# Project 2 Notebook

Introduction (40 points)

• 10 points for background on the protein/gene/species of interest and where the data is sourced from

### Introduction

• 10 points for specific, measurable, and clear scientific question

### Scientific Question

When examining dog breeds (canis lupus familiaris), will breeds of a similar size (e.g. Cocker Spaniel, English Cocker Spaniel) have more related genes and SNP's surrounding longevity than breeds of a different size (e.g.Doberman Pinscher, Miniature Pinscher)?

Note: I only selected 6 genes most closely associated with life span (HMGA2 , IGF1 (done) , IGSF1 (too big), IRS4 (too big), LCORL (done), and SMAD2 (too big) ). There are more genes involved in this, but these are the most significant.

Note: Size will be determined by the AKC. You can filter by all AKC recognized dog breeds by size. This is categorical data; if it is easier for me to work with numerical instead, I will instead use the ideal height and weight, as outlined in the official standard of each breed.

:when possible used regions by paper if regions too big used dimension of genes on ncbi

• 10 points for clear, specific, and measurable scientific hypothesis that is in the form of an if-then statement

### Scientific Hypothesis

If you examine canine breeds, then breeds of a similar size (e.g.Cocker Spaniel, English Cocker Spaniel) they will have more related SNPs and or fragments of genes surrounding longevity than breeds of a different size (e.g.Doberman Pinscher, Miniature Pinscher).

• 10 points for description of what analyses were done and how the data was downloaded for the project ## Analysis Performed:

#### SNP's

- EDA: Scatterplots of SNP nucleotide vs size to check for visible trends before analysis
- Multiple Sequence Alignment (of SNP+border sequences), which was then visualized with msaPrettyPrint() \_ Clustering of MSA results, which were visualized as Dendrograms

#### Gene Fragments:

 Multiple Sequence Alignment (of SNP+border sequences), which was then visualized with msaPrettyPrint() \_ Clustering of MSA results, which were visualized as Dendrograms

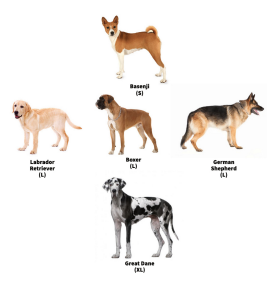
### **Data Sourcing**

- Dog breed information (sizing): American Kennel Club LINK \_ Data downloads:
- SNP and gene positionings:
- 25 points for definition of each of the packages loaded
- 5 points for correctly loading all of the packages needed

```
#for reading in fasta files
library("BiocManager")
#for reading in excel files
library("readxl")
#for multiple sequence alignment
library("msa")
## Loading required package: Biostrings
## Loading required package: BiocGenerics
## Attaching package: 'BiocGenerics'
## The following objects are masked from 'package:stats':
##
##
       IQR, mad, sd, var, xtabs
## The following objects are masked from 'package:base':
##
##
       anyDuplicated, append, as.data.frame, basename, cbind, colnames,
##
       dirname, do.call, duplicated, eval, evalq, Filter, Find, get, grep,
##
       grepl, intersect, is.unsorted, lapply, Map, mapply, match, mget,
##
       order, paste, pmax, pmax.int, pmin, pmin.int, Position, rank,
       rbind, Reduce, rownames, sapply, setdiff, sort, table, tapply,
##
       union, unique, unsplit, which.max, which.min
## Loading required package: S4Vectors
## Loading required package: stats4
##
## Attaching package: 'S4Vectors'
## The following objects are masked from 'package:base':
##
##
       expand.grid, I, unname
## Loading required package: IRanges
## Loading required package: XVector
## Loading required package: GenomeInfoDb
##
## Attaching package: 'Biostrings'
## The following object is masked from 'package:base':
##
##
       strsplit
## Attaching package: 'msa'
```

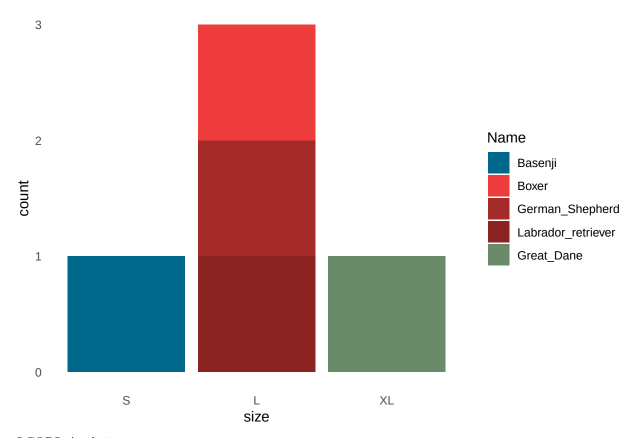
```
## The following object is masked from 'package:BiocManager':
##
##
       version
#for msa pretty print
library("tinytex")
#visualization of results
library("ggplot2")
#for clustering of DNA segs
library("DECIPHER")
## Loading required package: RSQLite
## Loading required package: parallel
#for cleaning up dendograms
library('dendextend')
##
## -----
## Welcome to dendextend version 1.15.2
## Type citation('dendextend') for how to cite the package.
## Type browseVignettes(package = 'dendextend') for the package vignette.
## The github page is: https://github.com/talgalili/dendextend/
##
## Suggestions and bug-reports can be submitted at: https://github.com/talgalili/dendextend/issues
## You may ask questions at stackoverflow, use the r and dendextend tags:
    https://stackoverflow.com/questions/tagged/dendextend
##
##
   To suppress this message use: suppressPackageStartupMessages(library(dendextend))
##
##
## Attaching package: 'dendextend'
## The following object is masked from 'package:Biostrings':
##
##
       nnodes
## The following object is masked from 'package:stats':
##
       cutree
```

First we will be performing MSA and clustering with various sequence data (SNPS and gene fragments). Unfortunately, the scope of this data is limited to the 5 breeds shown below



```
#global variable
alignment_name<<-""
#notebook functions
#align fasta from file_name with names from name file (visualization purposes) after alignment displays
mult_alingments<-function(file_name,fasta_names,name,big_aln=FALSE,dna_set=TRUE){
  #read in fasta for all dogs
  if(dna_set=="TRUE"){
  string_set<-readDNAStringSet(file=file_name,use.names=FALSE)</pre>
  }
  else{
      string_set<-readAAStringSet(file=file_name, use.names=FALSE)</pre>
  }
  #read in seq names as list
  table=read.table(fasta_names, header = FALSE, sep = "\n")[["V1"]]
  #update names for pretty print
  names(string_set)<-table</pre>
  #align unnamed seqs
  alignment<-msa(string_set,order="input")</pre>
  #if seq cant be display with msa pretty print, return
  if(big_aln==TRUE){
    return(alignment)
  #update global variable so multiple pretty print runs dont overrun eachother
  alignment_name<<-gsub(" ", "", paste(name,".pdf"), fixed = TRUE)</pre>
  #return pretty alignment, does not show up on my console
  msaPrettyPrint(alignment, file=alignment_name,output="pdf",
                 showNames="right",showLogo="top",askForOverwrite=FALSE,
                 showNumbering="none",paperWidth=6,paperHeight=3)
```

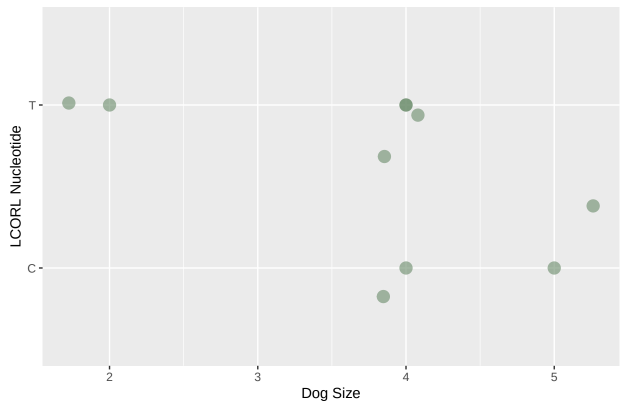
```
return(alignment_name)
#have figure with white background, no gridline and only axis ticks, no lines
tune_figure<-function(fig,addons){</pre>
  return(fig+theme_minimal()+theme(
    plot.background = element_blank(),
    panel.grid.major = element_blank(),
    panel.grid.minor = element blank(),
    panel.border = element_blank()))
}
#create dendogram based on fasta files, names of items clustered in fasta_names
create_dendogram<-function(fasta_path, fasta_names, fig_title){</pre>
  #grab DNA info from coallated file
  dna <- string_set<-readDNAStringSet(file=fasta_path,use.names=FALSE)</pre>
  #qet sequence names
  names(dna) = read.table(fasta\_names, \frac{header}{leasta\_names}, \frac{header}{leasta\_names} = FALSE, \frac{sep}{leasta\_names} = "\n")[["V1"]]
  #create distance matrix for clustering
  d1 <- DistanceMatrix(dna, type="dist")</pre>
  #form dendogram
  dendogram <- IdClusters (d1, method="complete", cutoff=0.05, showPlot=FALSE, type="dendrogram")
  #fix names being cut-off
  nodePar \leftarrow list(lab.cex = 0.6, pch = c(NA, 19),
                 cex = 0.7, col = "black")
#plot results
plot(as.dendrogram(dendogram), ylab = "Height", nodePar = nodePar, main=fig title)
  return(as.dendrogram(dendogram))
#visualize size breakdown of dogs
snps<-read excel("dog snps.xlsx")</pre>
#fix ordering of legend
snps$Name <- factor(snps$Name, levels = c("Basenji", "Boxer", "German_Shepherd", "Labrador_retriever", "G</pre>
p<-ggplot(data = snps, aes(size))+scale_x_discrete(limits = c("S","L","XL"))+geom_bar(aes(fill = Name))
tune_figure(p,add_ons)
```



## LCORL Analysis

```
#visualize LCORL SNP by size
p<-ggplot(data = snps, mapping = aes(y=lcorl,x=size_num))+geom_point(size=4,alpha=0.6,color="darkseagre
p+geom_jitter(size=4,alpha=0.6,color="darkseagreen4")</pre>
```

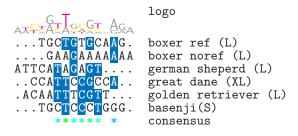
# LCORL SNP Distribution by Dog Size



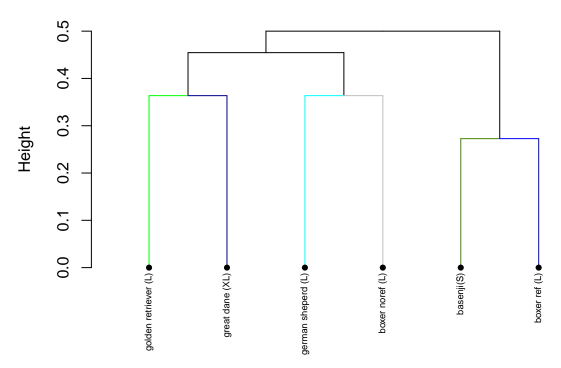
```
#LCORL CALL
alignment<-mult_alingments("fasta/LCORL_file.txt","fasta/names.txt","LCORL")</pre>
```

## use default substitution matrix
print(alignment\_name)

## [1] "LCORL.pdf"



# **LCORL Extended Fragment Dendogram**



## 'dendrogram' with 2 branches and 6 members total, at height 0.5

```
#IGF1 ANALYSIS
```

```
abbrev_x <- c("A","C","'G","'T")
print(length(abbrev_x))</pre>
```

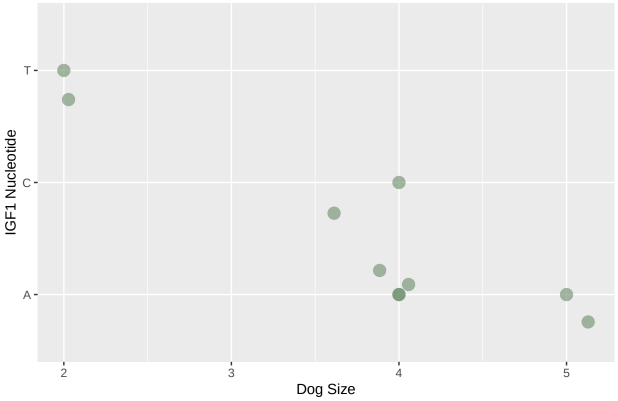
### ## [1] 4

```
print(length(seq(0,4,by=1)))
```

## ## [1] 5

```
#visualize IGF1 SNP by size
p<-ggplot(data = snps, mapping = aes(y=igf1,x=size_num))+geom_point(size=4,alpha=0.6,color="darkseagreenp+geom_jitter(size=4,alpha=0.6,color="darkseagreen4")</pre>
```

# IGF1 SNP Distribution by Dog Size



```
#IGF1 CALL
alignment<-mult_alingments("fasta/igf1.fasta","fasta/igf1_names.txt","igf1")</pre>
```

## use default substitution matrix

```
logo

TTCCTTTTCTA.... Boxer no ref (L)

....TCTGAAGAGTA
...GGCATTCCCCT.. Basenji (S)

AGGTCATGACT.... Great dane (XL)
..AATTCAGTGAA.. Labrador retriever (L)

* ****

Inon-conserved

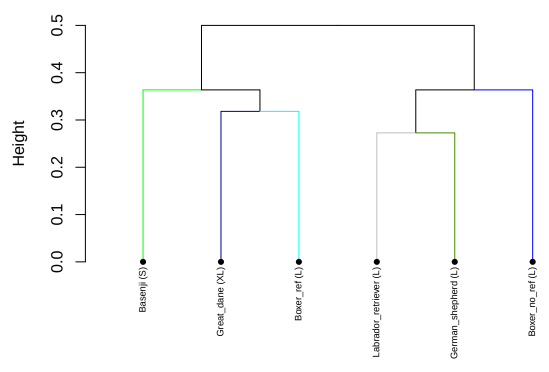
Inon-conserved

Inon-conserved
Inon-conserved
```

http://www.sthda.com/english/wiki/beautiful-dendrogram-visualizations-in-r-5-must-known-methods-unsupervised-machine-learning#plot.dendrogram-function for look and non cut off stuff

```
#Cluster IGF1 extended fragment
create_dendogram("fasta/igf1.fasta", "fasta/igf1_names.txt", "IGF1 Extended Fragment Dendogram")
```

# **IGF1 Extended Fragment Dendogram**



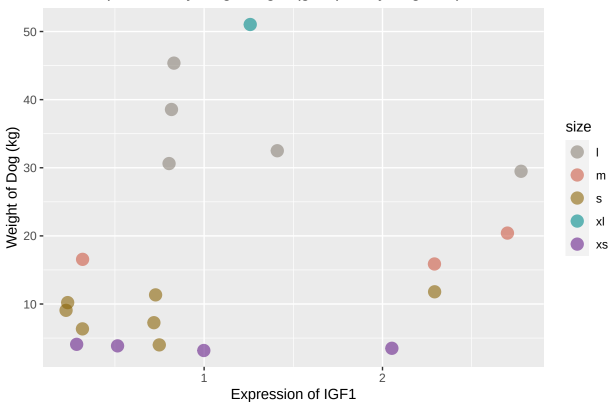
## 'dendrogram' with 2 branches and 6 members total, at height 0.5

The differences sequence-wise seem to be minor when looking at such a narrow scope and small subsection of dogs. Thankfully there is also expression data for the genes studied which covers more dog-breeds, depicted below



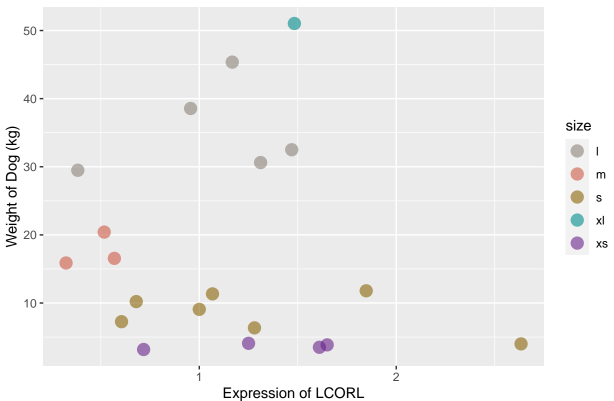
```
#expression data
expression<-read_excel("dog snps.xlsx",sheet="IGF1")
myColors <- c("antiquewhite4", " coral3", "darkgoldenrod4","darkcyan", "darkcrchid4")
p<-ggplot(data = expression, mapping = aes_string(x="norm_exp",y="weight_kg" ,col= "size"))+geom_point(p+scale_color_manual(values=myColors)</pre>
```





```
expression<-read_excel("dog snps.xlsx",sheet="LCORL")
p<-ggplot(data = expression, mapping = aes_string(x="norm_exp",y="weight_kg" ,col= "size"))+geom_point(
p+scale_color_manual(values=myColors)</pre>
```





```
expression<-read_excel("dog snps.xlsx",sheet="SMAD2")
p<-ggplot(data = expression, mapping = aes_string(x="norm_exp",y="weight_kg" ,col= "size"))+geom_point(
p+scale_color_manual(values=myColors)</pre>
```

