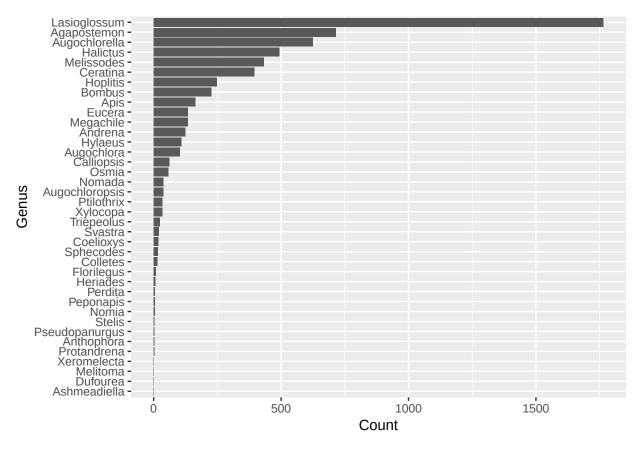
7350 Assignment 3

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April 8, 2019

```
library(tidyverse)
## -- Attaching packages ------ tidyverse 1.2.1 --
## v ggplot2 3.1.0
                        v purrr 0.2.5
## v tibble 2.0.1 v dplyr 0.8.0.1
## v tidyr 0.8.2 v stringr 1.3.1
           1.3.1
## v readr
                         v forcats 0.3.0
## -- Conflicts ----- tidyverse_conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                  masks stats::lag()
library(here)
## here() starts at C:/Users/summe/Downloads/Stat-7350-class
library(MASS)
##
## Attaching package: 'MASS'
## The following object is masked from 'package:dplyr':
##
       select
#here()
bees<-read_csv(here("Assignment 3","cleaned_bees_columns_removed.csv"),</pre>
               col_types = cols())
bees.rare.spe<-bees%>%filter(Species%in%c("sumptuosa", "pumila", "obliqua", "nasonii", "montivaga", "mesilla
bees%>%group_by(Genus)%>%summarise(Count = n())%>%
  ggplot(aes(x=reorder(Genus,Count),y=Count)) +
  geom col()+
  coord_flip()+
 labs(x ="Genus")
```

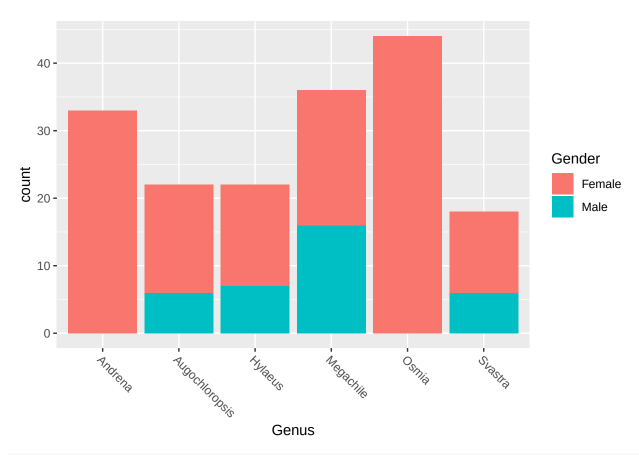


```
genus_count<-bees%>%count(Genus)%>%
  filter(n<=200)

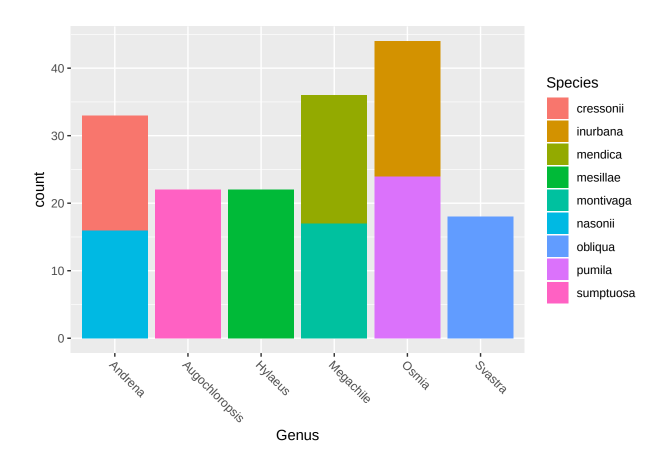
species_counts <- bees%>%
    count(Species)%>%
  filter(n>=15)%>%filter(n<=30)

bees_new<-bees%>%
  filter(Species%in%species_counts$Species)%>%
  filter(Genus%in%genus_count$Genus)

ggplot(data= bees_new, mapping= aes(x=Genus, fill = Gender)) +
    geom_bar()+
    theme(axis.text.x=element_text(angle = -45, hjust = 0))
```

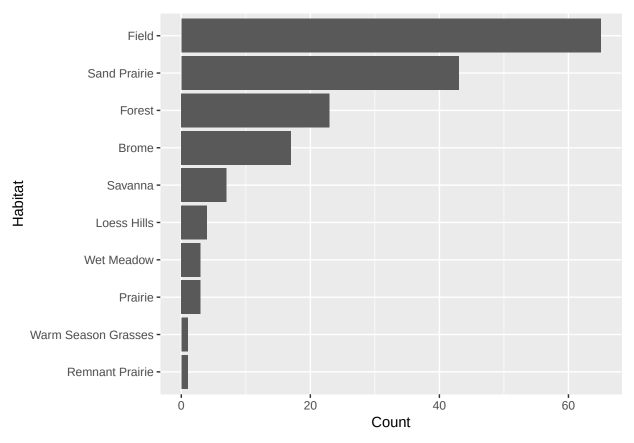


```
ggplot(data= bees_new, mapping= aes(x=Genus, fill = Species)) +
  geom_bar()+
  theme(axis.text.x=element_text(angle = -45, hjust = 0))
```



To find the specific habitat for those different species we study

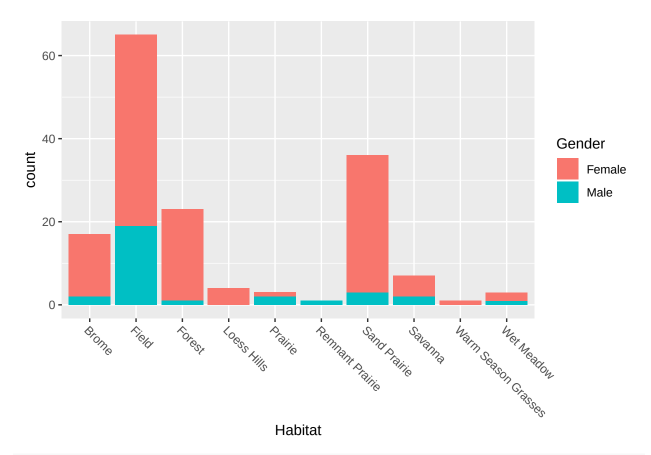
```
bees.rare.spe%>%group_by(Habitat)%>%
  summarise(Count = n())%>%drop_na()%>%
  ggplot(aes(x= reorder(Habitat,Count),y=Count))+
  geom_col()+
  coord_flip()+
  labs(x="Habitat")
```



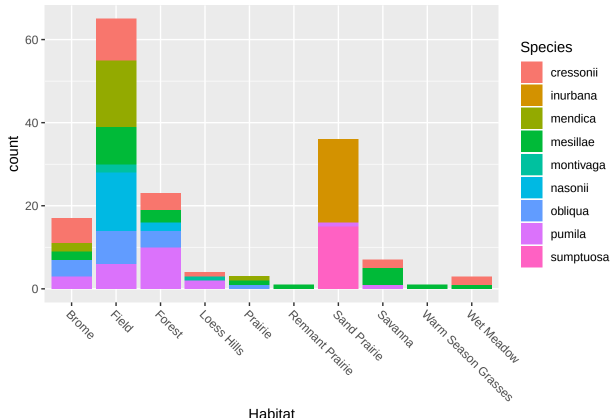
```
habitat_count_f<-bees.rare.spe%>%count(Habitat)%>%drop_na()
month_f<-bees.rare.spe%>%count(Month)%>%drop_na()

bees_new_f1<-bees.rare.spe%>%
    filter(Habitat%in%habitat_count_f$Habitat)%>%
    filter(Month%in%month_f$Month)

ggplot(bees_new_f1, aes(Habitat,fill = Gender)) +
    geom_bar()+
    theme(axis.text.x=element_text(angle = -45, hjust = 0))
```

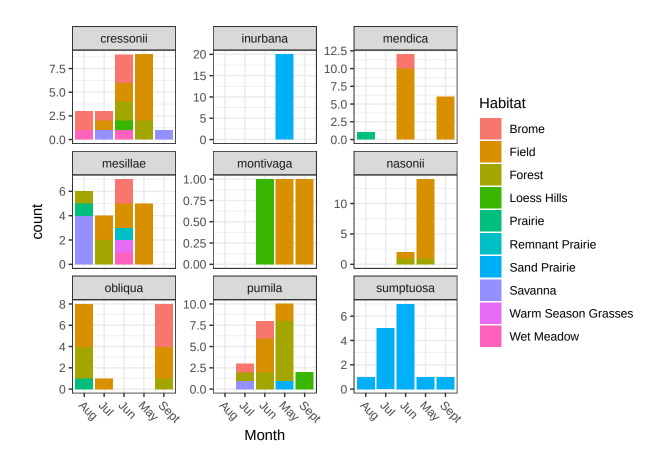


```
ggplot(bees_new_f1, aes(Habitat,fill = Species)) +
  geom_bar()+
  theme(axis.text.x=element_text(angle = -45, hjust = 0))
```



Habitat

```
ggplot(bees_new_f1, mapping = aes(x=Month, fill = Habitat)) +
  geom_bar()+
 facet_wrap(~ Species, scale = "free_y")+
  theme_bw()+
  theme(axis.text.x = element_text(angle = -45,hjust = 0))
```



Fisher Exact test

Independency between Habitat and Species

```
spec<-bees.rare.spe%>%count(Species)
hab<-bees.rare.spe%>%count(Habitat)
fisher.matr<-cbind(spec$n,hab$n)
fisher.test(fisher.matr,simulate.p.value = TRUE, B = 1e5)

##
## Fisher's Exact Test for Count Data with simulated p-value (based
## on 1e+05 replicates)
##
## data: fisher.matr
## p-value = 1e-05
## alternative hypothesis: two.sided</pre>
```

Independency between month and Species

```
spec<-bees.rare.spe%>%count(Species)
mont<-bees.rare.spe%>%count(Month)
```

```
fisher.matr<-cbind(spec$n,mont$n)
fisher.test(fisher.matr,simulate.p.value = TRUE, B = 1e5)

##
## Fisher's Exact Test for Count Data with simulated p-value (based
## on 1e+05 replicates)
##
## data: fisher.matr
## p-value = 1e-05
## alternative hypothesis: two.sided</pre>
```