Visualizations

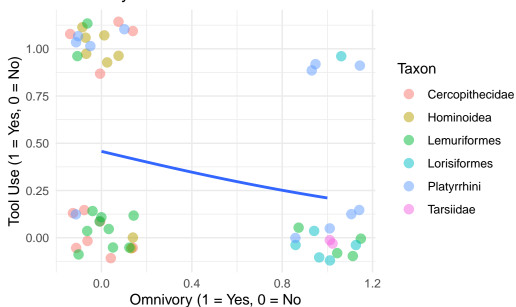
```
#load packages
  library(tidyverse)
-- Attaching packages ----- tidyverse 1.3.2 --
v ggplot2 3.3.6 v purrr 0.3.4
v tibble 3.1.8
               v dplyr 1.0.10
v tidyr 1.2.0
               v stringr 1.4.1
             v forcats 0.5.2
v readr
      2.1.2
-- Conflicts ----- tidyverse_conflicts() --
x dplyr::filter() masks stats::filter()
x dplyr::lag()
            masks stats::lag()
  library(tidymodels)
-- Attaching packages ----- tidymodels 1.0.0 --
      1.0.1 v rsample
1.0.0 v tune
v broom
                              1.1.0
v dials
                              1.0.0
1.0.1 v yardstick 1.0.0
v parsnip
v recipes
           1.0.1
-- Conflicts ----- tidymodels_conflicts() --
x scales::discard() masks purrr::discard()
x dplyr::filter() masks stats::filter()
x recipes::fixed() masks stringr::fixed()
x dplyr::lag()
               masks stats::lag()
x yardstick::spec() masks readr::spec()
x recipes::step() masks stats::step()
* Use suppressPackageStartupMessages() to eliminate package startup messages
```

```
library(ape)
Attaching package: 'ape'
The following object is masked from 'package:rsample':
    complement
The following object is masked from 'package:dials':
    degree
  library(caper)
Loading required package: MASS
Attaching package: 'MASS'
The following object is masked from 'package:dplyr':
    select
Loading required package: mvtnorm
  library(phytools)
Loading required package: maps
Attaching package: 'maps'
The following object is masked from 'package:purrr':
    map
  library(nlme)
```

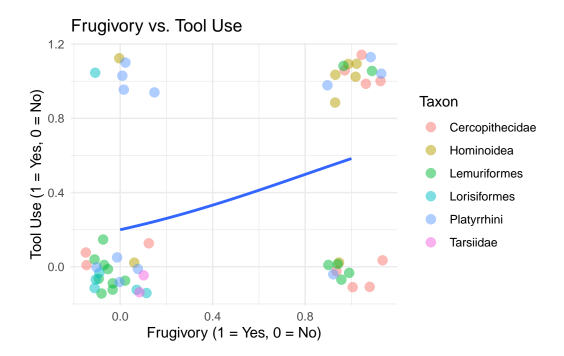
library(knitr)

```
Attaching package: 'nlme'
The following object is masked from 'package:dplyr':
    collapse
  library(gee)
  library(TreeTools)
Attaching package: 'TreeTools'
The following object is masked from 'package:phytools':
    as.multiPhylo
  library(patchwork)
Attaching package: 'patchwork'
The following object is masked from 'package:MASS':
    area
  library(phylolm)
  library(broom)
  library(readxl)
  #load data and phylogenetic tree
  tool_use_expanded <- read_xlsx("PrimateToolUseDataset.xlsx", sheet = "Dataset")</pre>
  tooluse.tree <- read.nexus("ToolUsePhylo54.nex")</pre>
  #EDA
  set.seed(13)
```

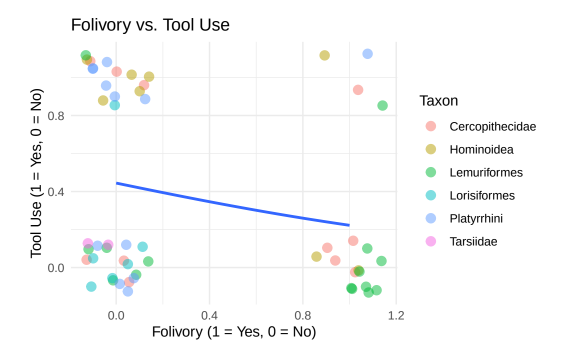
Omnivory vs. Tool Use

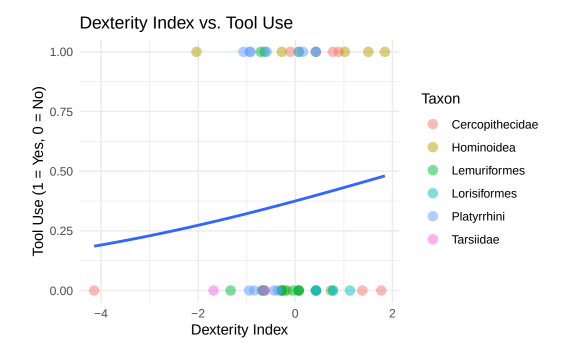


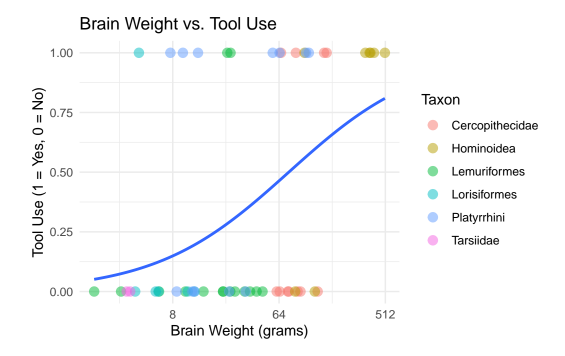
```
ggplot(tool_use_expanded, aes(x = Frugivore, y = Tool_Use)) +
geom_jitter(width = 0.15, height = 0.15, aes(color = Taxon), size = 3, alpha = 0.5) +
geom_smooth(method = glm, se = F, method.args= list(family="binomial")) + labs(title = "
y = "Tool Use (1 = Yes, 0 = No)",
x = "Frugivory (1 = Yes, 0 = No)") +
theme_minimal()
```



```
ggplot(tool_use_expanded, aes(x = Folivore, y = Tool_Use)) +
  geom_jitter(width = 0.15, height = 0.15, aes(color = Taxon), size = 3, alpha = 0.5) +
  geom_smooth(method = glm, se = F, method.args= list(family="binomial")) + labs(title =
      y = "Tool Use (1 = Yes, 0 = No)",
      x = "Folivory (1 = Yes, 0 = No)") + theme_minimal()
```







```
ggplot(tool_use_expanded, aes(Group_Size, Tool_Use)) +
  geom_point(aes(color = Taxon), size = 3, alpha = 0.5) + geom_smooth(method = glm, se = Fill the second black is a second black is
```

