ub %>% filter(SPECIES==spp, TIME=="Historic") %>% pull(SPECIES_OCCURRENCE_UB),
               birddata_ub %>% filter(SPECIES==spp, TIME=="Repeat") %>% pull(SPECIES_OCCURRENCE_UB),
              paired=T))
}
## [1] "Gray Jay"
##
```

```
## Paired t-test
##
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = 3.8584, df = 45, p-value = 0.0003613
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.05388656 0.17158307
## sample estimates:
## mean difference
##
         0.1127348
## [1] "Wilson's Warbler"
## Paired t-test
##
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = -0.48626, df = 45, p-value = 0.6291
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.04559386 0.02786020
## sample estimates:
## mean difference
      -0.008866828
##
##
## [1] "Savannah Sparrow"
## Paired t-test
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = -4.2372, df = 45, p-value = 0.0001106
\mbox{\tt \#\#} alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.24056787 -0.08555097
## sample estimates:
## mean difference
       -0.1630594
##
##
## [1] "Golden-crowned Kinglet"
##
## Paired t-test
##
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = 4.4154, df = 45, p-value = 6.256e-05
\#\# alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.06276711 0.16806013
## sample estimates:
## mean difference
##
         0.1154136
##
## [1] "Ruby-crowned Kinglet"
##
## Paired t-test
```

##

```
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = 2.9396, df = 45, p-value = 0.005172
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.02246134 0.12022453
## sample estimates:
## mean difference
##
       0.07134293
##
## [1] "Dark-eyed Junco"
## Paired t-test
##
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = 2.9979, df = 45, p-value = 0.004414
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.009185672 0.046795689
## sample estimates:
## mean difference
##
       0.02799068
##
## [1] "American Robin"
##
## Paired t-test
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = -3.9614, df = 45, p-value = 0.0002629
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.05735872 -0.01869235
## sample estimates:
## mean difference
##
       -0.03802554
## [1] "Hermit Thrush"
##
## Paired t-test
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = 3.1896, df = 45, p-value = 0.002595
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.02080370 0.09209571
## sample estimates:
## mean difference
##
         0.0564497
##
## [1] "Pine Siskin"
##
## Paired t-test
##
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = 4.1697, df = 45, p-value = 0.000137
```

```
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.03425685 0.09827318
## sample estimates:
## mean difference
       0.06626501
##
## [1] "American Pipit"
##
## Paired t-test
##
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = -2.5867, df = 45, p-value = 0.013
\mbox{\tt \#\#} alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.09428341 -0.01173342
## sample estimates:
## mean difference
       -0.05300842
##
##
## [1] "Golden-crowned Sparrow"
## Paired t-test
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = -2.4368, df = 45, p-value = 0.01884
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.089564645 -0.008505613
## sample estimates:
## mean difference
##
       -0.04903513
##
## [1] "Swainson's Thrush"
##
## Paired t-test
##
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = 2.6335, df = 45, p-value = 0.01154
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.02289099 0.17176561
## sample estimates:
## mean difference
##
         0.0973283
##
## [1] "Yellow-rumped Warbler"
##
## Paired t-test
##
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = 4.717, df = 45, p-value = 2.346e-05
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
```

```
## 0.07389142 0.18401529
## sample estimates:
## mean difference
##
         0.1289534
##
## [1] "Chipping Sparrow"
##
##
   Paired t-test
##
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = -3.7159, df = 45, p-value = 0.0005576
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -0.03720057 -0.01104835
## sample estimates:
## mean difference
##
       -0.02412446
##
## [1] "Varied Thrush"
##
   Paired t-test
##
##
## data: birddata_ub %>% filter(SPECIES == spp, TIME == "Historic") %>% pull(SPECIES_OCCURRENCE_UB) an
## t = -1.0155, df = 45, p-value = 0.3153
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
  -0.06735026 0.02219903
## sample estimates:
## mean difference
##
       -0.02257561
# Overall bird diversity
print(t.test(data_ub %>% select(STATION, TIME, SPECIES_DIV) %>% unique() %>% filter(TIME=="Historic") %
             data_ub %>% select(STATION, TIME, SPECIES_DIV) %>% unique() %>% filter(TIME=="Repeat") %>%
             paired=T))
##
##
   Paired t-test
##
## data: data_ub %>% select(STATION, TIME, SPECIES_DIV) %>% unique() %>% filter(TIME == "Historic") %>
## t = 2.369, df = 45, p-value = 0.02219
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## 0.002065134 0.025506872
## sample estimates:
## mean difference
##
          0.013786
At the species level, many species were affected and changed in probability of occurrence between the two time
points; some positively, some negatively. This is true for both the lower bound and upper bound predictions.
```

At the species level, many species were affected and changed in probability of occurrence between the two time points; some positively, some negatively. This is true for both the lower bound and upper bound predictions. Overall, based on the lower bound predictions, we cannot say that species diversity changed between the time steps (mean difference -0.004, CI=[-0.007,0.00008], t=-1.969, df=45, p-value=0.055). However, using the upper bound predictions, species diversity decreased slightly (mean difference 0.013, CI=[0.002,0.026], t=2.369, df=45, p-value=0.022).

Create a data frame with the t-test outputs to make it easier to compare.

```
birddata_lb <- data_lb %>%
  select(STATION, TIME, SPECIES, SPECIES_OCCURRENCE_LB) %>%
  unique()
df_lb <- data.frame(CODE=NA, ESTIMATE=NA, CI_LOW=NA, CI_HIGH=NA)</pre>
for (spp in species$CODE) {
  testoutput <- t.test(birddata_lb %>% filter(SPECIES==spp, TIME=="Historic") %>% pull(SPECIES_OCCURREN
                       birddata 1b %>% filter(SPECIES==spp, TIME=="Repeat") %>% pull(SPECIES OCCURRENCE
                       paired=T)
 row <- c(spp, testoutput$estimate, testoutput$conf.int[1], testoutput$conf.int[2])</pre>
 df_lb <- rbind(df_lb, row)</pre>
df_lb <- df_lb[-1,] %>% # remove the NA row
  mutate(DIRECTION_LB=ifelse(ESTIMATE<0 & CI_LOW<0 & CI_HIGH<0, "NEGATIVE",
                      ifelse(ESTIMATE>0 & CI_LOW>0 & CI_HIGH>0, "POSITIVE",
birddata_ub <- data_ub %>%
  select(STATION, TIME, SPECIES, SPECIES_OCCURRENCE_UB) %>%
  unique()
df_ub <- data.frame(CODE=NA, ESTIMATE=NA, CI_LOW=NA, CI_HIGH=NA)
for (spp in species$CODE) {
  testoutput <- t.test(birddata_ub %>% filter(SPECIES==spp, TIME=="Historic") %>% pull(SPECIES_OCCURREN
                       birddata_ub %>% filter(SPECIES==spp, TIME=="Repeat") %>% pull(SPECIES_OCCURRENCE
                       paired=T)
 row <- c(spp, testoutput$estimate, testoutput$conf.int[1], testoutput$conf.int[2])</pre>
  df_ub <- rbind(df_ub, row)</pre>
df_ub <- df_ub[-1,] %>% # remove the NA row
  mutate(DIRECTION_UB=ifelse(ESTIMATE<0 & CI_LOW<0 & CI_HIGH<0, "NEGATIVE",
                          ifelse(ESTIMATE>0 & CI_LOW>0 & CI_HIGH>0, "POSITIVE",
                                 "NULL")))
df_mean <- df %>%
  select(CODE, ESTIMATE, CI_LOW, CI_HIGH) %>%
  mutate(DIRECTION_MEAN=ifelse(ESTIMATE<0 & CI_LOW<0 & CI_HIGH<0, "NEGATIVE",
                          ifelse(ESTIMATE>0 & CI_LOW>0 & CI_HIGH>0, "POSITIVE",
                                 "NULL")))
df_merge <- left_join(df_mean, df_lb, by="CODE") %>%
  left_join(df_ub, by="CODE") %>%
  select(CODE, DIRECTION_LB, DIRECTION_MEAN, DIRECTION_UB) %>%
  distinct()
df_merge
      CODE DIRECTION_LB DIRECTION_MEAN DIRECTION_UB
##
## 1 GRAJ
               POSITIVE
                              POSITIVE
                                           POSITIVE
## 2 WIWA
               POSITIVE
                                  NULL
                                               NULL
## 3 SAVS
                              NEGATIVE
                                           NEGATIVE
               NEGATIVE
## 4 GCKI
               POSITIVE
                              POSITIVE
                                           POSITIVE
## 5 RCKI
                                           POSITIVE
               POSITIVE
                              POSITIVE
## 6 DEJU
              POSITIVE
                              POSITIVE
                                           POSITIVE
```

```
## 7 AMRO
               NEGATIVE
                              NEGATIVE
                                           NEGATIVE
## 8 HETH
               POSITIVE
                                           POSITIVE
                              POSITIVE
                                           POSITIVE
## 9 PISI
               POSITIVE
                              POSITIVE
## 10 AMPI
                   NULL
                              NEGATIVE
                                           NEGATIVE
## 11 GCSP
                   NULL
                                  NULL
                                           NEGATIVE
## 12 SWTH
                   NULL
                                  NULL
                                           POSITIVE
## 13 YRWA
               POSITIVE
                              POSITIVE
                                           POSITIVE
## 14 CHSP
                   NULL
                              NEGATIVE
                                           NEGATIVE
## 15 VATH
                   NULL
                                  NULL
                                                NULL
```

As we can see from the above table, for most species (10/15), there was no difference in conclusions. For 5 of the 15 species, the result changed from "no evidence of change" to either positive or negative. That being said, there are no situations in which the direction of change switched from "negative" to "positive" or vice-versa between the lower, mean and upper bound predictions.

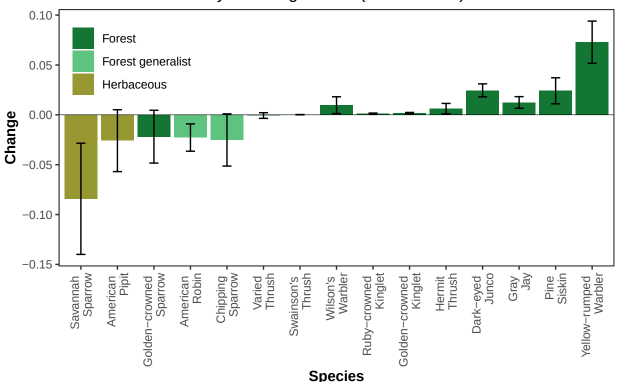
Let's prep for plotting.

```
df_lb <- left_join(df_lb, sonab, by="CODE")</pre>
df_lb$ESTIMATE <- as.numeric(df_lb$ESTIMATE)</pre>
df_lb$CI_LOW <- as.numeric(df_lb$CI_LOW)</pre>
df_lb$CI_HIGH <- as.numeric(df_lb$CI_HIGH)</pre>
df_lb$NAME <- factor(df_lb$NAME)</pre>
df_lb$BREEDING <- factor(df_lb$BREEDING)</pre>
df_lb$NONBREEDING <- factor(df_lb$NONBREEDING)</pre>
df_lb$CONSERVATION <- factor(df_lb$CONSERVATION)</pre>
df lb$NAMESEP <- factor(df lb$NAMESEP, levels=speciesorder)</pre>
df_ub <- left_join(df_ub, sonab, by="CODE")</pre>
df ub$ESTIMATE <- as.numeric(df ub$ESTIMATE)</pre>
df_ub$CI_LOW <- as.numeric(df_ub$CI_LOW)</pre>
df_ub$CI_HIGH <- as.numeric(df_ub$CI_HIGH)</pre>
df_ub$NAME <- factor(df_ub$NAME)</pre>
df_ub$BREEDING <- factor(df_ub$BREEDING)</pre>
df_ub$NONBREEDING <- factor(df_ub$NONBREEDING)</pre>
df_ub$CONSERVATION <- factor(df_ub$CONSERVATION)</pre>
df_ub$NAMESEP <- factor(df_ub$NAMESEP, levels=speciesorder)</pre>
```

Recreate Figure 4 from the main text with the lower and upper bound values to visualize these differences.

```
plot.background = element_rect(fill="transparent", colour=NA),
    panel.grid.minor = element_blank(),
    panel.grid.major = element_blank(),
    legend.title = element_blank(),
    legend.position = c(0.01,0.99),
    legend.background = element_blank(),
    legend.justification = c("left","top"))+
    geom_hline(yintercept = 0, linewidth=0.2)
```

Change in species occurrence by breeding habitat (lower bound)



```
plot5 <- ggplot(data=df_ub, aes(x=NAMESEP, y=ESTIMATE, fill=BREEDING)) +</pre>
  theme_bw() +
  geom_bar(stat="identity", position=position_dodge(0.9)) +
  geom_errorbar(aes(ymax=CI_HIGH, ymin=CI_LOW) ,position=position_dodge(0.9),width=0.25) +
  scale_fill_manual("Legend", values = c("Forest" = rgb(19/256,119/256,51/256),
                                         "Herbaceous" = rgb(153/256, 153/256, 50/256),
                                         "Forest generalist" = rgb(95/256,196/256,128/256))) +
  labs(x="Species",y="Change") +
  ggtitle("Change in species occurrence \nby breeding habitat (upper bound)") +
  theme(plot.title = element_text(hjust = 0.5),
        axis.text = element_text(),
        axis.text.x = element_text(angle=90, vjust=0.5, hjust=1),
        axis.title = element_text(face="bold"),
                   = element_text(),
        panel.background = element_rect(fill="transparent", colour=NA), # transparent background for gr
        plot.background = element_rect(fill="transparent", colour=NA),
```

```
panel.grid.minor = element_blank(),
    panel.grid.major = element_blank(),
    legend.title = element_blank(),
    legend.position = c(0.01,0.99),
    legend.background = element_blank(),
    legend.justification = c("left","top"))+
    geom_hline(yintercept = 0, linewidth=0.2)
```

Change in species occurrence by breeding habitat (upper bound)

