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**Toronto, Canada**

A report on

**Executive Summary Report 3**

**Subject**

Introduction to Analytics – ALY 6000

**Guided by**

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**Submitted by**

Name NUID Date of Submission

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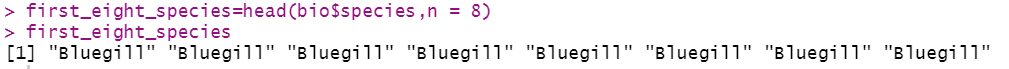
**Introduction**

The dataset of FSAdata(InchLake2) is primarily focuses on total weights and lengths of different fish species captured in InchLake2 from May, 2007 to May, 2008 by Derek H. Ogle, personal collection. In addition, this report consists of key findings, methodology, conclusion, bibliography, and appendix.

**Key findings**

The dataset has only 676 observations, and 7variables (NetID, FishID, Species, Length, Weight, and Year of capture).

The first five species of the dataset by analyzing it via head function



Group by count of all the species in the dataset by table function according to 676 observations

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As seen from the figure, Largemouth Bass accounted for 33 percent, which was the highest amongst all these species; while, Tadpole Madtom has the least amount of species 0.887574.

LargeMouth Bass species has the highest weight and length around 1070 and 429, respectively; whilst, Bluegill accounted for the lowest.

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Many species of the fish do not have tag number in the dataset.

Also, the data provided and executed in R language has several Null values that need to be replaced by mean or 0 in order to get the exact analysis and for model preparation.

**Methodology**

Data visualization in R language with InchLake2(InchBio.csv) dataset provided in this report for analysis.

Chart, bar chart

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Chart, waterfall chart

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By considering graphs, Largemouth bass and Bluegill has the most counts as compared to other species.

Fish relative frequency shows the proportion out of 676 observations.

This report includes descriptive as well as inferential statistics. Furthermore, Null values need to be fixed first before analyzing the data. After, the data was analyzed and processed in R such as (Exploratory Data Analytics, plotting via simple plot, and ggplot2, filtration by dplyr(chaining and piping).

**Conclusion**

Considering all the points, graphs, details generated above, it can be recapitulated that:

The pareto chart has all the species with two axis chart and cumulative frequency. The axis shown in the graph at left side depicts cumulative counts, and the right axis illustrates cumulative frequency.

Many observations do not have scale; thus, the data has to be strongly scaled before making analysis report.

Surprisingly, Pumkinseed and Tadpole Madtom has the least count of the species throughout the observations captured by the author.

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**Appendix**

#1 printing my name and load the necessary libraries.

print("Raj Tank") #printing my name

#installing libraries

install.packages("FSA")

install.packages("FSAdata")

install.packages("magrittr")

install.packages("dplyr")

install.packages("tidyr")

install.packages("plyr")

install.packages("tidyverse")

#importing those libaries

library(FSA)

library(FSAdata)

library(magrittr)

library(dplyr)

library(tidyr)

library(plyr)

library(tidyverse)

#2 importing the dataset "inchBio.csv".

#setting up the working directory

setwd("C:/Users/baps/Downloads")

bio=read.csv("inchBio.csv") #naming the table bio

bio

#3 Displaying the headtail structure of the dataset.

head\_tail=headtail(bio)

print(head\_tail)

#4 creating "<counts>", which counts and lists all the species records.

counts=bio$species

counts

#displaying the species

spec\_records=bio$species

spec\_records

#5 Depicting 8 species' names.

first\_eight\_species=head(bio$species,n = 8)

first\_eight\_species

#6 creating "<tmp>" that contains different species and records of it.

tmp=count(bio$species)

tmp

#7 creating a subset "<tmp2>" of just species variable, and display 5 records.

tmp2=bio$species

head(tmp2,n=5) # to get only 5 recors of the dataframe

#8 creating a table "<w>", and display the class of it.

w<-table(bio$species)

w

#checking the class of the variable

class(w)

#9 conver"<w>" to dataframe named "<t>",and print results.

# converting table into dataframe

t=as.data.frame(w)

t

class(t)

#10 Extracting and displaying the frequency values from "<t>" dataframe

extracted\_freq=t$Freq

#displaying the frequency

extracted\_freq

#11 creating a table named "<cSpec>" from the bio and that table displays the

#number of species

cSpec<-table(bio$species)

cSpec

class(cSpec)

#12 creating a table name "<cSpecPct>" that displays the species and percentage

#of records. Also confirm class of it

cSpecPct <- (table(bio$species)/676)\*100

print(cSpecPct)

class(cSpecPct)

#13 converting the table "<SpecPct>"to dataframe named"<u>" and confirm

#the class is dataframe

u=as.data.frame(cSpecPct)

class(u)

class(cSpec)

cSpec=as.data.frame(cSpec)

class(cSpec)

cSpec

class(cSpecPct)

cSpecPct=as.data.frame(cSpecPct)

cSpecPct

class(cSpecPct)

#14 creating a barplot of "<cSec>" with the following details mentioned below

ggplot(cSpec, aes(y=Freq, x=Var1)) +

geom\_bar(stat = "identity",fill="green") +

coord\_flip()+labs(y="Counts",x="Species",title = "Fish Counts")

#using barplot method

barplot(height = cSpec$Freq,names=cSpec$Var1,col = "green",

horiz = TRUE,ylab ="Species",xlab = "COUNTS",main = "Fish Counts",cex.lab =0.60)

#15 creating a braplot of "<cSpecPct>" with following details mentioned below

barplot(height = cSpecPct$Freq,names=cSpecPct$Var1,las=1,

col = "lightblue",ylim=c(0,40),cex.names = 0.60,

main ="Fish Relative Frequency")

#16 rearranging "<u>" cSpecPct dataframe in descneding order of relative

#frequency and save that object as "<d>"

d<-cSpec %>% arrange(desc(cSpec$Freq))

d

#17 rename <d> column and var 1 to species, and Freq to RelFreq

#renaming columns

colnames(d)[colnames(d)=="Var1"] <- "Species"

colnames(d)[colnames(d)=="Freq"] <- "RelFreq"

class(d)

d

#18 Add new variables to <d> and call them cumfreq, counts, and cumcounts

d=mutate(d,cumfreq=cumsum(RelFreq),counts=RelFreq\*676,

cumcounts=cumsum(counts))

print(d)

view(d)

#19 adding new variable ",def\_par>" to store parameter variable

def\_par=par(no.readonly = TRUE)

def\_par

#20 creating "<pc>" with following details mentioned below

pc<-barplot(d$counts,width = 1,ylab="Cumulative Counts",

main ="Species Pareto",,cex.names = 0.7,names.arg=d$Species,las=2,

border = NA,axes =F, space= 0.15,

ylim=c(0,3.5)\*max(d$counts,na.rm = TRUE))

#21 adding cumulative count line to "<pc>" with following details

lines(pc,d$cumcounts,type="b",cex=0.7,pch=16,col="cyan4")

#22 placing a grey box around pareto chart

box(col="grey")

#reference:https://www.statmethods.net/advgraphs/parameters.html

#23 adding left side axiswith following details

axis(at = c(0,d$cumcounts),side = 2,las=1, col.axis="grey62",cex.axis=0.8,

col = "grey62")

#24 adding axis on right side of the box with following details

axis(side = 4,cex.axis=0.80,col = "cyan4",col.axis="cyan4",at = c(0,d$cumcounts),

las=1,labels = paste(c(0,round(d$cumfreq\*100)),"%",sep=""))

#25 displaying the finished species pareto chart, having my last name on the plot

pc<-barplot(d$counts,width = 1,ylab="Cumulative Counts",

main ="Species Pareto(Raj Tank)",,cex.names = 0.7,names.arg=d$Species,las=2,

border = NA,axes =F, space= 0.15,

ylim=c(0,3.5)\*max(d$counts,na.rm = TRUE))

lines(pc,d$cumcounts,type="b",cex=0.7,pch=16,col="cyan4")

box(col="grey")

axis(at = c(0,d$cumcounts),side = 2,las=1, col.axis="grey62",cex.axis=0.8,

col = "grey62")

axis(side = 4,cex.axis=0.80,col = "cyan4",col.axis="cyan4",at = c(0,d$cumcounts),

las=1,labels = paste(c(0,round(d$cumfreq\*100)),"%",sep=""))

#26 commit my repo in [Github](https://github.com/rajtank/week_3_R_Project)