clustering

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2 step markov chain I guess

write up explanation of aggregation before vs after, conditional probabilities along with the tree would be a good idea This section is going cover two step clustering based on various clustering methods for comparison. Visit data will be combined into one bee result for each orchard, note in principle this is fundementally flawed as the data does depend on previous years and the assumption that independent of the past is false as pesticide last year affects bee population this year as would be expected. In these scenarios each implementation will have a maximum of 8 clusters. Two after pre-bloom, 4 after and 8 after post bloom.

```
load("data")
markov_data <- data_2012 %>%
  select(ends with(".pre"), ends with(".blm"), ends with(".pos"), orchard) %>%
  unique()
#Standardising the data for purpose of clustering
markov_data <- markov_data %>%
  mutate(eiqB11F.pre = (eiqB11F.pre - mean(eiqB11F.pre))/sd(eiqB11F.pre)) %%
  mutate(eiqB11I.pre= (eiqB11I.pre - mean(eiqB11I.pre))/sd(eiqB11I.pre)) %>%
  mutate(eiqB11F.blm = (eiqB11F.blm - mean(eiqB11F.blm))/sd(eiqB11F.blm)) %>%
  mutate(eiqB11I.blm = (eiqB11I.blm - mean(eiqB11I.blm))/sd(eiqB11I.blm)) %>%
  mutate(eiqB11T.blm= (eiqB11T.blm - mean(eiqB11T.blm))/sd(eiqB11T.blm)) %>%
  mutate(eiqB11F.pos = (eiqB11F.pos - mean(eiqB11F.pos))/sd(eiqB11F.pos)) %>%
  mutate(eiqB11I.pos = (eiqB11I.pos - mean(eiqB11I.pos))/sd(eiqB11I.pos)) %>%
  mutate(eiqB11T.pos = (eiqB11T.pos - mean(eiqB11T.pos))/sd(eiqB11T.pos))
#function to generate the agglomaerative clustering
aggl <- function(data){</pre>
  agnes(dist(data, method = "euclidian"),
                   diss=TRUE, method = "ward")
}
#function to reduce copy pasting for each node
aggl_c <- function(data, ends, c_num){</pre>
  #define temp dataset of this stage by cluster
   temp <- data %>%
    filter(cluster == c_num) %>%
    select(ends_with(ends))
#If statement as if 1 element in dataset it'll error
if(nrow(temp) > 1){
  temp$cluster <- cutree(aggl(temp) , k = 2)</pre>
```

```
output <- data %>%
   filter(cluster == c_num) %>%
   select(-cluster) %>%
   mutate(cluster = temp$cluster)
   #Else statement for if 1 element in dataset
    #If only 1 element in the dataset -> set the cluster number to 1
   output <- data %>%
      filter(cluster == c_num) %>%
      select(-cluster) %>%
      mutate(cluster = 1)
 }
  output
####Agglomerative heirachicale clustering
#pre-bloom step
temp <- markov_data %>%
  select(ends_with(".pre"))
temp$cluster <- cutree(aggl(temp) , k = 2)</pre>
aggl_node1 <- markov_data %>%
 mutate(cluster = temp$cluster)
#during bloom step 1
aggl_node2 <- aggl_c(aggl_node1, ".blm", 1)
#during bloom step 2
aggl_node3 <- aggl_c(aggl_node1, ".blm", 2)</pre>
#post bloom step 1
aggl_node4 <- aggl_c(aggl_node2, ".pos", 1)</pre>
#post bloom step 2
aggl_node5 <- aggl_c(aggl_node2, ".pos", 2)</pre>
#post bloom step 3
aggl_node6 <- aggl_c(aggl_node3, ".pos", 1)</pre>
#post bloom step 4
#Note this node only has 1 element in it so it is automatically set to 1
aggl_node7 <- aggl_c(aggl_node3, ".pos", 2)</pre>
orchard_node_agglom <- tibble(orchard = aggl_node4 %% filter(cluster == 1) %% select(orchard) %% as_
  bind_rows(tibble(orchard = aggl_node4 %>% filter(cluster == 2) %>% select(orchard) %>% as_vector, nod
  bind_rows(tibble(orchard = aggl_node5 %>% filter(cluster == 1) %>% select(orchard) %>% as_vector, nod
  bind_rows(tibble(orchard = aggl_node5 %>% filter(cluster == 2) %>% select(orchard) %>% as_vector, nod
  bind_rows(tibble(orchard = aggl_node6 %>% filter(cluster == 1) %>% select(orchard) %>% as_vector, nod
  bind_rows(tibble(orchard = aggl_node6 %>% filter(cluster == 2) %>% select(orchard) %>% as_vector, nod
  bind_rows(tibble(orchard = aggl_node7 %>% filter(cluster == 1) %>% select(orchard) %>% as_vector, nod
```

kmeans approach: Even running the 1001 times it is different everytime: store output later on but run it like 1million times at each stage.

```
#function to generate the kmeans clustering
k_clustering <- function(data, n = 1001){</pre>
#creating the dataset to then calculate the optimal cluster from
  for(i in 1:n){
  if(i == 1){
     k_list <- tibble(kmeans(data, 2)$cluster)</pre>
 }else{
     k_list <- bind_cols(k_list, tibble(kmeans(data, 2)$cluster))</pre>
 }
  apply(k_list[ ,1:length(k_list)], 1, mfv1) %>%
    enframe(value = "cluster") %>%
    select(cluster)
}
#pre-bloom step
#Getting the most common clustering
kmeans_c <- function(data, ends, c_num){</pre>
  data <- data %>%
    filter(cluster == c_num) %>%
    select(-cluster)
  if(nrow(data) > 2){
    temp <- data %>%
    select(ends_with(ends))
  cluster <- k_clustering(temp, 2)</pre>
 output <- bind_cols(data, cluster)</pre>
 }else{
    #Not more than 2 rows and get the error message as:
    #number of cluster centres must lie between 1 and nrow(x)
    output <- data %>%
      mutate(cluster = 1)
 }
output
temp <- k_clustering(markov_data %>% select(-orchard))
kmeans_node1 <- markov_data %>%
 bind_cols(temp)
#during bloom step 1
kmeans_node2 <- kmeans_c(kmeans_node1, ".blm", 1)</pre>
#during bloom step 2
```

```
kmeans_node3 <- kmeans_c(kmeans_node1, ".blm", 2)</pre>
#post bloom step 1
kmeans_node4 <- kmeans_c(kmeans_node2, ".pos", 1)</pre>
#post bloom step 2
kmeans node5 <- kmeans c(kmeans node2, ".pos", 2)
#post bloom step 3
kmeans_node6 <- kmeans_c(kmeans_node3, ".pos", 1)</pre>
#post bloom step 4
kmeans_node7 <- kmeans_c(kmeans_node3, ".pos", 2)</pre>
#End timepoint nodes
orchard_node_kmeans <- tibble(orchard = kmeans_node4 %% filter(cluster == 1) %% select(orchard) %% a
  bind_rows(tibble(orchard = kmeans_node4 %>% filter(cluster == 2) %>% select(orchard) %>% as_vector, n
  bind_rows(tibble(orchard = kmeans_node5 %>% filter(cluster == 1) %>% select(orchard) %>% as_vector, n
  bind_rows(tibble(orchard = kmeans_node5 %>% filter(cluster == 2) %>% select(orchard) %>% as_vector, n
  bind_rows(tibble(orchard = kmeans_node6 %>% filter(cluster == 1) %>% select(orchard) %>% as_vector, n
  bind_rows(tibble(orchard = kmeans_node6 %>% filter(cluster == 2) %>% select(orchard) %>% as_vector, n
  bind_rows(tibble(orchard = kmeans_node7 %>% filter(cluster == 1) %>% select(orchard) %>% as_vector, n
  bind_rows(tibble(orchard = kmeans_node7 %>% filter(cluster == 2) %>% select(orchard) %>% as_vector, n
Just a look at the difference in the final clusters between the two methods
clusterings <- orchard_node_agglom %>%
  arrange(orchard) %>%
  bind_cols(node_km = orchard_node_kmeans %>% arrange(orchard) %>%
              select(node) %>% as_vector)
clusterings
## # A tibble: 19 x 3
##
      orchard node node_km
##
      <fct> <dbl>
                      <dbl>
## 1 A
                          8
                  8
## 2 B
                  9
## 3 C
                  9
                           9
                  9
                           9
## 4 D
## 5 E
                  8
                           8
## 6 F
                  8
                           8
## 7 G
                           9
                  9
## 8 H.nooil
                  8
                           8
                           8
## 9 I
                  8
## 10 J
                  8
                          8
                  8
                          8
## 11 K
## 12 L
                  8
                          8
## 13 M
                  8
                          8
## 14 N
                  8
                          8
## 15 O.nooil
                 12
                          8
## 16 P
                 10
                         12
## 17 Q
                  8
                          8
## 18 R
                  8
                          8
```

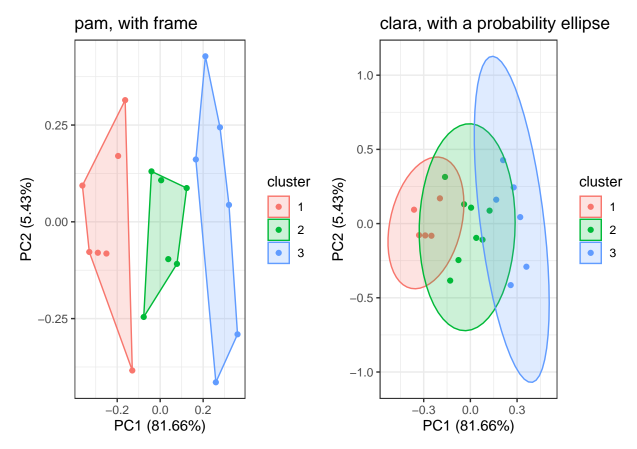
```
## 19 S 8 8
```

Looking at top two PCAs just to see how much variation is absorbed by these

```
p1 <- autoplot(pam(markov_data, 3), frame = TRUE) +
    theme_bw() +
    ggtitle("pam, with frame")

p2 <- autoplot(clara(markov_data, 3), frame = TRUE, frame.type = 'norm') +
    theme_bw() +
    ggtitle("clara, with a probability ellipse")

grid.arrange(p1, p2 ,nrow = 1)</pre>
```



Non-markov structuring

This section will approach looking at the data as one time point with 1 clustering point rather than 3 as above: Method Justification - Maximum linkage function is best for both standardised and unstandardised.

```
maximum_cluster <- data_2012 %>%
  arrange(orchard) %>%
  select(ends_with(".pre"),ends_with(".blm"), ends_with(".pos")) %>%
  filter(row_number() %% 2 == 1) %>%
  mutate(eiqB11F.pre = (eiqB11F.pre - mean(eiqB11F.pre))/sd(eiqB11F.pre)) %>%
  mutate(eiqB11I.pre= (eiqB11I.pre - mean(eiqB11I.pre))/sd(eiqB11I.pre)) %>%
  mutate(eiqB11F.blm = (eiqB11F.blm - mean(eiqB11F.blm))/sd(eiqB11F.blm)) %>%
  mutate(eiqB11I.blm = (eiqB11I.blm - mean(eiqB11I.blm))/sd(eiqB11I.blm)) %>%
```

```
mutate(eiqB11T.pos = (eiqB11T.pos - mean(eiqB11T.pos))/sd(eiqB11T.pos))
euclidean_cluster <- maximum_cluster</pre>
# matrix of methods to compare,
#rerun - without %>%mutate segment for non-standardised values
m <- c( "average", "single", "complete", "ward")</pre>
names(m) <- c( "average", "single", "complete", "ward")</pre>
distances <- c("euclidean", "maximum", "manhattan", "canberra",
               "minkowski")
names(distances) <- c("euclidean", "maximum", "manhattan",</pre>
                       "canberra", "minkowski")
clust_comps <- matrix(nrow = length(distances), ncol = length(m),</pre>
                      dimnames = list(distances,m))
# function to compute coefficient to see which is the best method
ac <- function(distance, linkage) {</pre>
  dista <- dist(maximum_cluster , method = distance)</pre>
  #Agglomerative Nesting form of Hierarchical Clustering
  agnes(dista, method = linkage)$ac
for(i in 1:length(distances)) {
 for(j in 1:length(m)) {
    clust_comps[i,j] <- ac(distances[i], m[j])</pre>
 }
}
#In future or in write up, perhaps change, needs more clustering analysis
#Look at literature in more detail to see other benefits/drawbacks and assumptions
#Then decide on actual best method, for now using euclidian as used that before
standardised_clust_comps <- clust_comps</pre>
Clustering code - Maximum
#NBclust function breaks when using index = all, instead using all the working indexes instead
fviz_nbclust(NbClust(maximum_cluster, distance = "maximum",
                                 min.nc = 2, max.nc = 8, method = "ward.D2", index = c("kl", "ch", "ccc"
## Among all indices:
## ========
```

mutate(eiqB11T.blm= (eiqB11T.blm - mean(eiqB11T.blm))/sd(eiqB11T.blm)) %>%
mutate(eiqB11F.pos = (eiqB11F.pos - mean(eiqB11F.pos))/sd(eiqB11F.pos)) %>%
mutate(eiqB11I.pos = (eiqB11I.pos - mean(eiqB11I.pos))/sd(eiqB11I.pos)) %>%

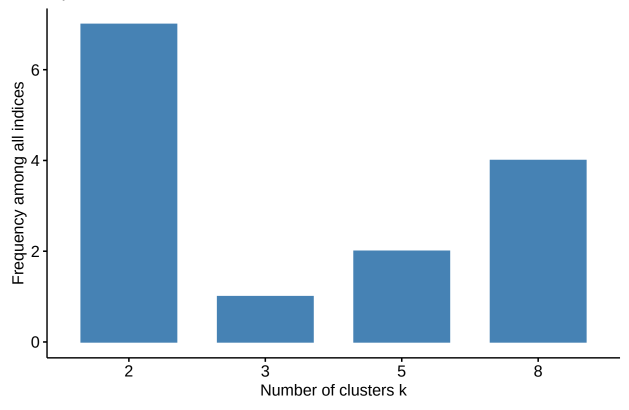
* According to the majority rule, the best number of clusters is 2 .

* 7 proposed 2 as the best number of clusters
* 1 proposed 3 as the best number of clusters
* 2 proposed 5 as the best number of clusters
* 4 proposed 8 as the best number of clusters

Conclusion

=========

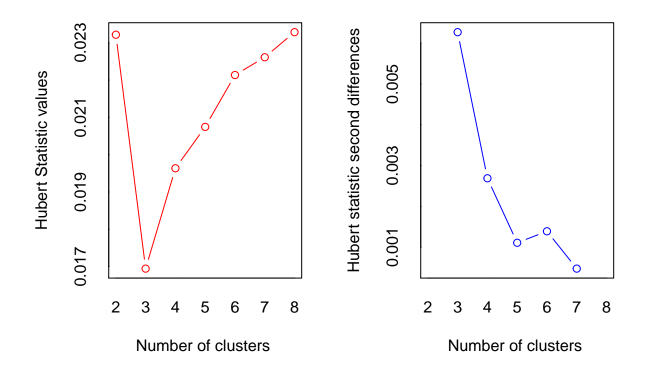
Optimal number of clusters -k = 2



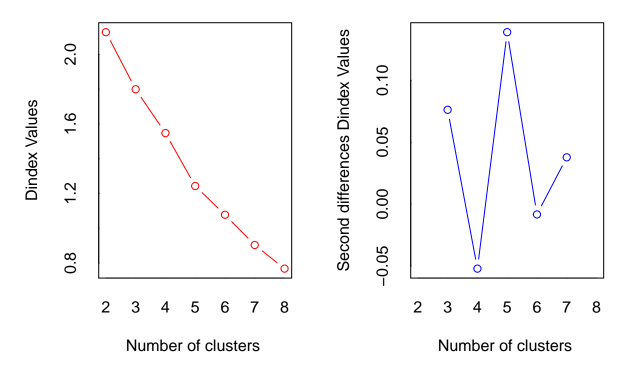
```
##
    eiqB11F.pre eiqB11I.pre eiqB11F.blm eiqB11I.blm eiqB11T.blm eiqB11F.pos
     0.56294059
              ## 1
## 2
     0.88788914 -0.3418658 -0.19022893 -0.30585457 -0.30620065
                                                     0.01058593
## 3
     0.88788914
              -0.3418658 -0.19022893 -0.30585457 -0.30620065
                                                     0.01058593
## 4
               -0.3418658 -0.19022893 -0.30585457 -0.30620065
     0.88788914
                                                     0.01058593
    -1.21724288
               -0.3418658 -0.47141495
                                 0.25463750 -0.26826632
                                                     0.05673110
## 6
    -1.78818873
               -0.3418658 -0.71087017
                                  ## 7
     0.92973857
                1.1014288
                        1.78834024
                                  0.06780681 -0.26972533
                                                     2.88563060
## 8
     0.01045773
              -0.3418658 2.26526351 -0.36201840 0.01478210
                                                     1.10101634
    -0.28688425
               ## 10 -0.05899826
              -0.3418658 -1.93232366 -1.17238220 -0.30620065
                                                     0.53623960
     1.50859712
                       0.15454685 -0.81247522 -0.29744657 -0.54716872
## 11
               -0.1356809
## 12
     1.44670216
              -0.3068769 0.15156608 -0.81247522 -0.30167771 -0.56622868
              -0.3418658 -0.18559218 -0.44683724 -0.30124000 -0.62240541
## 13 -0.33664640
## 14 -1.63608890
               ## 15 -1.58069994
               3.8693055 -1.36167172 -0.74943419 -0.30620065 -1.18316953
## 16 -0.14885146
              ## 17 -0.08097801 -0.3418658 -0.85792151 1.91089732 -0.28285645 -0.80197030
```

```
## 18 -0.02629240 -0.3418658 -0.15710926 -0.07432206 -0.05816852 -0.72071468
## 19 0.03876764 -0.3418658 1.45747473 -0.96491990 -0.30620065 1.85488796
      eiqB11I.pos eiqB11T.pos cluster
      0.07606715 -0.29236499
## 1
## 2
      1.78589159 -0.29236499
                                    1
## 3
      1.78589159 -0.29236499
                                    1
     1.78589159 -0.29236499
                                    1
## 5 -0.67656322 -0.29236499
                                    1
## 6
      0.61131654 -0.29236499
                                    1
## 7
      1.48904351 -0.23224030
                                    1
## 8 -0.90112187 -0.29236499
                                    1
## 9 -0.10337469 -0.17211561
                                    1
## 10 -1.07800026 -0.29236499
                                    1
## 11 -0.07517669 -0.16918270
                                    1
## 12 -0.07517669 0.04491986
                                    1
## 13 -0.78166487 -0.20291118
                                    1
## 14 -1.05339109 -0.29236499
                                    1
## 15 -0.68425358 -0.29236499
                                    2
## 16 -0.61093877 4.10700262
                                    2
## 17 -0.65349212 -0.29236499
                                    1
## 18 -0.28179116 0.03025530
                                    1
## 19 -0.55915698 -0.18971308
                                    1
#Clustering summaries based on non-markov approach (Before Adjustment)
maximum_non_markov_clusters <- data_2012 %>%
  group by(orchard) %>%
  summarise(mean_honey_ab = mean(apisAb),
           mean_wild_ab = mean(wildAbF),
            mean_social_rich = mean(socialRichF)) %>%
  mutate(cluster = maximum_cluster$cluster) %>%
  group_by(cluster) %>%
  summarise(mean_honey_ab = mean(mean_honey_ab),
            mean_wild_ab = mean(mean_wild_ab),
            mean_social_rich = mean(mean_social_rich),
           n = n()
```

Clustering code - euclidean



*** : The Hubert index is a graphical method of determining the number of clusters.
In the plot of Hubert index, we seek a significant knee that corresponds to a
significant increase of the value of the measure i.e the significant peak in Hubert
index second differences plot.
##



```
*** : The D index is a graphical method of determining the number of clusters.
                 In the plot of D index, we seek a significant knee (the significant peak in Dindex
##
                 second differences plot) that corresponds to a significant increase of the value of
##
                 the measure.
##
##
  ************************
## * Among all indices:
## * 8 proposed 2 as the best number of clusters
## * 5 proposed 3 as the best number of clusters
## * 7 proposed 5 as the best number of clusters
## * 4 proposed 8 as the best number of clusters
##
##
                    ***** Conclusion *****
##
  * According to the majority rule, the best number of clusters is 2
##
##
                    ************
## Among all indices:
## ========
## * 2 proposed 0 as the best number of clusters
## * 8 proposed 2 as the best number of clusters
## * 5 proposed 3 as the best number of clusters
## * 7 proposed 5 as the best number of clusters
## * 4 proposed 8 as the best number of clusters
```

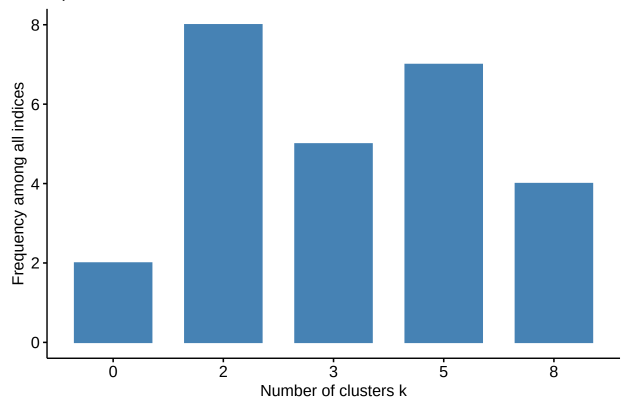
##

```
## Conclusion
```

=========

* According to the majority rule, the best number of clusters is 2 .

Optimal number of clusters -k = 2



```
eiqB11F.pre eiqB11I.pre eiqB11F.blm eiqB11I.blm eiqB11T.blm eiqB11F.pos
##
               ## 2
     0.88788914 - 0.3418658 - 0.19022893 - 0.30585457 - 0.30620065
                                                         0.01058593
## 3
      0.88788914 -0.3418658 -0.19022893 -0.30585457 -0.30620065
                                                         0.01058593
## 4
      0.88788914 \quad \hbox{-0.3418658} \quad \hbox{-0.19022893} \quad \hbox{-0.30585457} \quad \hbox{-0.30620065}
                                                         0.01058593
     -1.21724288
## 5
                -0.3418658 -0.47141495 0.25463750 -0.26826632
                                                         0.05673110
                -0.3418658 -0.71087017
## 6
     -1.78818873
                                    ## 7
      0.92973857
                 1.1014288 1.78834024
                                    0.06780681 -0.26972533
                                                         2.88563060
## 8
      0.01045773
                -0.3418658 2.26526351 -0.36201840
                                              0.01478210
## 9
     -0.28688425
                 0.2579449 -0.21970544
                                    0.87817061 -0.28577447 -0.22916831
## 10 -0.05899826
                -0.3418658 -1.93232366 -1.17238220 -0.30620065
                                                         0.53623960
                ## 11
     1.50859712
     1.44670216
                ## 13 -0.33664640
                -0.3418658 -0.18559218 -0.44683724 -0.30124000 -0.62240541
## 14 -1.63608890 -0.3418658 0.37048710 0.12282444 -0.30620065 -0.53312454
```

```
## 15 -1.58069994 3.8693055 -1.36167172 -0.74943419 -0.30620065 -1.18316953
## 16 -0.14885146 -0.3418658 0.30623494 -0.51446307 4.07083681 -0.08621861
## 17 -0.08097801 -0.3418658 -0.85792151 1.91089732 -0.28285645 -0.80197030
## 18 -0.02629240 -0.3418658 -0.15710926 -0.07432206 -0.05816852 -0.72071468
##
     eigB11I.pos eigB11T.pos cluster
     0.07606715 -0.29236499
## 1
## 2
      1.78589159 -0.29236499
## 3
     1.78589159 -0.29236499
                                  2
## 4
                                  2
     1.78589159 -0.29236499
## 5 -0.67656322 -0.29236499
                                  1
     0.61131654 -0.29236499
## 6
                                  1
## 7
      1.48904351 -0.23224030
                                  3
                                  3
## 8 -0.90112187 -0.29236499
## 9 -0.10337469 -0.17211561
                                  1
## 10 -1.07800026 -0.29236499
                                  1
                                  2
## 11 -0.07517669 -0.16918270
## 12 -0.07517669 0.04491986
                                  2
## 13 -0.78166487 -0.20291118
                                  1
## 14 -1.05339109 -0.29236499
                                  1
## 15 -0.68425358 -0.29236499
                                  4
## 16 -0.61093877 4.10700262
## 17 -0.65349212 -0.29236499
                                  1
## 18 -0.28179116 0.03025530
                                  1
                                  3
## 19 -0.55915698 -0.18971308
#Clustering summaries based on non-markov approach (Before Adjustment)
euclidean_non_markov_clusters <- data_2012 %>%
 group_by(orchard) %>%
 summarise(mean_honey_ab = mean(apisAb),
           mean_wild_ab = mean(wildAbF),
           mean_social_rich = mean(socialRichF)) %>%
 mutate(cluster = euclidean_cluster$cluster) %>%
 group_by(cluster) %>%
 summarise(mean_honey_ab = mean(mean_honey_ab),
           mean_wild_ab = mean(mean_wild_ab),
           mean_social_rich = mean(mean_social_rich),
           n = n()
```

Variable justification

```
bee_correlations <- data.frame(matrix(0, nrow = 0, ncol = 0))

for(i in 1:7){
   for(j in 1:7){
    bee_correlations[i,j] <- round(cor(data_2012[5+i], data_2012[5 + j]), 2)
   }
}
for(i in 1:7){
   names(bee_correlations)[i] <- colnames(data_2012[5 + i])
}
bee_correlations</pre>
```

```
apisAb.1 apisAb.2 apisAb.3 apisAb.4 apisAb.5 apisAb.6 apisAb.7 wildAbF.1
## 1
         1.00
                   1.00
                             1.00
                                       1.00
                                                1.00
                                                          1.00
                                                                    1.00
                                                                               0.29
## 2
         0.29
                   0.29
                             0.29
                                       0.29
                                                0.29
                                                          0.29
                                                                    0.29
                                                                               1.00
## 3
         0.19
                   0.19
                             0.19
                                       0.19
                                                0.19
                                                                    0.19
                                                                               0.91
                                                          0.19
## 4
         0.25
                   0.25
                             0.25
                                       0.25
                                                0.25
                                                          0.25
                                                                    0.25
                                                                               0.96
## 5
         0.13
                   0.13
                             0.13
                                       0.13
                                                0.13
                                                          0.13
                                                                    0.13
                                                                               0.92
## 6
         0.18
                   0.18
                             0.18
                                       0.18
                                                0.18
                                                          0.18
                                                                    0.18
                                                                               0.29
## 7
         0.22
                   0.22
                             0.22
                                       0.22
                                                0.22
                                                          0.22
                                                                    0.22
                                                                               0.31
     wildAbF.2 wildAbF.3 wildAbF.4 wildAbF.5 wildAbF.6 wildAbF.7 wildRichF.1
          0.29
                     0.29
                                0.29
                                           0.29
                                                      0.29
                                                                 0.29
## 1
                                                                              0.19
## 2
          1.00
                     1.00
                                1.00
                                           1.00
                                                      1.00
                                                                 1.00
                                                                              0.91
                     0.91
                                0.91
                                           0.91
                                                      0.91
                                                                              1.00
## 3
          0.91
                                                                 0.91
                                           0.96
## 4
          0.96
                     0.96
                                0.96
                                                      0.96
                                                                 0.96
                                                                              0.81
## 5
          0.92
                     0.92
                                0.92
                                           0.92
                                                      0.92
                                                                 0.92
                                                                              0.92
## 6
          0.29
                     0.29
                                0.29
                                           0.29
                                                      0.29
                                                                 0.29
                                                                              0.50
## 7
          0.31
                     0.31
                                0.31
                                           0.31
                                                      0.31
                                                                 0.31
                                                                              0.53
     wildRichF.2 wildRichF.3 wildRichF.4 wildRichF.5 wildRichF.6 wildRichF.7
##
            0.19
                          0.19
                                       0.19
                                                    0.19
                                                                 0.19
## 2
             0.91
                          0.91
                                       0.91
                                                    0.91
                                                                 0.91
                                                                              0.91
## 3
             1.00
                          1.00
                                       1.00
                                                    1.00
                                                                 1.00
                                                                              1.00
## 4
             0.81
                          0.81
                                       0.81
                                                    0.81
                                                                 0.81
                                                                              0.81
             0.92
                          0.92
                                       0.92
                                                    0.92
                                                                 0.92
                                                                              0.92
## 6
             0.50
                          0.50
                                       0.50
                                                    0.50
                                                                 0.50
                                                                              0.50
## 7
             0.53
                          0.53
                                       0.53
                                                    0.53
                                                                 0.53
                                                                              0.53
     solitaryAbF.1 solitaryAbF.2 solitaryAbF.3 solitaryAbF.4 solitaryAbF.5
## 1
               0.25
                              0.25
                                             0.25
                                                            0.25
                                                                            0.25
## 2
               0.96
                              0.96
                                             0.96
                                                             0.96
                                                                            0.96
## 3
               0.81
                              0.81
                                             0.81
                                                            0.81
                                                                            0.81
                                             1.00
                                                                            1.00
## 4
               1.00
                              1.00
                                                             1.00
                                                            0.92
                                                                            0.92
## 5
               0.92
                              0.92
                                             0.92
## 6
               0.02
                              0.02
                                             0.02
                                                            0.02
                                                                            0.02
## 7
               0.05
                              0.05
                                             0.05
                                                            0.05
                                                                            0.05
     solitaryAbF.6 solitaryAbF.7 solitaryRichF.1 solitaryRichF.2
## 1
               0.25
                              0.25
                                               0.13
## 2
                              0.96
               0.96
                                                0.92
                                                                 0.92
## 3
               0.81
                              0.81
                                               0.92
                                                                 0.92
## 4
               1.00
                              1.00
                                               0.92
                                                                 0.92
## 5
               0.92
                              0.92
                                                1.00
                                                                 1.00
## 6
               0.02
                              0.02
                                               0.13
                                                                 0.13
                              0.05
## 7
               0.05
                                               0.16
                                                                 0.16
     solitaryRichF.3 solitaryRichF.4 solitaryRichF.5 solitaryRichF.6
## 1
                 0.13
                                  0.13
                                                    0.13
                                                                     0.13
## 2
                 0.92
                                  0.92
                                                    0.92
                                                                     0.92
## 3
                                  0.92
                                                    0.92
                                                                     0.92
                 0.92
## 4
                                  0.92
                                                    0.92
                                                                     0.92
                 0.92
                                  1.00
## 5
                 1.00
                                                    1.00
                                                                     1.00
## 6
                 0.13
                                  0.13
                                                    0.13
                                                                     0.13
## 7
                 0.16
                                  0.16
                                                    0.16
                                                                     0.16
     solitaryRichF.7 socialAbF.1 socialAbF.2 socialAbF.3 socialAbF.4
## 1
                 0.13
                              0.18
                                           0.18
                                                        0.18
                                                                     0.18
## 2
                 0.92
                              0.29
                                           0.29
                                                        0.29
                                                                     0.29
## 3
                 0.92
                              0.50
                                           0.50
                                                        0.50
                                                                     0.50
## 4
                 0.92
                              0.02
                                           0.02
                                                        0.02
                                                                     0.02
## 5
                 1.00
                              0.13
                                           0.13
                                                        0.13
                                                                     0.13
```

## 7	##	6	0.1	1.00	1.00	1.00	1.00
## 1 0.18 0.18 0.18 0.18 0.22 0.22 ## 2 0.29 0.29 0.29 0.31 0.31 ## 3 0.50 0.50 0.50 0.50 0.53 0.53 ## 4 0.02 0.02 0.02 0.05 0.05 ## 5 0.13 0.13 0.13 0.16 0.16 ## 6 1.00 1.00 1.00 0.99 0.99 ## 7 0.99 0.99 0.99 1.00 1.00 1.00 ## * socialRichF.3 socialRichF.4 socialRichF.5 socialRichF.6 socialRichF.7 ## 1 0.22 0.22 0.22 0.22 0.22 0.22 ## 2 0.31 0.31 0.31 0.31 0.31 0.31 ## 3 0.53 0.53 0.53 0.53 0.53 0.53 ## 4 0.05 0.05 0.05 0.05 0.05 ## 5 0.16 0.16 0.16 0.16 0.16 ## 6 0.99 0.99 0.99 0.99 0.99 0.99	##	7	0.1	0.99	0.99	0.99	0.99
## 2 0.29 0.29 0.29 0.31 0.31 ## 3 0.50 0.50 0.50 0.50 0.53 0.53 ## 4 0.02 0.02 0.02 0.05 0.05 ## 5 0.13 0.13 0.13 0.16 0.16 ## 7 0.99 0.99 0.99 1.00 1.00 1.00 ## * * * * * * * * * * * * * * * * * *	##		socialAbF.5 so	ocialAbF.6 soci	ialAbF.7 socia	lRichF.1 socia	lRichF.2
## 3 0.50 0.50 0.50 0.50 0.53 0.53 ## 4 0.02 0.02 0.02 0.05 0.05 ## 5 0.13 0.13 0.13 0.16 0.16 ## 6 1.00 1.00 1.00 0.99 0.99 ## 7 0.99 0.99 0.99 1.00 1.00 ## socialRichF.3 socialRichF.4 socialRichF.5 socialRichF.6 socialRichF.7 ## 1 0.22 0.22 0.22 0.22 0.22 0.22 ## 2 0.31 0.31 0.31 0.31 0.31 0.31 ## 3 0.53 0.53 0.53 0.53 0.53 0.53 ## 4 0.05 0.05 0.05 0.05 0.05 ## 5 0.16 0.16 0.16 0.16 0.16 ## 6 0.99 0.99 0.99 0.99 0.99	##	1	0.18	0.18	0.18	0.22	0.22
## 4 0.02 0.02 0.02 0.05 0.05 ## 5 0.13 0.13 0.13 0.16 0.16 ## 6 1.00 1.00 1.00 0.99 0.99 ## 7 0.99 0.99 0.99 1.00 1.00 ## 1 0.22 0.22 0.22 0.22 0.22 ## 2 0.31 0.31 0.31 0.31 0.31 0.31 ## 3 0.53 0.53 0.53 0.53 0.53 ## 4 0.05 0.05 0.05 0.05 0.05 ## 5 0.16 0.16 0.16 0.16 0.16 ## 6 0.99 0.99 0.99 0.99 0.99	##	2	0.29	0.29	0.29	0.31	0.31
## 5	##	3	0.50	0.50	0.50	0.53	0.53
## 6	##	4	0.02	0.02	0.02	0.05	0.05
## 7 0.99 0.99 0.99 1.00 1.00 ## socialRichF.3 socialRichF.4 socialRichF.5 socialRichF.6 socialRichF.7 ## 1 0.22 0.22 0.22 0.22 0.22 ## 2 0.31 0.31 0.31 0.31 0.31 0.31 ## 3 0.53 0.53 0.53 0.53 0.53 0.53 ## 4 0.05 0.05 0.05 0.05 0.05 0.05 ## 5 0.16 0.16 0.16 0.16 0.16 ## 6 0.99 0.99 0.99 0.99	##	5	0.13	0.13	0.13	0.16	0.16
## socialRichF.3 socialRichF.4 socialRichF.5 socialRichF.6 socialRichF.7 ## 1 0.22 0.22 0.22 0.22 0.22 ## 2 0.31 0.31 0.31 0.31 0.31 0.31 ## 3 0.53 0.53 0.53 0.53 0.53 0.53 ## 4 0.05 0.05 0.05 0.05 0.05 0.05 ## 5 0.16 0.16 0.16 0.16 0.16 ## 6 0.99 0.99 0.99 0.99 0.99	##	6	1.00	1.00	1.00	0.99	0.99
## 1 0.22 0.22 0.22 0.22 0.22 0.22 ## 2 0.31 0.31 0.31 0.31 0.31 ## 3 0.53 0.53 0.53 0.53 0.53 0.53 ## 4 0.05 0.05 0.05 0.05 0.05 ## 5 0.16 0.16 0.16 0.16 0.16 ## 6 0.99 0.99 0.99 0.99 0.99	##	7	0.99	0.99	0.99	1.00	1.00
## 2 0.31 0.31 0.31 0.31 0.31 ## 3 0.53 0.53 0.53 0.53 0.53 ## 4 0.05 0.05 0.05 0.05 0.05 ## 5 0.16 0.16 0.16 0.16 0.16 ## 6 0.99 0.99 0.99 0.99 0.99	##		${\tt socialRichF.3}$	${\tt socialRichF.4}$	socialRichF.5	socialRichF.6	socialRichF.7
## 3 0.53 0.53 0.53 0.53 0.53 ## 4 0.05 0.05 0.05 0.05 0.05 ## 5 0.16 0.16 0.16 0.16 0.16 ## 6 0.99 0.99 0.99 0.99 0.99	##	1	0.22	0.22	0.22	0.22	0.22
## 4 0.05 0.05 0.05 0.05 0.05 ## 5 0.16 0.16 0.16 0.16 0.16 ## 6 0.99 0.99 0.99 0.99	##	2	0.31	0.31	0.31	0.31	0.31
## 5 0.16 0.16 0.16 0.16 0.16 ## 6 0.99 0.99 0.99 0.99	##	3	0.53	0.53	0.53	0.53	0.53
## 6 0.99 0.99 0.99 0.99	##	4	0.05	0.05	0.05	0.05	0.05
	##	5	0.16	0.16	0.16	0.16	0.16
## 7 1.00 1.00 1.00 1.00 1.00	##	6	0.99	0.99	0.99	0.99	0.99
	##	7	1.00	1.00	1.00	1.00	1.00

This demonstrates that to consider the affects on wild bees, just two variables need to be considered as there is very high correlation between the outcomes. WildAbF accounts a .9+ correlation with all wild variables except for the for the solitary bee variable. Then for this SocialRichF could be used.

Updating the bee values to match the environments

In this particular analysis, the interest is in the affects of pesticides on bee population. As such, it is necessary to correct, as best we can, for extra factors (confounding?) to make the bee values representative.

All these graphs were plotted as part of EDA by the use of lapply to the a basic plotting function. Here the relationships between some of the main extra factors (are they confounding?) will be examined.

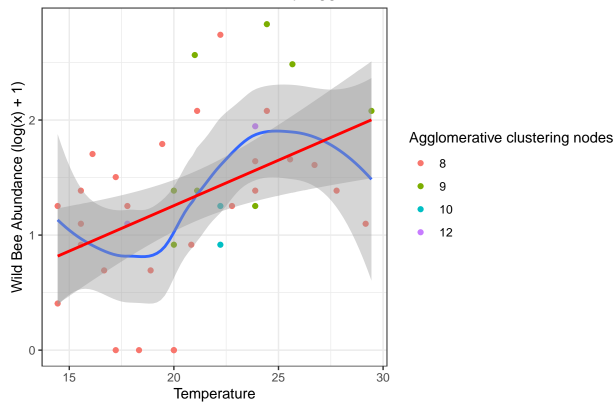
Temperature

It is known that temperature affects the speed at which bees fly, as such this will have an effect on bee abundance and possibly richness although more bees might not necessarily mean more species are observed. From the previous analysis carried out it is known that a $\log(x) + 1$ transposition of bee count allows for a "better" model fit and as such in general this will the case for our observations.

```
## Joining, by = "orchard"
wild_bee_abundance_agglom
```

```
## `geom_smooth()` using method = 'loess' and formula 'y ~ x'
```

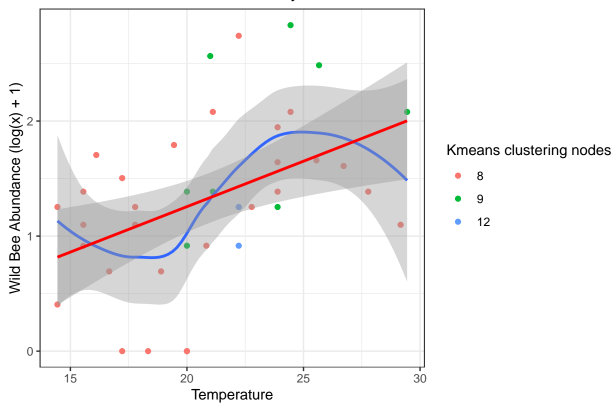
Wild Bee Abundance Coloured by Agglomerature Nodes



```
## Joining, by = "orchard"
wild_bee_abundance_kmeans
```

$geom_smooth()$ using method = 'loess' and formula 'y ~ x'

Wild Bee Abundance Coloured by Kmeans Nodes



```
##Linear model
lm_temp <- lm(log(data_2012$wildAbF + 1) ~ data_2012$temp)
temp_factor <- tidy(lm_temp) %>%
    slice(2) %>%
    select(estimate) %>%
    as_vector()

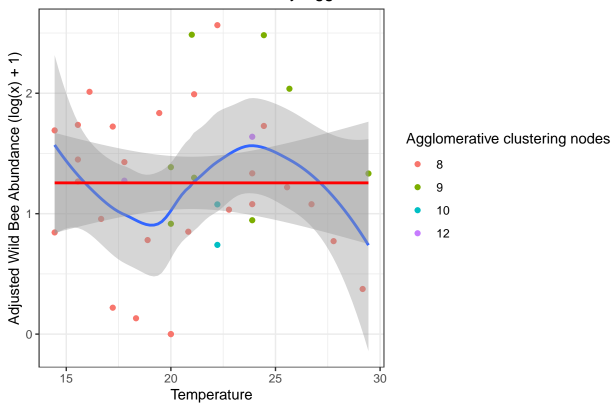
#Adjusting bee value as though temp is always 20
adjusted_data <- data_2012 %>%
    inner_join(clusterings) %>%
    mutate(adjusted_bees = (((20 - temp) * temp_factor) + log(wildAbF + 1)))
```

```
## Joining, by = "orchard"
```

```
wild_bee_abundance_agglom_adjusted <- adjusted_data %>%
    ggplot(aes(x = temp, y = adjusted_bees)) +
    geom_point(aes(colour = as_factor(node))) +
    geom_smooth() +
    geom_smooth(method = "lm", colour = "red") +
    labs(x = "Temperature", y = "Adjusted Wild Bee Abundance (log(x) + 1)",
        title = "Wild Bee Abundance Coloured by Agglomerature Nodes",
        colour = "Agglomerative clustering nodes") +
    theme_bw()
wild_bee_abundance_agglom_adjusted
```

`geom_smooth()` using method = 'loess' and formula 'y ~ x'

Wild Bee Abundance Coloured by Agglomerature Nodes



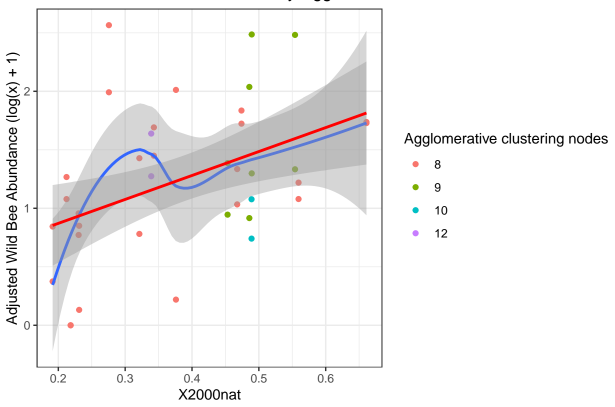
X2000nat

Based on previous research undertaken in the white paper it is known that the X2000nat variable has an effect on the the bee count so it also necessary to adjust for this. Wher Again the value will be adjust as though the X2000nat variable is constant. In this case the mean will be chosen as the constant value and the bee count will be adjusted for this in the same way as above.

```
wild_bee_abundance_agglom_x2000 <- adjusted_data %>%
    ggplot(aes(x = X2000nat, y = adjusted_bees)) +
    geom_point(aes(colour = as_factor(node))) +
    geom_smooth() +
    geom_smooth(method = "lm", colour = "red") +
    labs(x = "X2000nat", y = "Adjusted Wild Bee Abundance (log(x) + 1)",
        title = "Wild Bee Abundance Coloured by Agglomerature Nodes",
        colour = "Agglomerative clustering nodes") +
    theme_bw()
    wild_bee_abundance_agglom_x2000
```

`geom_smooth()` using method = 'loess' and formula 'y ~ x'

Wild Bee Abundance Coloured by Agglomerature Nodes



```
#Outcome is 0.388
mean_x2000 <- adjusted_data %>%
    summarise(mean_x2000 = mean(X2000nat)) %>%
    as_vector()

lm_X2000nat <- lm(adjusted_data$adjusted_bees ~ adjusted_data$X2000nat)
X2000_factor <- tidy(lm_X2000nat) %>%
    slice(2) %>%
    select(estimate) %>%
    as_vector()

#Adjusting the data for this variable

adjusted_data <- adjusted_data %>%
    mutate(adjusted_bees = ((mean_x2000 - X2000nat) * X2000_factor) + adjusted_bees)
```

So if you look at them individually, it can only be adjusted for one. So need to combine a linear model of all affecting factors to best adjust at the end.

Analysising other potential confounders

This section will look at finding other parameters to include in the bee variable, dataset used at this point is the original just with the bee variable considered going through a $\log(x + 1)$ transformation. The only bee variable to be considered at this point is the wild bee abundance, in future more could be considered in this

way or it would be expected to be similar for all bee variables and I could just use the variables decided upon in this way for all the bee variables considered.

Local Diversity

The aim here is to whether adding local diversity as a adjustment factor is going to be beneficial or not. This will be achieved by running a F test to see whether the variances are equal, a shapiro test to confirm the bee data can be assumed to be normal. Note, independence assumption is not validated, this is due more than 1 result being used from each orchard, I decided to include these as I thought the extra power gained from twice the observations was worth the penalty of this assumption.

```
#When adding temperature in other variables have an effect
logged data <- data 2012 %>%
  inner join(clusterings) %>%
  mutate(adjusted_bees = (((20 - temp) * temp_factor) + log(wildAbF + 1)))
## Joining, by = "orchard"
#Just logged
logged_data <- data_2012 %>%
  inner_join(clusterings) %>%
  mutate(adjusted_bees = log(wildAbF + 1))
## Joining, by = "orchard"
#X2000nat adjustment
logged data <- data 2012 %>%
  inner join(clusterings) %>%
  mutate(adjusted_bees = log(wildAbF + 1))%>%
  mutate(adjusted_bees = ((mean_x2000 - X2000nat) * X2000_factor) + adjusted_bees)
## Joining, by = "orchard"
#Testing underlying normal data assumption, again demonstrating why the transformation is neccessary:
#Not normal
shapiro.test(logged_data$wildAbF)
##
##
   Shapiro-Wilk normality test
##
## data: logged_data$wildAbF
## W = 0.79764, p-value = 9.361e-06
#After transformation -> normal
shapiro.test(logged_data$adjusted_bees)
##
##
   Shapiro-Wilk normality test
##
## data: logged_data$adjusted_bees
## W = 0.97681, p-value = 0.6046
#Checking if the two subsets have the same variance have the same variance
\#Outcome shows that they can be assumed to have the same variance as p.value > 0,05
simple <- logged_data %>% filter(local.diversity == 0) %>% select(adjusted_bees) %>% as_vector()
diverse <- logged_data %>% filter(local.diversity == 1) %>% select(adjusted_bees) %>% as_vector()
tidy(var.test(simple, diverse)) %>%
  select(statistic, p.value)
```

```
## Multiple parameters; naming those columns num.df, denom.df
## # A tibble: 1 x 2
     statistic p.value
         <dbl>
##
                 <db1>
## 1
          1.17
                 0.725
#Performing a two sample t-test on the data to see whether they are the same.
#Since HO: Means the same, H1: means are different ~ p.value not significant
#This implies local diversity does NOT have an effect on the original bee counts.
#I thought this was an interesting point and worth noting, same happens with Region.
tidy(t.test(wildAbF ~ local.diversity, logged_data, var.equal = TRUE))
## # A tibble: 1 x 9
    estimate1 estimate2 statistic p.value parameter conf.low conf.high method
##
         <dbl>
                   <dbl>
                             <dbl>
                                      <dbl>
                                                <dbl>
                                                          <dbl>
                                                                    <dbl> <chr>
                                                                     1.91 " Two~
## 1
           3.5
                    4.18
                            -0.532
                                      0.598
                                                          -3.27
## # ... with 1 more variable: alternative <chr>
#However the the adjusted version of bee count does list local diversity
#as a factor which has an effect on the log transformed bee counts
tidy(t.test(adjusted_bees ~ local.diversity, logged_data, var.equal = TRUE))
## # A tibble: 1 x 9
##
     estimate1 estimate2 statistic p.value parameter conf.low conf.high method
                   <dbl>
                             <dbl>
                                      <dbl>
                                                <dbl>
                                                          <dbl>
##
         <dbl>
                                                                    <dbl> <chr>
                                      0.376
                    1.41
                            -0.896
                                                        -0.617
                                                                    0.239 " Two~
## 1
         1.22
                                                   36
## # ... with 1 more variable: alternative <chr>
From This analysis, it can be concluded that local diversity does NOT play a factor in bee count ### Region
Region has 3 possible options - as such an ANOVA test will be used to see whether there is a difference
between the means in wild abundance based on region.
#Checking Anova assumption that the variances are the same
# HO: Variances the same: H1: Atleast one variance is different
leveneTest(adjusted_bees ~ region, data = logged_data)
## Levene's Test for Homogeneity of Variance (center = median)
        Df F value Pr(>F)
## group 2 0.1954 0.8234
\#Since\ pr > 0.05\ we\ can\ assume\ variances\ are\ the\ same
#Running the anova test
anova1 <- aov(adjusted_bees ~ region, data = logged_data)</pre>
summary(anova1)
##
               Df Sum Sq Mean Sq F value Pr(>F)
## region
                2 0.522 0.2611
                                   0.624 0.542
## Residuals
               35 14.652 0.4186
#probablity shows that region is not an affecting factor to wildAbF
#Checking that the following anova assumption is TRUE:
#Residuals of the response variable are normally distributed is NOT true
shapiro.test(residuals(anova1))
```

```
##
## Shapiro-Wilk normality test
##
## data: residuals(anova1)
## W = 0.96856, p-value = 0.3546
```

Day

Looking at the day to see whether that also has an effect, since the data can be paired by day 1 and day 2 a paired t-test will be carried out on this to test whether there is a difference in bee count. When using just the X2000nat variable, then it becomes very close to significant.

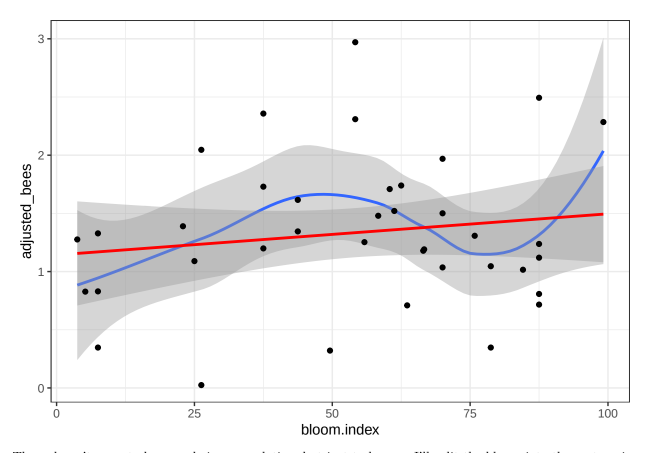
```
day_1 <- logged_data %>% filter(day == 1) %>% select(adjusted_bees) %>% as_vector()
day_2 <- logged_data %>% filter(day == 2) %>% select(adjusted_bees) %>% as_vector()
#This demonstrates that the variances come from the same distribution:
tidy(var.test(day 1, day 2)) %>%
  select(statistic, p.value)
## Multiple parameters; naming those columns num.df, denom.df
## # A tibble: 1 x 2
##
     statistic p.value
##
         <dbl>
                 <dbl>
         0.936
                 0.891
## 1
#Hence we can run a paired t-test with equal variances
tidy(t.test(adjusted_bees ~ day, logged_data, var.equal = TRUE, paired = TRUE))
## # A tibble: 1 x 8
##
     estimate statistic p.value parameter conf.low conf.high method
##
                          <dbl>
        <dbl>
                  <dbl>
                                    <dbl>
                                              <dbl>
                                                        <dbl> <chr>
## 1
       -0.342
                  -2.02 0.0588
                                       18
                                            -0.697
                                                       0.0142 Paire~
## # ... with 1 more variable: alternative <chr>
#Bit surprising but it demonstrates that there is no statistical significance between days
#So combining values to start with is probably appropriate
```

Bloom

First it shall be graphed to see if there is any obvious correlation.

```
logged_data %>%
  ggplot(aes(x = bloom.index, y = adjusted_bees)) +
  geom_smooth() +
  geom_smooth(method = "lm", colour = "red") +
  theme_bw() +
  geom_point()
```

`geom_smooth()` using method = 'loess' and formula 'y ~ x'



There doesn't seem to be any obvious correlation, but just to be sure I'll split the bloom into thee categories and test these with an Anova test.

```
logged_data <- logged_data %>%
          mutate(bloom_category = if_else(bloom.index <= 33, 1,</pre>
                                   if_else(bloom.index <= 66, 2, 3)))</pre>
#Running the anova test
anova1 <- aov(adjusted_bees ~ bloom_category, data = logged_data)</pre>
summary(anova1)
##
                  Df Sum Sq Mean Sq F value Pr(>F)
## bloom_category
                   1
                       0.19 0.1897
                                       0.456 0.504
## Residuals
                  36
                      14.98 0.4162
#As expected there is no evidence to suggest that bloom index (category)
#Has an effect on the bee counts (although only 1 DF in bloom_category thought it should be 2??)
```

Bloom is a continuous variable but for the sakes of deciding whether to include it or not

Final Confounders

Based on the analysis above, the confounders that could be taken into account are: Region and local diversity based on the statistical tests, temperature and X2000nat based on the literature.

```
#Checking the variables to see if any are heavily correlated (binary/categorical ones won't be)
#So checking other two and we get a low correlation so acceptable to use them together
cor(logged_data$temp, logged_data$X2000nat)
```

```
## [1] 0.2065843
f_lm <- lm(adjusted_bees ~ temp + X2000nat + local.diversity + region, data = logged_data)
#Interesting values here seem to suggest that only significant one is X2000nat
summary(f lm)
##
## Call:
## lm(formula = adjusted_bees ~ temp + X2000nat + local.diversity +
       region, data = logged_data)
##
##
## Residuals:
                 1Q
                      Median
##
       Min
                                            Max
## -1.23257 -0.35949 -0.06686 0.23707
                                       1.30560
##
## Coefficients:
##
                  Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                   -0.33682
                              0.61856 -0.545 0.58985
## temp
                   0.08343
                               0.02620
                                        3.184 0.00323 **
## X2000nat
                   0.09182
                              0.79392
                                        0.116 0.90865
## local.diversity 0.20663
                               0.28923
                                        0.714 0.48014
                              0.36305 -1.164 0.25290
## regionLO
                   -0.42271
## regionS
                   -0.08406
                              0.46728 -0.180 0.85837
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.5782 on 32 degrees of freedom
## Multiple R-squared: 0.2949, Adjusted R-squared: 0.1848
## F-statistic: 2.677 on 5 and 32 DF, p-value: 0.03941
```

Notes, from running the adjustments: When you use temperature as an original adjustment, only significant one in the model is X2000nat and when you use X2000nat in the original data only significant one is temp. When you use neither and just apply the linear model on the logged data then both temp and nat are significant -> suggesting that just these two variables would be best to adjust by. This suggest to me that fitting a joint lm using both temp and X2000nat to adjust the data is probably best to avoid overfitting and keep the model as parsimonious as possible. Despite region and local diversity being significant for temp adjustment originally, unlikely to be worth implementing but could be considered to cover model changes section. To take this further, could apply lasso regression on the full dataset to see what that comes up with as tends to have a lower MSE than a standard LM.

```
#Using the logged_data with no adjustments
logged_data <- data_2012 %>%
  inner_join(clusterings) %>%
  mutate(adjusted_bees = log(wildAbF + 1)) %>%
  mutate(adjusted_social = log(socialRichF + 1))

## Joining, by = "orchard"

#Defining linear model with temp and X2000nat as factors
temp_nat_lm <- tidy(lm(adjusted_bees ~ temp + X2000nat, data = logged_data))
#Extracting temp and nat estimate values
temp_factor <- temp_nat_lm %>%
  select(estimate) %>%
  slice(2) %>%
  as_vector()
nat_factor <- temp_nat_lm %>%
```

```
select(estimate) %>%
  slice(3) %>%
  as_vector()
#Now adjustments shall be made for these two variables:
#Temperature will be set to 20 degrees
#X2000 nat will be set to the meean x2000 nat value
temp nat adjusted <- logged data %>%
  mutate(adjusted_bees = (adjusted_bees + ((20 - temp) * temp_factor) + ((mean_x2000 - X2000nat) * X200
#Checking that after the adjustment they are not significant at all in the model
tidy(lm(adjusted_bees ~ temp + X2000nat, data = temp_nat_adjusted))
## # A tibble: 3 x 5
##
     term
                  estimate std.error statistic p.value
                     <dbl>
##
     <chr>
                               <dbl>
                                          <dbl>
                                                  <dbl>
## 1 (Intercept)
                 1.24e+ 0
                              0.549
                                       2.25e+ 0 0.0309
                              0.0249 -3.79e-16 1.000
## 2 temp
                 -9.42e-18
## 3 X2000nat
                  9.12e- 2
                              0.735
                                     1.24e- 1 0.902
#As expected now the bee abundance has been adjusted for these two parameters
The same process can be applied to the variable SocialRichF, the other bee variable which needs to be
considered. Since we know that only temp and nat were statistic on the wildAbF we can go straight to the
LM and test that (May need to check this with Julia, if not just cp previous analysis...)
#Looking at the model ~ only X2000nat significant
lm(adjusted_social ~ temp + X2000nat + local.diversity + region, data = logged_data) %>%
  summary
##
## Call:
## lm(formula = adjusted_social ~ temp + X2000nat + local.diversity +
##
       region, data = logged_data)
##
## Residuals:
##
        Min
                  1Q
                       Median
                                     30
                                             Max
## -0.51626 -0.27605 -0.06653 0.19262 0.72315
##
## Coefficients:
##
                    Estimate Std. Error t value Pr(>|t|)
                               0.381099 -0.753 0.45689
## (Intercept)
                   -0.287012
## temp
                    0.007198
                               0.016144
                                           0.446
                                                 0.65869
## X2000nat
                    1.507687
                               0.489142
                                          3.082 0.00421 **
## local.diversity -0.164463
                               0.178196 -0.923
                                                  0.36295
                   -0.094068
## regionLO
                               0.223680 -0.421 0.67690
## regionS
                    0.165221
                               0.287892
                                          0.574 0.57005
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.3562 on 32 degrees of freedom
## Multiple R-squared: 0.3068, Adjusted R-squared: 0.1985
## F-statistic: 2.833 on 5 and 32 DF, p-value: 0.03153
social_lm<- tidy(lm(adjusted_social ~ X2000nat, data = logged_data))</pre>
```

```
social_nat_factor <- social_lm %>%
   select(estimate) %>%
   slice(2) %>%
   as_vector()

#Adjustment for social added
temp_nat_adjusted <- temp_nat_adjusted %>%
   mutate(adjusted_social = (adjusted_social + ((mean_x2000 - X2000nat) * X2000_factor)))
```

Clustering Summarising

Functions involved with summarising the clusters:

```
#Gets the name of a variable as a string ~ now implemented in clusterise but keeping
get_name <- function(x) {</pre>
  deparse(substitute(x))
#Gets the bee variables that I am summarising for each cluster - makes it easy to change in future
clusterise <- function(bee_data, cluster_data, v_name = deparse(substitute(cluster_data))){</pre>
  inner_join(bee_data, cluster_data) %>%
    summarise(mean_honey_ab = mean(mean_honey_ab),
              mean_wild_ab = mean(mean_wild_ab),
              mean social rich = mean(mean social rich),
              n = n()) \%
   mutate(cluster = v name)
}
#The function to output the summaries of both agglom and kmeans clustering
cluster_summarise <- function(bee_data){</pre>
output <- list()</pre>
output$agglom_bees <- clusterise(bee_data, aggl_node1) %>%
  bind_rows(clusterise(bee_data, aggl_node2)) %>%
  bind_rows(clusterise(bee_data, aggl_node3)) %>%
  bind_rows(clusterise(bee_data, aggl_node4)) %>%
  bind_rows(clusterise(bee_data, aggl_node5)) %>%
  bind_rows(clusterise(bee_data, aggl_node6)) %>%
  bind_rows(clusterise(bee_data, aggl_node7)) %>%
  bind_rows(clusterise(bee_data, aggl_node4 %>% filter(cluster == 1), "agglom_node8")) %>%
  bind_rows(clusterise(bee_data, aggl_node4 %>% filter(cluster == 2), "agglom_node9")) %>%
  bind_rows(clusterise(bee_data, aggl_node5 %>% filter(cluster == 1), "agglom_node10")) %>%
  bind_rows(clusterise(bee_data, aggl_node5 %>% filter(cluster == 2), "agglom_node11")) %>%
  bind_rows(clusterise(bee_data, aggl_node6 %>% filter(cluster == 1), "agglom_node12")) %>%
  bind_rows(clusterise(bee_data, aggl_node6 %>% filter(cluster == 2), "agglom_node13")) %>%
  #Only 1 result for cluster as none in the latter cluster
  #Unable to to bind as no common variables for an empty node15
  bind_rows(clusterise(bee_data, aggl_node7 %% filter(cluster == 1), "agglom_node14"))
output$kmeans_bees <- clusterise(bee_data, kmeans_node1) %>%
  bind_rows(clusterise(bee_data, kmeans_node2)) %>%
  bind_rows(clusterise(bee_data, kmeans_node3)) %>%
```

```
bind_rows(clusterise(bee_data, kmeans_node4)) %>%
  bind_rows(clusterise(bee_data, kmeans_node5)) %>%
  bind_rows(clusterise(bee_data, kmeans_node6)) %>%
  bind_rows(clusterise(bee_data, kmeans_node7)) %>%
  bind_rows(clusterise(bee_data, kmeans_node4 %>% filter(cluster == 1), "kmeans_node8")) %>%
  bind_rows(clusterise(bee_data, kmeans_node4 %>% filter(cluster == 2), "kmeans_node9")) %>%
  bind_rows(clusterise(bee_data, kmeans_node5 %% filter(cluster == 1), "kmeans_node10")) %>%
  bind rows(clusterise(bee data, kmeans node5 %% filter(cluster == 2), "kmeans node11")) %>%
  bind_rows(clusterise(bee_data, kmeans_node6 %>% filter(cluster == 1), "kmeans_node12")) %>%
  bind_rows(clusterise(bee_data, kmeans_node6 %>% filter(cluster == 2), "kmeans_node13")) %>%
  bind_rows(clusterise(bee_data, kmeans_node7 %>% filter(cluster == 1), "kmeans_node14")) %>%
  bind_rows(clusterise(bee_data, kmeans_node7 %>% filter(cluster == 2), "kmeans_node15"))
#Return a list containing two tibbles one for each type of clustering algorithm
output
}
Original summarising without any variable correction
#original Bee data (logged has no changes to it)
bee_values <- logged_data %>%
  group_by(orchard) %>%
  summarise(mean_honey_ab = mean(apisAb),
            mean_wild_ab = mean(adjusted_bees),
            mean_social_rich = mean(adjusted_social)
original_bee_summary <- cluster_summarise(bee_data = bee_values)</pre>
original_bee_summary$agglom_bees
## # A tibble: 14 x 5
      mean_honey_ab mean_wild_ab mean_social_rich
##
                                                       n cluster
                                             <dbl> <int> <chr>
##
              <dbl>
                           <dbl>
## 1
               6.56
                            1.33
                                             0.379
                                                      19 aggl_node1
## 2
               6.44
                            1.32
                                             0.389
                                                      18 aggl_node2
## 3
               8.75
                            1.52
                                             0.203
                                                      1 aggl_node3
## 4
               6.70
                            1.34
                                             0.391
                                                      17 aggl_node4
## 5
                            1.08
                                             0.347
                                                       1 aggl_node5
## 6
               8.75
                                             0.203
                            1.52
                                                       1 aggl_node6
## 7
             {\tt NaN}
                                                       0 aggl_node7
                          NaN
## 8
               6.01
                                             0.358
                            1.17
                                                      13 agglom_node8
## 9
               8.94
                            1.86
                                             0.499
                                                       4 agglom_node9
## 10
               2
                            1.08
                                             0.347
                                                       1 agglom_node10
## 11
                          NaN
                                           NaN
                                                       0 agglom_node11
             NaN
## 12
               8.75
                            1.52
                                             0.203
                                                       1 agglom_node12
## 13
                                                       0 agglom node13
             NaN
                          NaN
                                          NaN
## 14
                          NaN
                                          NaN
             NaN
                                                       0 agglom_node14
#Kmeans
original_bee_summary$kmeans_bees
## # A tibble: 15 x 5
```

n cluster

mean_honey_ab mean_wild_ab mean_social_rich

##		<dbl> <db< th=""><th>l> <dbl></dbl></th><th><int></int></th><th><chr></chr></th></db<></dbl>	l> <dbl></dbl>	<int></int>	<chr></chr>
##	1	6.56 1.	33 0.379	19	kmeans_node1
##	2	6.81 1.	35 0.381	18	kmeans_node2
##	3	2 1.	0.347	1	kmeans_node3
##	4	6.81 1.	35 0.381	18	kmeans_node4
##	5	NaN NaN	NaN	0	kmeans_node5
##	6	2 1.	0.347	1	kmeans_node6
##	7	NaN NaN	NaN	0	kmeans_node7
##	8	6.21 1.	20 0.347	14	kmeans_node8
##	9	8.94 1.	36 0.499	4	kmeans_node9
##	10	NaN NaN	NaN	0	kmeans_node10
##	11	NaN NaN	NaN	0	kmeans_node11
##	12	2 1.	0.347	1	$kmeans_node12$
##	13	NaN NaN	NaN	0	kmeans_node13
##	14	NaN NaN	NaN	0	kmeans_node14
##	15	NaN NaN	NaN	0	kmeans_node15

##Original summarising of non-markov approach maximum cluster

```
##
     eiqB11F.pre eiqB11I.pre eiqB11F.blm eiqB11I.blm eiqB11T.blm eiqB11F.pos
## 1
      0.56294059
                  -0.3418658 -0.19022893 -0.30585457 -0.30620065
## 2
      0.88788914
                                                                0.01058593
                  -0.3418658 -0.19022893 -0.30585457 -0.30620065
## 3
      0.88788914
                                                                0.01058593
## 4
      0.88788914
                  -0.3418658 -0.19022893 -0.30585457 -0.30620065
                                                                0.01058593
## 5
     -1.21724288
                  -0.3418658 -0.47141495
                                         0.25463750 -0.26826632
                                                                0.05673110
## 6
     -1.78818873
                  -0.3418658 -0.71087017
                                         2.78888695 0.36494510 -0.33951546
## 7
      0.92973857
                   1.1014288
                             1.78834024
                                        0.06780681 -0.26972533
                                                                2.88563060
## 8
      0.01045773
                  -0.3418658
                            2.26526351 -0.36201840 0.01478210
## 9
     -0.28688425
                   ## 10 -0.05899826
                  -0.3418658 -1.93232366 -1.17238220 -0.30620065
                                                                0.53623960
                            0.15454685 -0.81247522 -0.29744657 -0.54716872
## 11
      1.50859712
                  -0.1356809
      1.44670216
                  -0.3068769
                             0.15156608 -0.81247522 -0.30167771 -0.56622868
                  -0.3418658 -0.18559218 -0.44683724 -0.30124000 -0.62240541
## 13 -0.33664640
## 14 -1.63608890
                  -0.3418658
                             0.37048710 0.12282444 -0.30620065 -0.53312454
                   3.8693055 -1.36167172 -0.74943419 -0.30620065 -1.18316953
## 15 -1.58069994
## 16 -0.14885146
                  -0.3418658 0.30623494 -0.51446307 4.07083681 -0.08621861
                  -0.3418658 -0.85792151 1.91089732 -0.28285645 -0.80197030
## 17 -0.08097801
                  -0.3418658 -0.15710926 -0.07432206 -0.05816852 -0.72071468
## 18 -0.02629240
##
                 -0.3418658
                            1.45747473 -0.96491990 -0.30620065
  19
     0.03876764
                                                               1.85488796
##
     eiqB11I.pos eiqB11T.pos cluster
##
  1
      0.07606715 -0.29236499
##
  2
      1.78589159 -0.29236499
                                  1
## 3
      1.78589159 -0.29236499
                                  1
## 4
      1.78589159 -0.29236499
                                  1
## 5
     -0.67656322 -0.29236499
                                  1
## 6
      0.61131654 -0.29236499
                                  1
## 7
      1.48904351 -0.23224030
                                  1
     -0.90112187 -0.29236499
## 8
                                  1
     -0.10337469 -0.17211561
                                  1
## 10 -1.07800026 -0.29236499
                                  1
## 11 -0.07517669 -0.16918270
                                  1
## 12 -0.07517669 0.04491986
                                  1
## 13 -0.78166487 -0.20291118
                                  1
## 14 -1.05339109 -0.29236499
                                  1
```

```
## 15 -0.68425358 -0.29236499
## 16 -0.61093877 4.10700262
                                     2
## 17 -0.65349212 -0.29236499
                                     1
## 18 -0.28179116 0.03025530
                                     1
## 19 -0.55915698 -0.18971308
                                     1
#Clustering summaries based on non-markov approach
non_markov_clusters <- logged_data %>%
  group_by(orchard) %>%
  summarise(mean_honey_ab = mean(apisAb),
            mean_wild_ab = mean(adjusted_bees),
            mean_social_rich = mean(adjusted_social)) %>%
  mutate(cluster = maximum_cluster$cluster) %>%
  group_by(cluster) %>%
  summarise(mean_honey_ab = mean(mean_honey_ab),
            mean_wild_ab = mean(mean_wild_ab),
            mean_social_rich = mean(mean_social_rich),
            n = n()
After Adjustment:
adjusted_bee_values <- temp_nat_adjusted %>%
  group_by(orchard) %>%
  summarise(mean_honey_ab = mean(apisAb),
            mean_wild_ab = mean(adjusted_bees),
            mean_social_rich = mean(adjusted_social)
adjusted_bee_summary <- cluster_summarise(bee_data = adjusted_bee_values)</pre>
#Agglom
adjusted_bee_summary$agglom_bees
## # A tibble: 14 x 5
##
      mean_honey_ab mean_wild_ab mean_social_rich
                                                       n cluster
##
              <dbl>
                           <dbl>
                                             <dbl> <int> <chr>
## 1
               6.56
                           1.27
                                             0.379
                                                      19 aggl_node1
## 2
               6.44
                           1.25
                                             0.383
                                                      18 aggl_node2
## 3
               8.75
                           1.57
                                             0.303
                                                      1 aggl_node3
## 4
               6.70
                           1.28
                                             0.397
                                                      17 aggl_node4
## 5
                           0.735
               2
                                             0.140
                                                       1 aggl_node5
## 6
               8.75
                           1.57
                                             0.303
                                                       1 aggl_node6
## 7
                                                       0 aggl_node7
             {\tt NaN}
                         NaN
                                          NaN
## 8
               6.01
                           1.24
                                             0.434
                                                      13 agglom_node8
## 9
               8.94
                                             0.278
                           1.44
                                                       4 agglom_node9
                           0.735
                                             0.140
## 10
               2
                                                       1 agglom_node10
## 11
             NaN
                         {\tt NaN}
                                           {\tt NaN}
                                                       0 agglom_node11
## 12
               8.75
                          1.57
                                             0.303
                                                       1 agglom_node12
## 13
             NaN
                         NaN
                                           NaN
                                                       0 agglom_node13
## 14
             NaN
                         NaN
                                           NaN
                                                       0 agglom_node14
#Kmeans
adjusted_bee_summary$kmeans_bees
## # A tibble: 15 x 5
##
      mean_honey_ab mean_wild_ab mean_social_rich
                                                       n cluster
```

```
##
                <dbl>
                              <dbl>
                                                  <dbl> <int> <chr>
##
                 6.56
                              1.27
    1
                                                  0.379
                                                            19 kmeans_node1
##
   2
                6.81
                              1.30
                                                  0.392
                                                            18 kmeans node2
##
   3
                2
                              0.735
                                                  0.140
                                                             1 kmeans_node3
##
    4
                6.81
                              1.30
                                                  0.392
                                                            18 kmeans node4
   5
                                                             0 kmeans node5
##
                                               \mathtt{NaN}
              NaN
                            NaN
                                                             1 kmeans node6
##
   6
                2
                              0.735
                                                 0.140
    7
##
              NaN
                            NaN
                                               {\tt NaN}
                                                             0 kmeans node7
##
    8
                6.21
                              1.26
                                                  0.425
                                                            14 kmeans node8
                                                 0.278
   9
##
                8.94
                              1.44
                                                             4 kmeans_node9
## 10
              NaN
                            NaN
                                               NaN
                                                             0 kmeans_node10
                                                             0 kmeans_node11
## 11
              {\tt NaN}
                            \mathtt{NaN}
                                               \mathtt{NaN}
## 12
                2
                              0.735
                                                 0.140
                                                             1 kmeans_node12
## 13
              NaN
                            NaN
                                               NaN
                                                             0 kmeans_node13
## 14
              NaN
                                                             0 kmeans_node14
                            NaN
                                               NaN
## 15
              NaN
                            NaN
                                               NaN
                                                             0 kmeans_node15
```

```
##Non-Markov
##Maximum
#Clustering summaries based on non-markov approach
maximum_non_markov_clusters <- temp_nat_adjusted %>%
  group_by(orchard) %>%
  summarise(mean_honey_ab = mean(apisAb),
            mean_wild_ab = mean(adjusted_bees),
            mean_social_rich = mean(adjusted_social)) %>%
  mutate(cluster = maximum_cluster$cluster) %>%
  group by(cluster) %>%
  summarise(mean_honey_ab = mean(mean_honey_ab),
            mean_wild_ab = mean(mean_wild_ab),
            mean_social_rich = mean(mean_social_rich),
            n=n())
##euclidean:
euclidean_non_markov_clusters <- temp_nat_adjusted %>%
  group_by(orchard) %>%
  summarise(mean honey ab = mean(apisAb),
            mean_wild_ab = mean(adjusted_bees),
            mean_social_rich = mean(adjusted_social)) %>%
  mutate(cluster = euclidean cluster$cluster) %>%
  group by(cluster) %>%
  summarise(mean_honey_ab = mean(mean_honey_ab),
            mean_wild_ab = mean(mean_wild_ab),
            mean_social_rich = mean(mean_social_rich),
            n=n()
```

Although honey bee abundance is one of the summarised variables, this is likely to be related directly to the hive acr variable.

Comparing cluster values before and after adjustments, for now just looking at the agglomerative node clustering selection as it is, in my opinion, better than Kmeans.

```
##agglom
adjusted_bee_summary$agglom_bees %>%
select(mean_wild_ab, mean_social_rich ,cluster) %>%
rename(adjusted = mean_wild_ab) %>%
rename(adjusted_social = mean_social_rich) %>%
```

```
inner_join(original_bee_summary$agglom_bees %>%
               select(mean_wild_ab, mean_social_rich, cluster)) %>%
  select(cluster,mean_wild_ab, adjusted, mean_social_rich, adjusted_social)
## Joining, by = "cluster"
## # A tibble: 14 x 5
##
                    mean_wild_ab adjusted mean_social_rich adjusted_social
      cluster
##
      <chr>
                            <dbl>
                                      <dbl>
                                                       <dbl>
                                                                        <dbl>
                             1.33
                                     1.27
                                                                        0.379
## 1 aggl node1
                                                       0.379
                                                                        0.383
## 2 aggl_node2
                             1.32
                                     1.25
                                                       0.389
                             1.52
                                                                        0.303
## 3 aggl_node3
                                     1.57
                                                       0.203
## 4 aggl_node4
                             1.34
                                     1.28
                                                       0.391
                                                                        0.397
## 5 aggl_node5
                             1.08
                                     0.735
                                                       0.347
                                                                        0.140
## 6 aggl_node6
                             1.52
                                     1.57
                                                       0.203
                                                                        0.303
## 7 aggl_node7
                           {\tt NaN}
                                   NaN
                                                     NaN
                                                                      NaN
## 8 agglom_node8
                                                       0.358
                                                                        0.434
                             1.17
                                     1.24
## 9 agglom_node9
                             1.86
                                     1.44
                                                       0.499
                                                                        0.278
                             1.08
                                     0.735
                                                       0.347
                                                                        0.140
## 10 agglom_node10
## 11 agglom_node11
                           NaN
                                   NaN
                                                     NaN
                                                                      NaN
## 12 agglom_node12
                             1.52
                                                       0.203
                                                                        0.303
                                     1.57
## 13 agglom_node13
                           NaN
                                   NaN
                                                     NaN
                                                                      NaN
## 14 agglom_node14
                           NaN
                                                                      NaN
                                   NaN
                                                     NaN
```

I guess potentially can see two paths one with a mean slightly lower than the other if you think of the structure.

Looking at clusters

The aim of this section is to look at the clusterings from a low, medium, high point of view...

```
#Non_markov clustering summary maximum
maximum_final_standardised <- maximum_cluster %>%
  group by(cluster) %>%
  summarise_all(funs(mean)) %>%
  inner_join(maximum_non_markov_clusters)
## Joining, by = "cluster"
#Non_markov clustering summary maximum
euclidean_final_standardised <- euclidean_cluster %>%
  group_by(cluster) %>%
  summarise_all(funs(mean)) %>%
  inner_join(euclidean_non_markov_clusters)
## Joining, by = "cluster"
#Cluster 1 -> Low insecticide, High Fungicide, low thinner
#Cluster 2 -> High insecticide, Low Fungicide, low (no) thinner
#Cluster 3 -> middle insecticide, High Fungicide, low thinner
#Cluster 4 -> Low insecticide, High Fungicide, High thinner
#Agglom clustering summary (end stage)
bee_summary_agglom <- adjusted_bee_summary$agglom_bees %>%
 slice(8:14) %>%
```

```
select(-cluster) %>%
  mutate(cluster = c(1,2,3,4,5,6,7))
pesticide_summary_agglom <- aggl_c(aggl_node2, ".pos", 1) %>%
  bind_rows(aggl_c(aggl_node2, ".pos", 2) %>%
  mutate(cluster = if_else(cluster == 1, 3, 4))) %>%
  bind_rows(aggl_c(aggl_node3, ".pos", 1)%>%
  mutate(cluster = if_else(cluster == 1, 5, 6))) %>%
  bind_rows(aggl_c(aggl_node3, ".pos", 2)%>%
  mutate(cluster = if_else(cluster ==1, 7, 8))) %>%
  group_by(cluster) %>%
  summarise_all(funs(mean))
##
agglom_summary_standardised <- bee_summary_agglom %>%
  inner_join(pesticide_summary_agglom) %>%
  select(-orchard)
## Joining, by = "cluster"
 # select(-orchard) %>%
  # #1 for high, O for low, just using bloom
  \# mutate(pest\_rating = c(0,0,1,1,0,0,1)) \%
  # mutate(insect_rating = c(1,0,1,0,0,1,1)) %>%
  # mutate(thinner\_rating = c(1,0,0,1,0,1,0))
bee_summary_kmeans <- adjusted_bee_summary$kmeans_bees %>%
  slice(8:14) %>%
  select(-cluster) %>%
 mutate(cluster = c(1,2,3,4,5,6,7))
pesticide_summary_kmeans <- aggl_c(kmeans_node2, ".pos", 1) %>%
  bind_rows(aggl_c(kmeans_node2, ".pos", 2) %>%
  mutate(cluster = if_else(cluster == 1, 3, 4))) %>%
  bind_rows(aggl_c(kmeans_node3, ".pos", 1)%>%
  mutate(cluster = if_else(cluster == 1, 5, 6))) %>%
  bind_rows(aggl_c(kmeans_node3, ".pos", 2)%>%
  mutate(cluster = if_else(cluster ==1, 7, 8))) %>%
  group_by(cluster) %>%
  summarise_all(funs(mean))
kmeans summary standardised <- bee summary kmeans %>%
  inner_join(pesticide_summary_kmeans) %>%
 select(-orchard)
## Joining, by = "cluster"
```

```
#write.csv(agglom_summary, "agglom_summary.csv")
#Overall
#HLH
#HHL
#HHL
#HLH
#LLL
#LHH
#LLL
#Just Bloom
#LHH
#LLL
#HHL
#HLH
#LLL
#LHH
#HHL
##Just bloom and just fungicide and insecticide
```

Looking at the difference between standardised data and non-standardised data: This section is non-standardised

write up explanation of aggregation before vs after, conditional probabilities along with the tree would be a good idea This section is going cover two step clustering based on various clustering methods for comparison. Visit data will be combined into one bee result for each orchard, note in principle this is fundementally flawed as the data does depend on previous years and the assumption that independent of the past is false as pesticide last year affects bee population this year as would be expected. In these scenarios each implementation will have a maximum of 8 clusters. Two after pre-bloom, 4 after and 8 after post bloom.

```
load("data")
markov data <- data 2012 %>%
  select(ends_with(".pre"), ends_with(".blm"), ends_with(".pos"), orchard) %>%
  unique()
#function to generate the agglomaerative clustering
aggl <- function(data){</pre>
  agnes(dist(data, method = "euclidian"),
                    diss=TRUE, method = "ward")
}
#function to reduce copy pasting for each node
aggl_c <- function(data, ends, c_num){</pre>
  #define temp dataset of this stage by cluster
    temp <- data %>%
    filter(cluster == c_num) %>%
    select(ends_with(ends))
#If statement as if 1 element in dataset it'll error
if(nrow(temp) > 1){
  temp$cluster <- cutree(aggl(temp) , k = 2)</pre>
```

```
output <- data %>%
   filter(cluster == c_num) %>%
   select(-cluster) %>%
   mutate(cluster = temp$cluster)
   #Else statement for if 1 element in dataset
    #If only 1 element in the dataset -> set the cluster number to 1
   output <- data %>%
      filter(cluster == c_num) %>%
      select(-cluster) %>%
      mutate(cluster = 1)
 }
  output
####Agglomerative heirachicale clustering
#pre-bloom step
temp <- markov_data %>%
  select(ends_with(".pre"))
temp$cluster <- cutree(aggl(temp) , k = 2)</pre>
aggl_node1 <- markov_data %>%
 mutate(cluster = temp$cluster)
#during bloom step 1
aggl_node2 <- aggl_c(aggl_node1, ".blm", 1)
#during bloom step 2
aggl_node3 <- aggl_c(aggl_node1, ".blm", 2)</pre>
#post bloom step 1
aggl_node4 <- aggl_c(aggl_node2, ".pos", 1)</pre>
#post bloom step 2
aggl_node5 <- aggl_c(aggl_node2, ".pos", 2)</pre>
#post bloom step 3
aggl_node6 <- aggl_c(aggl_node3, ".pos", 1)</pre>
#post bloom step 4
#Note this node only has 1 element in it so it is automatically set to 1
aggl_node7 <- aggl_c(aggl_node3, ".pos", 2)</pre>
orchard_node_agglom <- tibble(orchard = aggl_node4 %% filter(cluster == 1) %% select(orchard) %% as_
  bind_rows(tibble(orchard = aggl_node4 %>% filter(cluster == 2) %>% select(orchard) %>% as_vector, nod
  bind_rows(tibble(orchard = aggl_node5 %>% filter(cluster == 1) %>% select(orchard) %>% as_vector, nod
  bind_rows(tibble(orchard = aggl_node5 %>% filter(cluster == 2) %>% select(orchard) %>% as_vector, nod
  bind_rows(tibble(orchard = aggl_node6 %>% filter(cluster == 1) %>% select(orchard) %>% as_vector, nod
  bind_rows(tibble(orchard = aggl_node6 %>% filter(cluster == 2) %>% select(orchard) %>% as_vector, nod
  bind_rows(tibble(orchard = aggl_node7 %>% filter(cluster == 1) %>% select(orchard) %>% as_vector, nod
```

kmeans approach: Even running the 1001 times it is different everytime: store output later on but run it like 1million times at each stage.

```
#function to generate the kmeans clustering
k_clustering <- function(data, n = 1001){</pre>
#creating the dataset to then calculate the optimal cluster from
  for(i in 1:n){
  if(i == 1){
     k_list <- tibble(kmeans(data, 2)$cluster)</pre>
 }else{
     k_list <- bind_cols(k_list, tibble(kmeans(data, 2)$cluster))</pre>
 }
  apply(k_list[ ,1:length(k_list)], 1, mfv1) %>%
    enframe(value = "cluster") %>%
    select(cluster)
}
#pre-bloom step
#Getting the most common clustering
kmeans_c <- function(data, ends, c_num){</pre>
  data <- data %>%
    filter(cluster == c_num) %>%
    select(-cluster)
  if(nrow(data) > 2){
    temp <- data %>%
    select(ends_with(ends))
  cluster <- k_clustering(temp, 2)</pre>
 output <- bind_cols(data, cluster)</pre>
 }else{
    #Not more than 2 rows and get the error message as:
    #number of cluster centres must lie between 1 and nrow(x)
    output <- data %>%
      mutate(cluster = 1)
 }
output
temp <- k_clustering(markov_data %>% select(-orchard))
kmeans_node1 <- markov_data %>%
 bind_cols(temp)
#during bloom step 1
kmeans_node2 <- kmeans_c(kmeans_node1, ".blm", 1)</pre>
#during bloom step 2
```

```
kmeans_node3 <- kmeans_c(kmeans_node1, ".blm", 2)</pre>
#post bloom step 1
kmeans_node4 <- kmeans_c(kmeans_node2, ".pos", 1)</pre>
#post bloom step 2
kmeans_node5 <- kmeans_c(kmeans_node2, ".pos", 2)</pre>
#post bloom step 3
kmeans_node6 <- kmeans_c(kmeans_node3, ".pos", 1)</pre>
#post bloom step 4
kmeans_node7 <- kmeans_c(kmeans_node3, ".pos", 2)</pre>
#End timepoint nodes
orchard_node_kmeans <- tibble(orchard = kmeans_node4 %% filter(cluster == 1) %% select(orchard) %% a
  bind_rows(tibble(orchard = kmeans_node4 %>% filter(cluster == 2) %>% select(orchard) %>% as_vector, n
  bind_rows(tibble(orchard = kmeans_node5 %>% filter(cluster == 1) %>% select(orchard) %>% as_vector, n
  bind_rows(tibble(orchard = kmeans_node5 %>% filter(cluster == 2) %>% select(orchard) %>% as_vector, n
  bind_rows(tibble(orchard = kmeans_node6 %>% filter(cluster == 1) %>% select(orchard) %>% as_vector, n
  bind_rows(tibble(orchard = kmeans_node6 %>% filter(cluster == 2) %>% select(orchard) %>% as_vector, n
  bind_rows(tibble(orchard = kmeans_node7 %>% filter(cluster == 1) %>% select(orchard) %>% as_vector, n
  bind_rows(tibble(orchard = kmeans_node7 %>% filter(cluster == 2) %>% select(orchard) %>% as_vector, n
Just a look at the difference in the final clusters between the two methods
clusterings <- orchard_node_agglom %>%
  arrange(orchard) %>%
  bind_cols(node_km = orchard_node_kmeans %>% arrange(orchard) %>%
              select(node) %>% as_vector)
clusterings
## # A tibble: 19 x 3
##
      orchard node node_km
      <fct> <dbl>
                      <dbl>
##
## 1 A
                  8
                         10
## 2 B
                  9
                         10
## 3 C
                  9
                         10
                  9
## 4 D
                         10
## 5 E
                 12
                         12
## 6 F
                 13
                         13
## 7 G
                          8
                 10
## 8 H.nooil
                 11
                          9
## 9 I
                  8
                          10
## 10 J
                  8
                          9
## 11 K
                  8
                         10
## 12 L
                  8
                         10
## 13 M
                  8
                         10
                 14
## 14 N
                         12
## 15 O.nooil
                 12
                         12
## 16 P
                  8
                         10
## 17 Q
                  8
                          9
                  8
## 18 R
                         10
```

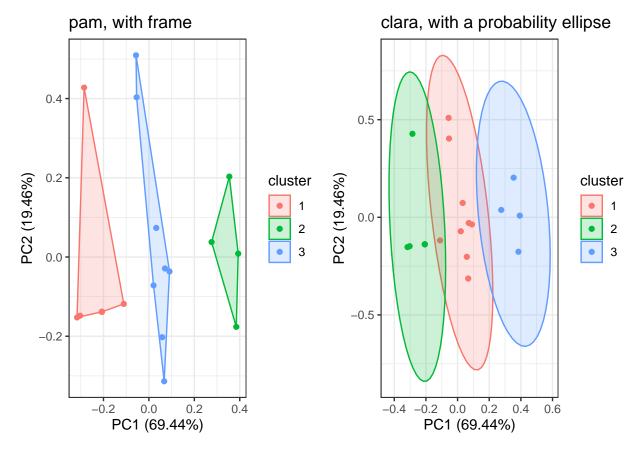
```
## 19 S 11 9
```

Looking at top two PCAs just to see how much variation is absorbed by these

```
p1 <- autoplot(pam(markov_data, 3), frame = TRUE) +
    theme_bw() +
    ggtitle("pam, with frame")

p2 <- autoplot(clara(markov_data, 3), frame = TRUE, frame.type = 'norm') +
    theme_bw() +
    ggtitle("clara, with a probability ellipse")

grid.arrange(p1, p2 ,nrow = 1)</pre>
```



Non-markov structuring

This section will approach looking at the data as one time point with 1 clustering point rather than 3 as above: Method Justification - Maximum linkage function is best for both standardised and unstandardised.

```
maximum_cluster <- data_2012 %>%
   arrange(orchard) %>%
   select(ends_with(".pre"),ends_with(".blm"), ends_with(".pos")) %>%
   filter(row_number() %% 2 == 1)

euclidean_cluster <- maximum_cluster

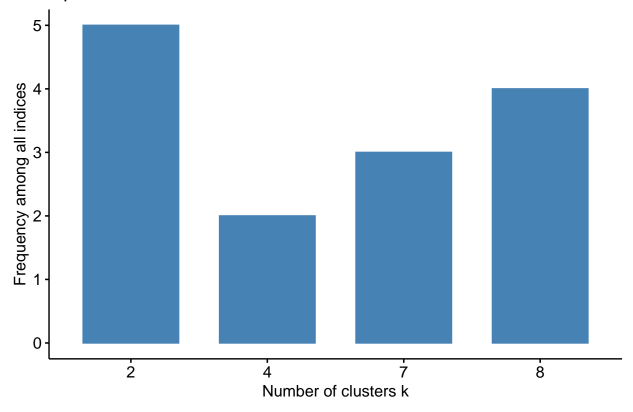
# matrix of methods to compare,</pre>
```

```
m <- c( "average", "single", "complete", "ward")
names(m) <- c( "average", "single", "complete", "ward")</pre>
distances <- c("euclidean", "maximum", "manhattan", "canberra",
               "minkowski")
names(distances) <- c("euclidean", "maximum", "manhattan",</pre>
                       "canberra", "minkowski")
clust_comps <- matrix(nrow = length(distances), ncol = length(m),</pre>
                      dimnames = list(distances,m))
# function to compute coefficient to see which is the best method
ac <- function(distance, linkage) {</pre>
 dista <- dist(maximum_cluster , method = distance)</pre>
  #Agglomerative Nesting form of Hierarchical Clustering
  agnes(dista, method = linkage)$ac
for(i in 1:length(distances)) {
  for(j in 1:length(m)) {
    clust_comps[i,j] <- ac(distances[i], m[j])</pre>
 }
}
#In future or in write up, perhaps change, needs more clustering analysis
#Look at literature in more detail to see other benefits/drawbacks and assumptions
#Then decide on actual best method, for now using euclidian as used that before
clust_comps = clust_comps
Clustering code - Maximum
#NBclust function breaks when using index = all, instead using all the working indexes instead
fviz nbclust(NbClust(maximum cluster, distance = "maximum",
```

* According to the majority rule, the best number of clusters is 2.

#rerun - without %>%mutate segment for non-standardised values

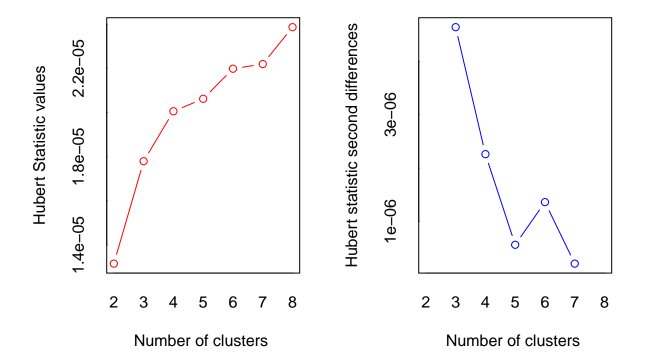
Optimal number of clusters -k = 2



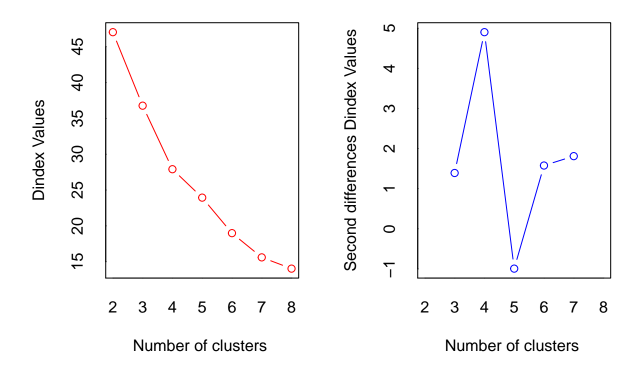
##		eiqB11F.pre	eiqB11I.pre	eiqB11F.blm	eiqB11I.blm	eiqB11T.blm	eiqB11F.pos
##	1	143.09	0.000	58.10	19.04	0.0440	14.48
##	2	161.57	0.000	53.16	9.36	0.0000	31.37
##	3	161.57	0.000	53.16	9.36	0.0000	31.37
##	4	161.57	0.000	53.16	9.36	0.0000	31.37
##	5	41.85	0.000	44.67	14.25	0.0260	32.29
##	6	9.38	0.000	37.44	36.36	0.4600	24.39
##	7	163.95	2.310	112.90	12.62	0.0250	88.69
##	8	111.67	0.000	127.30	8.87	0.2200	53.11
##	9	94.76	0.960	52.27	19.69	0.0140	26.59
##	10	107.72	0.000	0.56	1.80	0.0000	41.85
##	11	196.87	0.330	63.57	4.94	0.0060	20.25
##	12	193.35	0.056	63.48	4.94	0.0031	19.87
##	13	91.93	0.000	53.30	8.13	0.0034	18.75
##	14	18.03	0.000	70.09	13.10	0.0000	20.53
##	15	21.18	6.740	17.79	5.49	0.0000	7.57
##	16	102.61	0.000	68.15	7.54	3.0000	29.44
##	17	106.47	0.000	33.00	28.70	0.0160	15.17

```
## 18
           109.58
                         0.000
                                     54.16
                                                  11.38
                                                             0.1700
                                                                           16.79
## 19
           113.28
                         0.000
                                    102.91
                                                   3.61
                                                             0.0000
                                                                           68.14
##
      eiqB11I.pos eiqB11T.pos cluster
## 1
            24.31
                         0.000
                                     1
## 2
            57.66
                         0.000
                                     1
## 3
            57.66
                         0.000
                                     1
## 4
            57.66
                         0.000
                                     1
                                     2
## 5
             9.63
                         0.000
## 6
            34.75
                         0.000
                                     3
## 7
                                     4
            51.87
                         0.041
## 8
             5.25
                         0.000
                                     5
                                     6
## 9
            20.81
                         0.082
                                     7
## 10
             1.80
                         0.000
                                     8
## 11
            21.36
                         0.084
## 12
            21.36
                         0.230
                                     8
## 13
             7.58
                         0.061
                                     6
## 14
             2.28
                         0.000
                                     2
                                     3
## 15
             9.48
                         0.000
## 16
            10.91
                         3.000
                                     6
## 17
            10.08
                         0.000
                                     6
## 18
            17.33
                         0.220
                                     6
## 19
            11.92
                         0.070
                                     5
#Clustering summaries based on non-markov approach (Before Adjustment)
maximum_non_markov_clusters <- data_2012 %>%
  group by(orchard) %>%
  summarise(mean_honey_ab = mean(apisAb),
            mean_wild_ab = mean(wildAbF),
            mean_social_rich = mean(socialRichF)) %>%
  mutate(cluster = maximum_cluster$cluster) %>%
  group_by(cluster) %>%
  summarise(mean_honey_ab = mean(mean_honey_ab),
            mean_wild_ab = mean(mean_wild_ab),
            mean_social_rich = mean(mean_social_rich),
            n=n()
```

Clustering code - euclidean

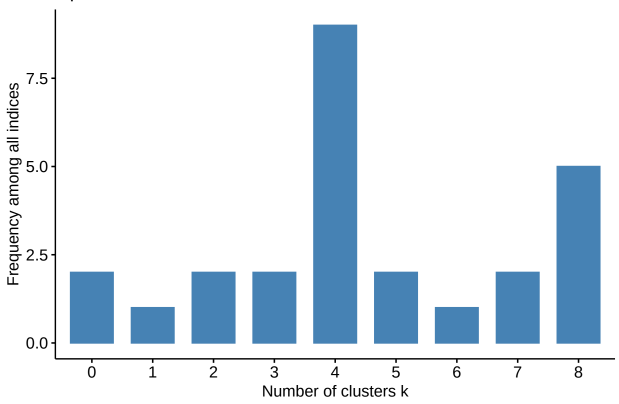


*** : The Hubert index is a graphical method of determining the number of clusters.
In the plot of Hubert index, we seek a significant knee that corresponds to a
significant increase of the value of the measure i.e the significant peak in Hubert
index second differences plot.
##



```
*** : The D index is a graphical method of determining the number of clusters.
                In the plot of D index, we seek a significant knee (the significant peak in Dindex
##
                second differences plot) that corresponds to a significant increase of the value of
##
                the measure.
##
##
## * Among all indices:
## * 2 proposed 2 as the best number of clusters
## * 2 proposed 3 as the best number of clusters
## * 9 proposed 4 as the best number of clusters
## * 2 proposed 5 as the best number of clusters
\#\# * 1 proposed 6 as the best number of clusters
## * 2 proposed 7 as the best number of clusters
## * 5 proposed 8 as the best number of clusters
##
##
                   ***** Conclusion *****
##
## * According to the majority rule, the best number of clusters is 4
##
##
## Among all indices:
## * 2 proposed 0 as the best number of clusters
## * 1 proposed 1 as the best number of clusters
## * 2 proposed 2 as the best number of clusters
```

Optimal number of clusters -k = 4



##		eiqB11F.pre	eiqB11I.pre	$\verb"eiqB11F.blm"$	${\tt eiqB11I.blm}$	${\tt eiqB11T.blm}$	eiqB11F.pos
##	1	143.09	0.000	58.10	19.04	0.0440	14.48
##	2	161.57	0.000	53.16	9.36	0.0000	31.37
##	3	161.57	0.000	53.16	9.36	0.0000	31.37
##	4	161.57	0.000	53.16	9.36	0.0000	31.37
##	5	41.85	0.000	44.67	14.25	0.0260	32.29
##	6	9.38	0.000	37.44	36.36	0.4600	24.39
##	7	163.95	2.310	112.90	12.62	0.0250	88.69

```
## 9
            94.76
                         0.960
                                      52.27
                                                   19.69
                                                               0.0140
                                                                             26.59
                         0.000
## 10
           107.72
                                       0.56
                                                    1.80
                                                               0.0000
                                                                             41.85
                                                    4.94
                                                                             20.25
## 11
           196.87
                         0.330
                                      63.57
                                                               0.0060
## 12
           193.35
                         0.056
                                      63.48
                                                    4.94
                                                               0.0031
                                                                             19.87
## 13
            91.93
                         0.000
                                                               0.0034
                                                                             18.75
                                      53.30
                                                    8.13
## 14
            18.03
                                                   13.10
                                                               0.0000
                                                                             20.53
                         0.000
                                      70.09
## 15
            21.18
                         6.740
                                      17.79
                                                    5.49
                                                               0.0000
                                                                             7.57
## 16
           102.61
                         0.000
                                      68.15
                                                    7.54
                                                               3.0000
                                                                             29.44
## 17
           106.47
                         0.000
                                      33.00
                                                   28.70
                                                               0.0160
                                                                             15.17
## 18
           109.58
                         0.000
                                      54.16
                                                   11.38
                                                               0.1700
                                                                             16.79
## 19
           113.28
                         0.000
                                                    3.61
                                                               0.0000
                                                                             68.14
                                     102.91
##
      eiqB11I.pos eiqB11T.pos cluster
## 1
            24.31
                         0.000
                                      1
## 2
            57.66
                         0.000
                                      1
## 3
            57.66
                         0.000
                                      1
## 4
            57.66
                         0.000
                                      1
                                      2
## 5
             9.63
                         0.000
## 6
            34.75
                         0.000
                                      2
## 7
                                      3
            51.87
                         0.041
## 8
             5.25
                         0.000
                                      3
## 9
            20.81
                         0.082
                                      4
## 10
             1.80
                         0.000
                                      4
## 11
            21.36
                         0.084
                                      1
## 12
            21.36
                         0.230
                                      1
## 13
             7.58
                         0.061
                                      4
## 14
             2.28
                         0.000
                                      2
## 15
                         0.000
                                      2
             9.48
## 16
                                      4
            10.91
                         3.000
## 17
            10.08
                         0.000
                                      4
## 18
            17.33
                         0.220
                                      4
## 19
            11.92
                         0.070
                                      3
#Clustering summaries based on non-markov approach (Before Adjustment)
#Cluster numbers are 6 ,4 ,3 ,6
euclidean_non_markov_clusters <- data_2012 %>%
  group_by(orchard) %>%
  summarise(mean_honey_ab = mean(apisAb),
            mean_wild_ab = mean(wildAbF),
            mean_social_rich = mean(socialRichF)) %>%
  mutate(cluster = euclidean_cluster$cluster) %>%
  group_by(cluster) %>%
  summarise(mean_honey_ab = mean(mean_honey_ab),
            mean wild ab = mean(mean wild ab),
            mean social rich = mean(mean social rich),
            n=n()
euclidean_non_markov_clusters
## # A tibble: 4 x 5
##
     cluster mean_honey_ab mean_wild_ab mean_social_rich
                                                                 n
##
       <int>
                      <dbl>
                                    <dbl>
                                                      <dbl> <int>
## 1
           1
                       8.31
                                     4.62
                                                      0.542
                                                                 6
## 2
           2
                       7.94
                                     5.5
                                                      0.688
                                                                 4
## 3
           3
                       3.75
                                     2.88
                                                      0.625
                                                                 3
## 4
           4
                       5.29
                                     2.60
                                                      0.569
                                                                 6
```

8

111.67

0.000

127.30

8.87

0.2200

53.11

Variable justification

```
bee correlations <- data.frame(matrix(0, nrow = 0, ncol = 0))
for(i in 1:7){
  for(j in 1:7){
  bee correlations[i,j] <- round(cor(data 2012[5+i], data 2012[5 + j]), 2)
  }
}
for(i in 1:7){
names(bee_correlations)[i] <- colnames(data_2012[5 +i])</pre>
}
bee_correlations
     apisAb.1 apisAb.2 apisAb.3 apisAb.4 apisAb.5 apisAb.6 apisAb.7 wildAbF.1
##
## 1
                   1.00
                             1.00
                                       1.00
                                                1.00
                                                          1.00
                                                                    1.00
## 2
         0.29
                   0.29
                             0.29
                                       0.29
                                                0.29
                                                          0.29
                                                                    0.29
                                                                               1.00
## 3
         0.19
                   0.19
                             0.19
                                       0.19
                                                                               0.91
                                                0.19
                                                          0.19
                                                                    0.19
## 4
                             0.25
                                       0.25
                                                                               0.96
         0.25
                   0.25
                                                0.25
                                                          0.25
                                                                    0.25
## 5
         0.13
                   0.13
                             0.13
                                                0.13
                                                          0.13
                                                                               0.92
                                       0.13
                                                                    0.13
                                                                    0.18
## 6
         0.18
                   0.18
                                                0.18
                                                          0.18
                                                                               0.29
                             0.18
                                      0.18
## 7
         0.22
                   0.22
                             0.22
                                       0.22
                                                0.22
                                                          0.22
                                                                    0.22
                                                                               0.31
##
     wildAbF.2 wildAbF.3 wildAbF.4 wildAbF.5 wildAbF.6 wildAbF.7 wildRichF.1
## 1
          0.29
                     0.29
                                0.29
                                           0.29
                                                      0.29
                                                                 0.29
                                                                              0.19
## 2
          1.00
                     1.00
                                1.00
                                           1.00
                                                      1.00
                                                                 1.00
                                                                              0.91
## 3
          0.91
                     0.91
                                0.91
                                           0.91
                                                      0.91
                                                                 0.91
                                                                              1.00
## 4
          0.96
                     0.96
                                0.96
                                           0.96
                                                      0.96
                                                                 0.96
                                                                              0.81
## 5
          0.92
                     0.92
                                0.92
                                           0.92
                                                      0.92
                                                                 0.92
                                                                              0.92
## 6
          0.29
                     0.29
                                0.29
                                           0.29
                                                      0.29
                                                                 0.29
                                                                              0.50
## 7
                                0.31
                                           0.31
                                                      0.31
                                                                 0.31
                                                                              0.53
          0.31
                     0.31
     wildRichF.2 wildRichF.3 wildRichF.4 wildRichF.5 wildRichF.6 wildRichF.7
                          0.19
## 1
             0.19
                                      0.19
                                                    0.19
                                                                 0.19
                                                                              0.19
## 2
             0.91
                          0.91
                                       0.91
                                                    0.91
                                                                 0.91
                                                                              0.91
## 3
             1.00
                          1.00
                                       1.00
                                                    1.00
                                                                 1.00
                                                                              1.00
## 4
                          0.81
                                       0.81
                                                    0.81
             0.81
                                                                 0.81
                                                                              0.81
## 5
                          0.92
                                       0.92
                                                    0.92
                                                                 0.92
             0.92
                                                                              0.92
## 6
             0.50
                          0.50
                                       0.50
                                                    0.50
                                                                 0.50
                                                                              0.50
## 7
             0.53
                          0.53
                                       0.53
                                                    0.53
                                                                 0.53
                                                                              0.53
     solitaryAbF.1 solitaryAbF.2 solitaryAbF.3 solitaryAbF.4 solitaryAbF.5
## 1
               0.25
                              0.25
                                             0.25
                                                            0.25
                                                                           0.25
## 2
                              0.96
                                                            0.96
               0.96
                                             0.96
                                                                           0.96
## 3
                              0.81
               0.81
                                             0.81
                                                            0.81
                                                                           0.81
## 4
               1.00
                              1.00
                                             1.00
                                                            1.00
                                                                            1.00
## 5
               0.92
                              0.92
                                             0.92
                                                            0.92
                                                                           0.92
## 6
               0.02
                              0.02
                                             0.02
                                                            0.02
                                                                            0.02
## 7
               0.05
                              0.05
                                             0.05
                                                            0.05
                                                                            0.05
##
     solitaryAbF.6 solitaryAbF.7 solitaryRichF.1 solitaryRichF.2
## 1
               0.25
                              0.25
                                               0.13
                                                                 0.13
## 2
               0.96
                              0.96
                                               0.92
                                                                 0.92
## 3
               0.81
                              0.81
                                               0.92
                                                                 0.92
## 4
               1.00
                                                                 0.92
                              1.00
                                               0.92
## 5
               0.92
                              0.92
                                               1.00
                                                                 1.00
## 6
               0.02
                              0.02
                                               0.13
                                                                 0.13
## 7
               0.05
                              0.05
                                               0.16
                                                                 0.16
```

##		solitaryRichF.3	solitaryRichF	.4 solitaryR	ichF.5 solita	ryRichF.6
##	1	0.13 0		13	0.13	0.13
##	2	0.92	0.9	92	0.92	0.92
##	3	0.92	0.9	92	0.92	0.92
##	4	0.92	0.9	92	0.92	0.92
##	5	1.00	1.0	00	1.00	1.00
##	6	0.13	0.3	13	0.13	0.13
##	7	0.16	0.3	16	0.16	0.16
##		${\tt solitaryRichF.7}$	socialAbF.1 so	ocialAbF.2 s	ocialAbF.3 so	cialAbF.4
##		0.13	0.18	0.18	0.18	0.18
##	2	0.92	0.29	0.29	0.29	0.29
##	3	0.92	0.50	0.50	0.50	0.50
##	_	0.92	0.02	0.02	0.02	0.02
##	5	1.00	0.13	0.13	0.13	0.13
##		0.13	1.00	1.00	1.00	1.00
##	7	0.16	0.99	0.99	0.99	0.99
##		socialAbF.5 soc				alRichF.2
##		0.18	0.18	0.18	0.22	0.22
##		0.29	0.29	0.29	0.31	0.31
	3	0.50	0.50	0.50	0.53	0.53
##		0.02	0.02	0.02	0.05	0.05
##		0.13	0.13	0.13	0.16	0.16
##	6	1.00	1.00	1.00	0.99	0.99
##	7	0.99	0.99	0.99	1.00	1.00
##		socialRichF.3 so				
##		0.22	0.22	0.22		
##				0.31		
##		0.53	0.53	0.53		
##		1 0.05 0.05		0.05		
##		0.16	0.16	0.16		
	C	0.99	0.99	0.99	0.9	9 0.99
## ##		1.00	1.00	1.00		

This demonstrates that to consider the affects on wild bees, just two variables need to be considered as there is very high correlation between the outcomes. WildAbF accounts a .9+ correlation with all wild variables except for the for the solitary bee variable. Then for this SocialRichF could be used.

Updating the bee values to match the environments

In this particular analysis, the interest is in the affects of pesticides on bee population. As such, it is necessary to correct, as best we can, for extra factors (confounding?) to make the bee values representative.

All these graphs were plotted as part of EDA by the use of lapply to the a basic plotting function. Here the relationships between some of the main extra factors (are they confounding?) will be examined.

Temperature

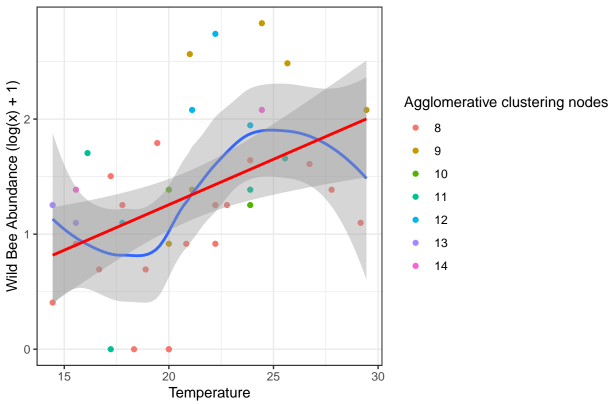
It is known that temperature affects the speed at which bees fly, as such this will have an effect on bee abundance and possibly richness although more bees might not necessarily mean more species are observed. From the previous analysis carried out it is known that a $\log(x) + 1$ transposition of bee count allows for a "better" model fit and as such in general this will the case for our observations.

```
##agglom
wild_bee_abundance_agglom <- data_2012 %>%
```

```
## Joining, by = "orchard"
wild_bee_abundance_agglom
```

`geom_smooth()` using method = 'loess' and formula 'y ~ x'

Wild Bee Abundance Coloured by Agglomerature Nodes

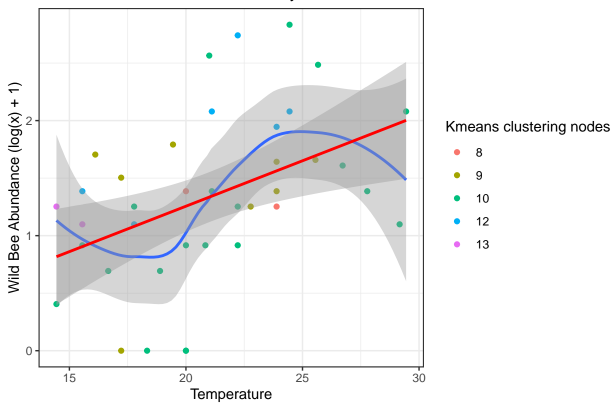


```
##kmeans
wild_bee_abundance_kmeans <- data_2012 %>%
inner_join(orchard_node_kmeans) %>%
ggplot(aes(x = temp, y = log(wildAbF + 1))) +
geom_point(aes(colour = as_factor(node))) +
geom_smooth() +
geom_smooth(method = "lm", colour = "red") +
labs(x = "Temperature", y = "Wild Bee Abundance (log(x) + 1)",
    title = "Wild Bee Abundance Coloured by Kmeans Nodes",
    colour = "Kmeans clustering nodes") +
theme_bw()
```

```
## Joining, by = "orchard"
wild_bee_abundance_kmeans
```

$geom_smooth()$ using method = 'loess' and formula 'y ~ x'

Wild Bee Abundance Coloured by Kmeans Nodes



```
##Linear model
lm_temp <- lm(log(data_2012$wildAbF + 1) ~ data_2012$temp)
temp_factor <- tidy(lm_temp) %>%
    slice(2) %>%
    select(estimate) %>%
    as_vector()

#Adjusting bee value as though temp is always 20
adjusted_data <- data_2012 %>%
    inner_join(clusterings) %>%
    mutate(adjusted_bees = (((20 - temp) * temp_factor) + log(wildAbF + 1)))
```

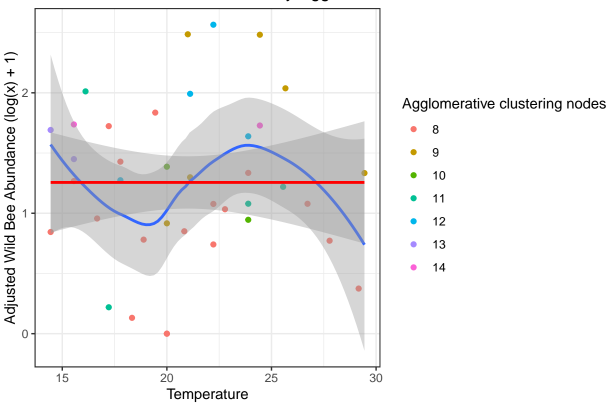
Joining, by = "orchard"

```
wild_bee_abundance_agglom_adjusted <- adjusted_data %>%
    ggplot(aes(x = temp, y = adjusted_bees)) +
    geom_point(aes(colour = as_factor(node))) +
    geom_smooth() +
    geom_smooth(method = "lm", colour = "red") +
    labs(x = "Temperature", y = "Adjusted Wild Bee Abundance (log(x) + 1)",
        title = "Wild Bee Abundance Coloured by Agglomerature Nodes",
        colour = "Agglomerative clustering nodes") +
```

```
theme_bw()
wild_bee_abundance_agglom_adjusted
```

$geom_smooth()$ using method = 'loess' and formula 'y ~ x'

Wild Bee Abundance Coloured by Agglomerature Nodes



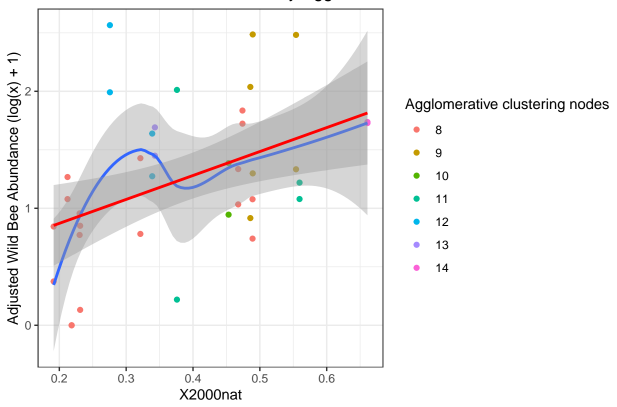
X2000nat

Based on previous research undertaken in the white paper it is known that the X2000nat variable has an effect on the the bee count so it also necessary to adjust for this. Wher Again the value will be adjust as though the X2000nat variable is constant. In this case the mean will be chosen as the constant value and the bee count will be adjusted for this in the same way as above.

```
wild_bee_abundance_agglom_x2000 <- adjusted_data %>%
    ggplot(aes(x = X2000nat, y = adjusted_bees)) +
    geom_point(aes(colour = as_factor(node))) +
    geom_smooth() +
    geom_smooth(method = "lm", colour = "red") +
    labs(x = "X2000nat", y = "Adjusted Wild Bee Abundance (log(x) + 1)",
        title = "Wild Bee Abundance Coloured by Agglomerature Nodes",
        colour = "Agglomerative clustering nodes") +
    theme_bw()
    wild_bee_abundance_agglom_x2000
```

`geom_smooth()` using method = 'loess' and formula 'y ~ x'

Wild Bee Abundance Coloured by Agglomerature Nodes



```
#Outcome is 0.388
mean_x2000 <- adjusted_data %>%
    summarise(mean_x2000 = mean(X2000nat)) %>%
    as_vector()

lm_X2000nat <- lm(adjusted_data$adjusted_bees ~ adjusted_data$X2000nat)
X2000_factor <- tidy(lm_X2000nat) %>%
    slice(2) %>%
    select(estimate) %>%
    select(estimate) %>%
    as_vector()

#Adjusting the data for this variable

adjusted_data <- adjusted_data %>%
    mutate(adjusted_bees = ((mean_x2000 - X2000nat) * X2000_factor) + adjusted_bees)
```

So if you look at them individually, it can only be adjusted for one. So need to combine a linear model of all affecting factors to best adjust at the end.

Analysising other potential confounders

This section will look at finding other parameters to include in the bee variable, dataset used at this point is the original just with the bee variable considered going through a log(x + 1) transformation. The only bee variable to be considered at this point is the wild bee abundance, in future more could be considered in this

way or it would be expected to be similar for all bee variables and I could just use the variables decided upon in this way for all the bee variables considered.

Local Diversity

The aim here is to whether adding local diversity as a adjustment factor is going to be beneficial or not. This will be achieved by running a F test to see whether the variances are equal, a shapiro test to confirm the bee data can be assumed to be normal. Note, independence assumption is not validated, this is due more than 1 result being used from each orchard, I decided to include these as I thought the extra power gained from twice the observations was worth the penalty of this assumption.

```
#When adding temperature in other variables have an effect
logged data <- data 2012 %>%
  inner join(clusterings) %>%
  mutate(adjusted_bees = (((20 - temp) * temp_factor) + log(wildAbF + 1)))
## Joining, by = "orchard"
#Just logged
logged_data <- data_2012 %>%
  inner_join(clusterings) %>%
  mutate(adjusted_bees = log(wildAbF + 1))
## Joining, by = "orchard"
#X2000nat adjustment
logged data <- data 2012 %>%
  inner join(clusterings) %>%
  mutate(adjusted_bees = log(wildAbF + 1))%>%
  mutate(adjusted_bees = ((mean_x2000 - X2000nat) * X2000_factor) + adjusted_bees)
## Joining, by = "orchard"
#Testing underlying normal data assumption, again demonstrating why the transformation is neccessary:
#Not normal
shapiro.test(logged_data$wildAbF)
##
##
   Shapiro-Wilk normality test
##
## data: logged_data$wildAbF
## W = 0.79764, p-value = 9.361e-06
#After transformation -> normal
shapiro.test(logged_data$adjusted_bees)
##
##
   Shapiro-Wilk normality test
##
## data: logged_data$adjusted_bees
## W = 0.97681, p-value = 0.6046
#Checking if the two subsets have the same variance have the same variance
\#Outcome shows that they can be assumed to have the same variance as p.value > 0,05
simple <- logged_data %>% filter(local.diversity == 0) %>% select(adjusted_bees) %>% as_vector()
diverse <- logged_data %>% filter(local.diversity == 1) %>% select(adjusted_bees) %>% as_vector()
tidy(var.test(simple, diverse)) %>%
  select(statistic, p.value)
```

```
## Multiple parameters; naming those columns num.df, denom.df
## # A tibble: 1 x 2
     statistic p.value
##
         <dbl>
                 <db1>
## 1
          1.17
                 0.725
#Performing a two sample t-test on the data to see whether they are the same.
#Since HO: Means the same, H1: means are different ~ p.value not significant
#This implies local diversity does NOT have an effect on the original bee counts.
#I thought this was an interesting point and worth noting, same happens with Region.
tidy(t.test(wildAbF ~ local.diversity, logged_data, var.equal = TRUE))
## # A tibble: 1 x 9
     estimate1 estimate2 statistic p.value parameter conf.low conf.high method
##
         <dbl>
                   <dbl>
                             <dbl>
                                      <dbl>
                                                <dbl>
                                                          <dbl>
                                                                    <dbl> <chr>
                                      0.598
                                                                     1.91 " Two~
## 1
           3.5
                    4.18
                            -0.532
                                                          -3.27
## # ... with 1 more variable: alternative <chr>
#However the the adjusted version of bee count does list local diversity
#as a factor which has an effect on the log transformed bee counts
tidy(t.test(adjusted_bees ~ local.diversity, logged_data, var.equal = TRUE))
## # A tibble: 1 x 9
##
     estimate1 estimate2 statistic p.value parameter conf.low conf.high method
         <dbl>
                   <dbl>
                             <dbl>
                                      <dbl>
                                                <dbl>
                                                          <dbl>
##
                                                                    <dbl> <chr>
          1.22
                                      0.376
                    1.41
                            -0.896
                                                        -0.617
                                                                    0.239 " Two~
## 1
                                                   36
## # ... with 1 more variable: alternative <chr>
From This analysis, it can be concluded that local diversity does NOT play a factor in bee count ### Region
Region has 3 possible options - as such an ANOVA test will be used to see whether there is a difference
between the means in wild abundance based on region.
#Checking Anova assumption that the variances are the same
# HO: Variances the same: H1: Atleast one variance is different
leveneTest(adjusted_bees ~ region, data = logged_data)
## Levene's Test for Homogeneity of Variance (center = median)
        Df F value Pr(>F)
## group 2 0.1954 0.8234
\#Since\ pr > 0.05\ we\ can\ assume\ variances\ are\ the\ same
#Running the anova test
anova1 <- aov(adjusted_bees ~ region, data = logged_data)</pre>
summary(anova1)
##
               Df Sum Sq Mean Sq F value Pr(>F)
## region
                2 0.522 0.2611
                                   0.624 0.542
## Residuals
               35 14.652 0.4186
#probablity shows that region is not an affecting factor to wildAbF
#Checking that the following anova assumption is TRUE:
#Residuals of the response variable are normally distributed is NOT true
shapiro.test(residuals(anova1))
```

```
##
## Shapiro-Wilk normality test
##
## data: residuals(anova1)
## W = 0.96856, p-value = 0.3546
```

Day

Looking at the day to see whether that also has an effect, since the data can be paired by day 1 and day 2 a paired t-test will be carried out on this to test whether there is a difference in bee count. When using just the X2000nat variable, then it becomes very close to significant.

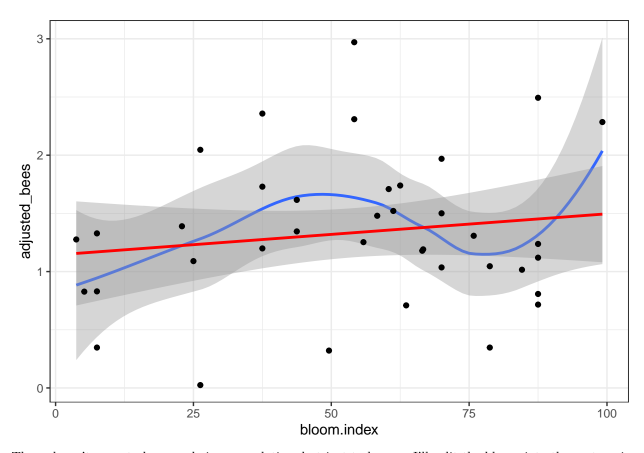
```
day_1 <- logged_data %>% filter(day == 1) %>% select(adjusted_bees) %>% as_vector()
day_2 <- logged_data %>% filter(day == 2) %>% select(adjusted_bees) %>% as_vector()
#This demonstrates that the variances come from the same distribution:
tidy(var.test(day 1, day 2)) %>%
  select(statistic, p.value)
## Multiple parameters; naming those columns num.df, denom.df
## # A tibble: 1 x 2
##
     statistic p.value
##
         <dbl>
                 <dbl>
         0.936
                 0.891
## 1
#Hence we can run a paired t-test with equal variances
tidy(t.test(adjusted_bees ~ day, logged_data, var.equal = TRUE, paired = TRUE))
## # A tibble: 1 x 8
##
     estimate statistic p.value parameter conf.low conf.high method
##
                          <dbl>
        <dbl>
                  <dbl>
                                    <dbl>
                                              <dbl>
                                                        <dbl> <chr>
## 1
       -0.342
                  -2.02 0.0588
                                       18
                                             -0.697
                                                       0.0142 Paire~
## # ... with 1 more variable: alternative <chr>
#Bit surprising but it demonstrates that there is no statistical significance between days
#So combining values to start with is probably appropriate
```

Bloom

First it shall be graphed to see if there is any obvious correlation.

```
logged_data %>%
  ggplot(aes(x = bloom.index, y = adjusted_bees)) +
  geom_smooth() +
  geom_smooth(method = "lm", colour = "red") +
  theme_bw() +
  geom_point()
```

`geom_smooth()` using method = 'loess' and formula 'y ~ x'



There doesn't seem to be any obvious correlation, but just to be sure I'll split the bloom into thee categories and test these with an Anova test.

```
logged_data <- logged_data %>%
          mutate(bloom_category = if_else(bloom.index <= 33, 1,</pre>
                                   if_else(bloom.index <= 66, 2, 3)))</pre>
#Running the anova test
anova1 <- aov(adjusted_bees ~ bloom_category, data = logged_data)</pre>
summary(anova1)
##
                  Df Sum Sq Mean Sq F value Pr(>F)
## bloom_category
                   1
                       0.19 0.1897
                                       0.456 0.504
## Residuals
                  36
                      14.98 0.4162
#As expected there is no evidence to suggest that bloom index (category)
#Has an effect on the bee counts (although only 1 DF in bloom_category thought it should be 2??)
```

Bloom is a continuous variable but for the sakes of deciding whether to include it or not

Final Confounders

Based on the analysis above, the confounders that could be taken into account are: Region and local diversity based on the statistical tests, temperature and X2000nat based on the literature.

```
#Checking the variables to see if any are heavily correlated (binary/categorical ones won't be)
#So checking other two and we get a low correlation so acceptable to use them together
cor(logged_data$temp, logged_data$X2000nat)
```

```
## [1] 0.2065843
f_lm <- lm(adjusted_bees ~ temp + X2000nat + local.diversity + region, data = logged_data)
#Interesting values here seem to suggest that only significant one is X2000nat
summary(f lm)
##
## Call:
## lm(formula = adjusted_bees ~ temp + X2000nat + local.diversity +
       region, data = logged_data)
##
##
## Residuals:
                 1Q
                      Median
##
       Min
                                            Max
## -1.23257 -0.35949 -0.06686 0.23707
                                       1.30560
##
## Coefficients:
##
                  Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                   -0.33682
                              0.61856 -0.545 0.58985
                   0.08343
                                        3.184 0.00323 **
## temp
                               0.02620
## X2000nat
                   0.09182
                               0.79392
                                        0.116 0.90865
## local.diversity 0.20663
                               0.28923
                                        0.714 0.48014
                               0.36305 -1.164 0.25290
## regionLO
                   -0.42271
## regionS
                   -0.08406
                               0.46728 -0.180 0.85837
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.5782 on 32 degrees of freedom
## Multiple R-squared: 0.2949, Adjusted R-squared: 0.1848
```

Notes, from running the adjustments: When you use temperature as an original adjustment, only significant one in the model is X2000nat and when you use X2000nat in the original data only significant one is temp. When you use neither and just apply the linear model on the logged data then both temp and nat are significant -> suggesting that just these two variables would be best to adjust by. This suggest to me that fitting a joint lm using both temp and X2000nat to adjust the data is probably best to avoid overfitting and keep the model as parsimonious as possible. Despite region and local diversity being significant for temp adjustment originally, unlikely to be worth implementing but could be considered to cover model changes section. To take this further, could apply lasso regression on the full dataset to see what that comes up with as tends to have a lower MSE than a standard LM.

F-statistic: 2.677 on 5 and 32 DF, p-value: 0.03941

```
#Using the logged_data with no adjustments
logged_data <- data_2012 %>%
  inner_join(clusterings) %>%
  mutate(adjusted_bees = log(wildAbF + 1)) %>%
  mutate(adjusted_social = log(socialRichF + 1))

## Joining, by = "orchard"

#Defining linear model with temp and X2000nat as factors
temp_nat_lm <- tidy(lm(adjusted_bees ~ temp + X2000nat, data = logged_data))
#Extracting temp and nat estimate values
temp_factor <- temp_nat_lm %>%
  select(estimate) %>%
  slice(2) %>%
  as_vector()
nat_factor <- temp_nat_lm %>%
```

```
select(estimate) %>%
  slice(3) %>%
  as_vector()
#Now adjustments shall be made for these two variables:
#Temperature will be set to 20 degrees
#X2000 nat will be set to the meean x2000 nat value
temp nat adjusted <- logged data %>%
  mutate(adjusted_bees = (adjusted_bees + ((20 - temp) * temp_factor) + ((mean_x2000 - X2000nat) * X200
#Checking that after the adjustment they are not significant at all in the model
tidy(lm(adjusted_bees ~ temp + X2000nat, data = temp_nat_adjusted))
## # A tibble: 3 x 5
##
     term
                  estimate std.error statistic p.value
                     <dbl>
##
     <chr>>
                               <dbl>
                                          <dbl>
                                                  <dbl>
## 1 (Intercept)
                 1.24e+ 0
                              0.549
                                       2.25e+ 0 0.0309
                              0.0249 -3.79e-16 1.000
## 2 temp
                 -9.42e-18
## 3 X2000nat
                  9.12e- 2
                              0.735
                                     1.24e- 1 0.902
#As expected now the bee abundance has been adjusted for these two parameters
The same process can be applied to the variable SocialRichF, the other bee variable which needs to be
considered. Since we know that only temp and nat were statistic on the wildAbF we can go straight to the
LM and test that (May need to check this with Julia, if not just cp previous analysis...)
#Looking at the model ~ only X2000nat significant
lm(adjusted_social ~ temp + X2000nat + local.diversity + region, data = logged_data) %>%
  summary
##
## Call:
## lm(formula = adjusted_social ~ temp + X2000nat + local.diversity +
##
       region, data = logged_data)
##
## Residuals:
##
        Min
                  1Q
                       Median
                                     30
                                             Max
## -0.51626 -0.27605 -0.06653 0.19262 0.72315
##
## Coefficients:
##
                    Estimate Std. Error t value Pr(>|t|)
                               0.381099 -0.753 0.45689
## (Intercept)
                   -0.287012
## temp
                    0.007198
                               0.016144
                                           0.446
                                                 0.65869
## X2000nat
                    1.507687
                               0.489142
                                          3.082 0.00421 **
## local.diversity -0.164463
                               0.178196 -0.923
                                                  0.36295
                   -0.094068
## regionLO
                               0.223680
                                         -0.421 0.67690
## regionS
                    0.165221
                               0.287892
                                          0.574 0.57005
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.3562 on 32 degrees of freedom
## Multiple R-squared: 0.3068, Adjusted R-squared: 0.1985
## F-statistic: 2.833 on 5 and 32 DF, p-value: 0.03153
social_lm<- tidy(lm(adjusted_social ~ X2000nat, data = logged_data))</pre>
```

```
social_nat_factor <- social_lm %>%
   select(estimate) %>%
   slice(2) %>%
   as_vector()

#Adjustment for social added
temp_nat_adjusted <- temp_nat_adjusted %>%
   mutate(adjusted_social = (adjusted_social + ((mean_x2000 - X2000nat) * X2000_factor)))
```

Clustering Summarising

Functions involved with summarising the clusters:

```
#Gets the name of a variable as a string ~ now implemented in clusterise but keeping
get_name <- function(x) {</pre>
  deparse(substitute(x))
#Gets the bee variables that I am summarising for each cluster - makes it easy to change in future
clusterise <- function(bee_data, cluster_data, v_name = deparse(substitute(cluster_data))){</pre>
  inner_join(bee_data, cluster_data) %>%
    summarise(mean_honey_ab = mean(mean_honey_ab),
              mean_wild_ab = mean(mean_wild_ab),
              mean social rich = mean(mean social rich),
              n = n()) \%
   mutate(cluster = v name)
}
#The function to output the summaries of both agglom and kmeans clustering
cluster_summarise <- function(bee_data){</pre>
output <- list()</pre>
output$agglom_bees <- clusterise(bee_data, aggl_node1) %>%
  bind_rows(clusterise(bee_data, aggl_node2)) %>%
  bind_rows(clusterise(bee_data, aggl_node3)) %>%
  bind_rows(clusterise(bee_data, aggl_node4)) %>%
  bind_rows(clusterise(bee_data, aggl_node5)) %>%
  bind_rows(clusterise(bee_data, aggl_node6)) %>%
  bind_rows(clusterise(bee_data, aggl_node7)) %>%
  bind_rows(clusterise(bee_data, aggl_node4 %>% filter(cluster == 1), "agglom_node8")) %>%
  bind_rows(clusterise(bee_data, aggl_node4 %>% filter(cluster == 2), "agglom_node9")) %>%
  bind_rows(clusterise(bee_data, aggl_node5 %>% filter(cluster == 1), "agglom_node10")) %>%
  bind_rows(clusterise(bee_data, aggl_node5 %>% filter(cluster == 2), "agglom_node11")) %>%
  bind_rows(clusterise(bee_data, aggl_node6 %>% filter(cluster == 1), "agglom_node12")) %>%
  bind_rows(clusterise(bee_data, aggl_node6 %>% filter(cluster == 2), "agglom_node13")) %>%
  #Only 1 result for cluster as none in the latter cluster
  #Unable to to bind as no common variables for an empty node15
  bind_rows(clusterise(bee_data, aggl_node7 %% filter(cluster == 1), "agglom_node14"))
output$kmeans_bees <- clusterise(bee_data, kmeans_node1) %>%
  bind_rows(clusterise(bee_data, kmeans_node2)) %>%
  bind_rows(clusterise(bee_data, kmeans_node3)) %>%
```

```
bind_rows(clusterise(bee_data, kmeans_node4)) %>%
  bind_rows(clusterise(bee_data, kmeans_node5)) %>%
  bind_rows(clusterise(bee_data, kmeans_node6)) %>%
  bind_rows(clusterise(bee_data, kmeans_node7)) %>%
  bind_rows(clusterise(bee_data, kmeans_node4 %>% filter(cluster == 1), "kmeans_node8")) %>%
  bind_rows(clusterise(bee_data, kmeans_node4 %>% filter(cluster == 2), "kmeans_node9")) %>%
  bind_rows(clusterise(bee_data, kmeans_node5 %>% filter(cluster == 1), "kmeans_node10")) %>%
  bind rows(clusterise(bee data, kmeans node5 %% filter(cluster == 2), "kmeans node11")) %>%
  bind rows(clusterise(bee data, kmeans node6 %% filter(cluster == 1), "kmeans node12")) %>%
  bind_rows(clusterise(bee_data, kmeans_node6 %>% filter(cluster == 2), "kmeans_node13")) %>%
  bind_rows(clusterise(bee_data, kmeans_node7 %>% filter(cluster == 1), "kmeans_node14")) %>%
  bind_rows(clusterise(bee_data, kmeans_node7 %>% filter(cluster == 2), "kmeans_node15"))
#Return a list containing two tibbles one for each type of clustering algorithm
output
}
Original summarising without any variable correction
#original Bee data (logged has no changes to it)
bee_values <- logged_data %>%
  group_by(orchard) %>%
  summarise(mean_honey_ab = mean(apisAb),
            mean_wild_ab = mean(adjusted_bees),
            mean_social_rich = mean(adjusted_social)
original_bee_summary <- cluster_summarise(bee_data = bee_values)</pre>
original_bee_summary$agglom_bees
## # A tibble: 14 x 5
      mean_honey_ab mean_wild_ab mean_social_rich
##
                                                      n cluster
##
              <dbl>
                           <dbl>
                                            <dbl> <int> <chr>
## 1
               6.56
                           1.33
                                            0.379
                                                     19 aggl_node1
## 2
               6.19
                           1.23
                                            0.386
                                                     15 aggl_node2
## 3
               7.94
                           1.71
                                            0.353
                                                      4 aggl_node3
## 4
               6.80
                           1.23
                                            0.376
                                                     12 aggl_node4
## 5
               3.75
                           1.23
                                            0.423
                                                      3 aggl_node5
                                            0.135
## 6
               8.75
                           1.70
                                                      3 aggl_node6
## 7
               5.5
                           1.73
                                            1.01
                                                      1 aggl_node7
               5.46
                           0.963
## 8
                                            0.325
                                                      9 agglom_node8
## 9
              10.8
                           2.04
                                            0.530
                                                      3 agglom_node9
## 10
               3.25
                           1.32
                                            0.405
                                                      1 agglom_node10
## 11
               4
                           1.19
                                            0.432
                                                      2 agglom_node11
## 12
               8
                           1.97
                                            0.101
                                                      2 agglom_node12
## 13
              10.2
                                            0.203
                                                       1 agglom node13
                           1.18
## 14
               5.5
                           1.73
                                            1.01
                                                       1 agglom_node14
#Kmeans
original_bee_summary$kmeans_bees
## # A tibble: 15 x 5
```

n cluster

mean_honey_ab mean_wild_ab mean_social_rich

##		<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<int></int>	<chr></chr>
##	1	6.56	1.33	0.379	19	kmeans_node1
##	2	6.19	1.23	0.386	15	kmeans_node2
##	3	7.94	1.71	0.353	4	kmeans_node3
##	4	6.5	1.36	0.516	5	kmeans_node4
##	5	6.04	1.17	0.320	10	kmeans_node5
##	6	7.94	1.71	0.353	4	kmeans_node6
##	7	NaN	NaN	NaN	0	kmeans_node7
##	8	3.25	1.32	0.405	1	kmeans_node8
##	9	7.31	1.37	0.544	4	kmeans_node9
##	10	6.04	1.17	0.320	10	${\tt kmeans_node10}$
##	11	NaN	NaN	NaN	0	kmeans_node11
##	12	7.17	1.89	0.403	3	${\tt kmeans_node12}$
##	13	10.2	1.18	0.203	1	kmeans_node13
##	14	NaN	NaN	NaN	0	${\tt kmeans_node14}$
##	15	NaN	NaN	NaN	0	kmeans_node15

$\hbox{\it\#\#Original summarising of non-markov approach}\\ \hbox{\it maximum_cluster}$

##		eigB11F.pre	eiqB11I.pre	eigB11F.blm	eigB11I.blm	eigB11T.blm	eigB11F.pos
##	1	143.09	0.000	58.10	19.04	0.0440	14.48
##	2	161.57	0.000	53.16	9.36	0.0000	31.37
##	3	161.57	0.000	53.16	9.36	0.0000	31.37
##	4	161.57	0.000	53.16	9.36	0.0000	31.37
##	5	41.85	0.000	44.67	14.25	0.0260	32.29
##	6	9.38	0.000	37.44	36.36	0.4600	24.39
##	7	163.95	2.310	112.90	12.62	0.0250	88.69
##	8	111.67	0.000	127.30	8.87	0.2200	53.11
##	9	94.76	0.960	52.27	19.69	0.0140	26.59
##		107.72	0.000	0.56	1.80	0.0000	41.85
##		196.87	0.330	63.57	4.94	0.0060	20.25
##		193.35	0.056	63.48	4.94	0.0031	19.87
##		91.93	0.000	53.30	8.13	0.0034	18.75
##		18.03	0.000	70.09	13.10	0.0000	20.53
	15	21.18	6.740	17.79	5.49	0.0000	7.57
##		102.61	0.000	68.15	7.54	3.0000	29.44
##		106.47	0.000	33.00	28.70	0.0160	15.17
##		109.58	0.000	54.16	11.38	0.1700	16.79
##	19	113.28	0.000	102.91	3.61	0.0000	68.14
##			eiqB11T.pos				
##		24.31	0.000	1			
##		57.66	0.000	1			
##		57.66	0.000	1			
##		57.66	0.000	1			
##		9.63	0.000	2			
##		34.75	0.000	3			
##		51.87	0.041	4			
##		5.25	0.000	5			
##		20.81	0.082	6			
##		1.80	0.000	7			
##		21.36	0.084	8			
##		21.36	0.230	8			
##		7.58	0.061	6			
##	14	2.28	0.000	2			

```
## 15
             9.48
                        0.000
## 16
            10.91
                        3.000
                                    6
            10.08
                        0.000
## 17
                                    6
            17.33
                        0.220
                                    6
## 18
## 19
            11.92
                        0.070
                                    5
#Clustering summaries based on non-markov approach
non_markov_clusters <- logged_data %>%
  group_by(orchard) %>%
  summarise(mean_honey_ab = mean(apisAb),
            mean_wild_ab = mean(adjusted_bees),
            mean_social_rich = mean(adjusted_social)) %>%
  mutate(cluster = maximum_cluster$cluster) %>%
  group_by(cluster) %>%
  summarise(mean_honey_ab = mean(mean_honey_ab),
            mean_wild_ab = mean(mean_wild_ab),
            mean_social_rich = mean(mean_social_rich),
            n=n()
After Adjustment:
adjusted_bee_values <- temp_nat_adjusted %>%
  group_by(orchard) %>%
  summarise(mean_honey_ab = mean(apisAb),
            mean_wild_ab = mean(adjusted_bees),
            mean_social_rich = mean(adjusted_social)
adjusted_bee_summary <- cluster_summarise(bee_data = adjusted_bee_values)</pre>
#Agglom
adjusted_bee_summary$agglom_bees
## # A tibble: 14 x 5
##
      mean_honey_ab mean_wild_ab mean_social_rich
                                                       n cluster
##
              <dbl>
                           <dbl>
                                            <dbl> <int> <chr>
## 1
               6.56
                           1.27
                                            0.379
                                                      19 aggl_node1
## 2
               6.19
                           1.15
                                            0.395
                                                      15 aggl_node2
                                            0.319
## 3
               7.94
                           1.72
                                                      4 aggl_node3
## 4
               6.80
                           1.19
                                            0.426
                                                      12 aggl_node4
## 5
               3.75
                           1.01
                                            0.270
                                                      3 aggl_node5
## 6
               8.75
                           1.90
                                            0.276
                                                      3 aggl_node6
## 7
               5.5
                                            0.449
                                                      1 aggl_node7
                           1.17
## 8
               5.46
                           1.06
                                            0.474
                                                      9 agglom_node8
## 9
              10.8
                                            0.281
                           1.56
                                                      3 agglom_node9
               3.25
                                            0.272
## 10
                           1.06
                                                      1 agglom_node10
## 11
               4
                           0.980
                                            0.269
                                                      2 agglom_node11
## 12
               8
                           2.05
                                            0.266
                                                       2 agglom_node12
              10.2
## 13
                           1.59
                                            0.295
                                                       1 agglom_node13
## 14
               5.5
                           1.17
                                            0.449
                                                       1 agglom_node14
#Kmeans
adjusted_bee_summary$kmeans_bees
## # A tibble: 15 x 5
##
      mean_honey_ab mean_wild_ab mean_social_rich
                                                      n cluster
```

```
##
               <dbl>
                             <dbl>
                                                <dbl> <int> <chr>
##
                6.56
                              1.27
   1
                                                0.379
                                                         19 kmeans_node1
##
   2
                6.19
                              1.15
                                                0.395
                                                         15 kmeans node2
                7.94
##
   3
                              1.72
                                                0.319
                                                          4 kmeans_node3
##
    4
                6.5
                              1.13
                                                0.356
                                                          5 kmeans node4
   5
                6.04
                                                0.414
                                                       10 kmeans node5
##
                              1.16
                                                0.319
                                                          4 kmeans node6
##
   6
                7.94
                              1.72
   7
##
              NaN
                            NaN
                                             NaN
                                                          0 kmeans node7
##
    8
                3.25
                              1.06
                                                0.272
                                                          1 kmeans node8
##
   9
                7.31
                              1.15
                                                0.377
                                                           4 kmeans_node9
## 10
                6.04
                              1.16
                                                0.414
                                                         10 kmeans_node10
                                                           0 kmeans_node11
## 11
              \mathtt{NaN}
                            {\tt NaN}
                                              \mathtt{NaN}
                7.17
## 12
                              1.76
                                                0.327
                                                           3 kmeans_node12
## 13
               10.2
                              1.59
                                                0.295
                                                           1 kmeans_node13
## 14
              NaN
                                                           0 kmeans_node14
                            NaN
                                              NaN
## 15
              NaN
                            NaN
                                              NaN
                                                           0 kmeans_node15
```

```
##Non-Markov
##Maxi.mum
#Clustering summaries based on non-markov approach
maximum_non_markov_clusters <- temp_nat_adjusted %>%
  group by (orchard) %>%
  summarise(mean_honey_ab = mean(apisAb),
            mean_wild_ab = mean(adjusted_bees),
            mean_social_rich = mean(adjusted_social)) %>%
  mutate(cluster = maximum_cluster$cluster) %>%
  group by(cluster) %>%
  summarise(mean_honey_ab = mean(mean_honey_ab),
            mean_wild_ab = mean(mean_wild_ab),
            mean_social_rich = mean(mean_social_rich),
            n=n())
##euclidean:
euclidean_non_markov_clusters <- temp_nat_adjusted %>%
  group_by(orchard) %>%
  summarise(mean honey ab = mean(apisAb),
            mean_wild_ab = mean(adjusted_bees),
            mean_social_rich = mean(adjusted_social)) %>%
  mutate(cluster = euclidean cluster$cluster) %>%
  group by(cluster) %>%
  summarise(mean_honey_ab = mean(mean_honey_ab),
            mean_wild_ab = mean(mean_wild_ab),
            mean_social_rich = mean(mean_social_rich),
            n=n()
```

Although honey bee abundance is one of the summarised variables, this is likely to be related directly to the hive acr variable.

Comparing cluster values before and after adjustments, for now just looking at the agglomerative node clustering selection as it is, in my opinion, better than Kmeans.

```
##agglom
adjusted_bee_summary$agglom_bees %>%
select(mean_wild_ab, mean_social_rich ,cluster) %>%
rename(adjusted = mean_wild_ab) %>%
rename(adjusted_social = mean_social_rich) %>%
```

```
inner_join(original_bee_summary$agglom_bees %>%
                select(mean_wild_ab, mean_social_rich, cluster)) %>%
  select(cluster,mean_wild_ab, adjusted, mean_social_rich, adjusted_social)
## Joining, by = "cluster"
## # A tibble: 14 x 5
##
                     mean_wild_ab adjusted mean_social_rich adjusted_social
      cluster
##
      <chr>
                            <dbl>
                                      <dbl>
                                                        <dbl>
                                                                         <dbl>
                                      1.27
                                                                         0.379
##
    1 aggl node1
                            1.33
                                                        0.379
##
    2 aggl_node2
                            1.23
                                      1.15
                                                        0.386
                                                                         0.395
##
    3 aggl_node3
                            1.71
                                      1.72
                                                        0.353
                                                                         0.319
##
   4 aggl_node4
                            1.23
                                      1.19
                                                        0.376
                                                                         0.426
##
    5 aggl_node5
                            1.23
                                      1.01
                                                        0.423
                                                                         0.270
   6 aggl_node6
##
                            1.70
                                      1.90
                                                        0.135
                                                                         0.276
##
   7 aggl_node7
                            1.73
                                      1.17
                                                        1.01
                                                                         0.449
##
                            0.963
                                      1.06
                                                        0.325
                                                                         0.474
   8 agglom_node8
##
  9 agglom_node9
                            2.04
                                      1.56
                                                        0.530
                                                                         0.281
## 10 agglom_node10
                            1.32
                                      1.06
                                                        0.405
                                                                         0.272
## 11 agglom_node11
                            1.19
                                      0.980
                                                        0.432
                                                                         0.269
                            1.97
                                      2.05
                                                                         0.266
## 12 agglom_node12
                                                        0.101
## 13 agglom_node13
                                                        0.203
                                                                         0.295
                            1.18
                                      1.59
## 14 agglom_node14
                            1.73
                                      1.17
                                                        1.01
                                                                         0.449
##Kmeans
adjusted_bee_summary$kmeans_bees %>%
  select(mean_wild_ab, mean_social_rich ,cluster) %>%
  rename(adjusted = mean_wild_ab) %>%
  rename(adjusted_social = mean_social_rich) %>%
  inner join(original bee summary$kmeans bees %>%
               select(mean_wild_ab, mean_social_rich, cluster)) %>%
  select(cluster,mean_wild_ab, adjusted, mean_social_rich, adjusted_social)
## Joining, by = "cluster"
## # A tibble: 15 x 5
##
      cluster
                     mean_wild_ab adjusted mean_social_rich adjusted_social
##
      <chr>
                            <dbl>
                                      <dbl>
                                                        <dbl>
                                                                         <dbl>
                             1.33
                                       1.27
                                                        0.379
                                                                         0.379
##
   1 kmeans_node1
##
    2 kmeans_node2
                             1.23
                                       1.15
                                                        0.386
                                                                         0.395
    3 kmeans_node3
                             1.71
                                                        0.353
                                                                         0.319
##
                                       1.72
##
   4 kmeans_node4
                             1.36
                                       1.13
                                                        0.516
                                                                         0.356
##
   5 kmeans node5
                             1.17
                                       1.16
                                                        0.320
                                                                         0.414
   6 kmeans_node6
                                                                         0.319
##
                             1.71
                                       1.72
                                                        0.353
##
   7 kmeans node7
                           NaN
                                     NaN
                                                      NaN
                                                                      NaN
## 8 kmeans_node8
                             1.32
                                       1.06
                                                        0.405
                                                                         0.272
## 9 kmeans_node9
                             1.37
                                       1.15
                                                        0.544
                                                                         0.377
## 10 kmeans_node10
                             1.17
                                       1.16
                                                        0.320
                                                                         0.414
## 11 kmeans_node11
                           NaN
                                     NaN
                                                      NaN
                                                                      NaN
## 12 kmeans_node12
                             1.89
                                       1.76
                                                        0.403
                                                                         0.327
                                       1.59
## 13 kmeans_node13
                             1.18
                                                        0.203
                                                                         0.295
## 14 kmeans_node14
                           NaN
                                     NaN
                                                      NaN
                                                                       NaN
## 15 kmeans_node15
                           NaN
                                     NaN
                                                      NaN
                                                                      NaN
```

I guess potentially can see two paths one with a mean slightly lower than the other if you think of the structure.

Looking at clusters

The aim of this section is to look at the clusterings from a low, medium, high point of view...

```
#Non_markov clustering summary maximum
maximum_final <- maximum_cluster %>%
 group_by(cluster) %>%
 summarise_all(funs(mean)) %>%
 inner_join(maximum_non_markov_clusters)
## Joining, by = "cluster"
#Non_markov clustering summary euclidean
euclidean final <- euclidean cluster %>%
 group_by(cluster) %>%
 summarise all(funs(mean)) %>%
 inner_join(euclidean_non_markov_clusters)
## Joining, by = "cluster"
#Cluster 1 -> Low insecticide, High Fungicide, low thinner
#Cluster 2 -> High insecticide, Low Fungicide, low (no) thinner
#Cluster 3 -> middle insecticide, High Fungicide, low thinner
#Cluster 4 -> Low insecticide, High Fungicide, High thinner
#Agglom clustering summary (end stage)
bee_summary_agglom <- adjusted_bee_summary$agglom_bees %>%
 slice(8:14) %>%
 select(-cluster) %>%
 mutate(cluster = c(1,2,3,4,5,6,7))
bind_rows(aggl_c(aggl_node2, ".pos", 2) %>%
 mutate(cluster = if_else(cluster == 1, 3, 4))) %>%
 bind_rows(aggl_c(aggl_node3, ".pos", 1)%>%
 mutate(cluster = if_else(cluster == 1, 5, 6))) %>%
 bind_rows(aggl_c(aggl_node3, ".pos", 2)%>%
 mutate(cluster = if_else(cluster ==1, 7, 8))) %>%
 group_by(cluster) %>%
 summarise_all(funs(mean))
##final agglom
agglom_summary <- bee_summary_agglom %>%
 inner_join(pesticide_summary_agglom) %>%
 select(-orchard) #%>%
## Joining, by = "cluster"
 #1 for high, O for low, just using bloom
 # mutate(pest_rating = c(0,0,1,1,0,0,1)) %>%
 # mutate(insect_rating = c(1,0,1,0,0,1,1)) %>%
 # mutate(thinner\_rating = c(1,0,0,1,0,1,0))
#Kmeans
```

```
bee_summary_kmeans <- adjusted_bee_summary$kmeans_bees %>%
  slice(8:14) %>%
  select(-cluster) %>%
  mutate(cluster = c(1,2,3,4,5,6,7))
pesticide_summary_kmeans <- aggl_c(kmeans_node2, ".pos", 1) %>%
  bind_rows(aggl_c(kmeans_node2, ".pos", 2) %>%
  mutate(cluster = if_else(cluster == 1, 3, 4))) %>%
  bind_rows(aggl_c(kmeans_node3, ".pos", 1)%>%
  mutate(cluster = if_else(cluster == 1, 5, 6))) %>%
  bind_rows(aggl_c(kmeans_node3, ".pos", 2)%>%
  mutate(cluster = if_else(cluster ==1, 7, 8))) %>%
  group_by(cluster) %>%
  summarise_all(funs(mean))
##
kmeans_summary <- bee_summary_kmeans %>%
  inner_join(pesticide_summary_kmeans) %>%
  select(-orchard)
## Joining, by = "cluster"
#write.csv(agglom_summary, "agglom_summary.csv")
#Overall
#HLH
#HHL
#HHL
#HLH
#LLL
#T.HH
#LLL
#Just Bloom
#LHH
#LLL
#HHL
#HLH
\#LLL
#LHH
#HHL
##Just bloom and just fungicide and insecticide
```

##Final summaries between models understand why there are more clusters in the non-standarised version? Put back into unit people are more familiar sense. Transform back, probs not due to?

```
round_df <- function(df, digits = 2) {
   #Using a temp variable to not round cluster and n as these are integers
temp <- df %>%
   mutate(cluster = as.character(cluster)) %>%
   mutate(n = as.character(n))
nums <- vapply(temp, is.numeric , FUN.VALUE = logical(1))</pre>
```

```
df[,nums] <- round(df[,nums], digits = digits)</pre>
  df
}
#Need a different rounding function when cluster/n not present
round_df_2 <- function(df, digits = 2) {</pre>
  nums <- vapply(df, is.numeric , FUN.VALUE = logical(1))</pre>
  df[,nums] <- round(df[,nums], digits = digits)</pre>
  df
}
apple_order <- function(data){</pre>
  data %>%
    select(cluster, n, mean_wild_ab, mean_social_rich, contains("pre"), contains("blm"), contains("pos"
}
#Rounding data for shiny app display
maximum_final_standardised <- round_df(maximum_final_standardised) %>%
  apple order()
euclidean_final_standardised <- round_df(euclidean_final_standardised) %>%
  apple_order()
#no n
agglom_summary_standardised <- round_df(agglom_summary_standardised) %>%
  apple_order()
#no n
kmeans_summary_standardised <- round_df(kmeans_summary_standardised) %>%
  apple_order()
# 8 clusters in this method - 6 different options so this is the cluster that will used - missing optio
maximum_final <- round_df(maximum_final) %>%
  apple_order()
maximum final app <- maximum final %>%
  mutate(fung_level = c("high","low","low","high","high","low","low","high")) %>%
  mutate(insect_level = c("low", "high", "high", "high", "low", "high", "low", "low", "low")) %>%
  mutate(thinner_level = c("low","low","high","low","high","low","high"))
protocol_summary <- maximum_final_app %>%
  group_by(fung_level, insect_level, thinner_level) %>%
  summarise(mean_wild_ab = mean(mean_wild_ab),
            mean_social_rich = mean(mean_social_rich)) %>%
  ungroup() %>%
  mutate(unlogged_ab = exp(mean_wild_ab) + 1) %>%
  mutate(unlogged_rich = exp(mean_social_rich) + 1) %>%
  round df 2()
```

```
euclidean_final <- round_df(euclidean_final) %>%
  apple_order()
#7 Clusters in this method - 5 diff options
agglom_summary <- round_df(agglom_summary) %>%
  mutate(fung_level = c(1,1,1,1,0,0,0)) %>%
  mutate(insect_level = c(1,1,1,0,0,1,0)) %>%
 mutate(thinner_level = c(1,0,1,1,0,1,0)) %>%
  apple_order()
#Removes previous NA inserted due to joining.
kmeans_summary <- round_df(kmeans_summary) %>%
  na.omit() %>%
  apple_order()
#Displaying table outputs
maximum_final_standardised
## # A tibble: 2 x 12
##
                 n mean_wild_ab mean_social_rich eiqB11F.pre eiqB11I.pre
     cluster
##
       <int> <int>
                          <dbl>
                                            <dbl>
                                                        <dbl>
## 1
                           1.28
                                             0.4
                                                         0.1
                                                                    -0.21
           1
                17
## 2
           2
                 2
                           1.15
                                             0.22
                                                        -0.86
                                                                     1.76
## # ... with 6 more variables: eigB11F.blm <dbl>, eigB11I.blm <dbl>,
       eiqB11T.blm <dbl>, eiqB11F.pos <dbl>, eiqB11I.pos <dbl>,
## #
       eiqB11T.pos <dbl>
euclidean final standardised
## # A tibble: 5 x 12
##
                 n mean_wild_ab mean_social_rich eiqB11F.pre eiqB11I.pre
     cluster
##
       <int> <int>
                          <dbl>
                                            <dbl>
                                                        <dbl>
                                                                    <dbl>
## 1
           1
                 9
                           1.44
                                            0.49
                                                        -0.54
                                                                   -0.28
           2
## 2
                 5
                           1.17
                                             0.3
                                                         1.12
                                                                   -0.290
## 3
           3
                 3
                           1.01
                                             0.27
                                                         0.33
                                                                    0.14
## 4
           4
                 1
                           1.57
                                             0.3
                                                        -1.58
                                                                    3.87
           5
                           0.74
                                                        -0.15
## 5
                 1
                                             0.14
                                                                   -0.34
## # ... with 6 more variables: eigB11F.blm <dbl>, eigB11I.blm <dbl>,
       eiqB11T.blm <dbl>, eiqB11F.pos <dbl>, eiqB11I.pos <dbl>,
       eigB11T.pos <dbl>
agglom_summary_standardised
## # A tibble: 4 x 12
##
     cluster
                 n mean_wild_ab mean_social_rich eiqB11F.pre eiqB11I.pre
##
       <dbl> <int>
                          <dbl>
                                                        <dbl>
                                            <dbl>
                                                                    <dbl>
## 1
           1
                13
                           1.24
                                             0.43
                                                        -0.14
                                                                    -0.28
           2
                                             0.28
## 2
                 4
                           1.44
                                                         0.9
                                                                     0.02
## 3
           3
                 1
                           0.74
                                             0.14
                                                        -0.15
                                                                    -0.34
           5
                           1.57
                                             0.3
                                                                     3.87
                 1
                                                        -1.58
## # ... with 6 more variables: eiqB11F.blm dbl>, eiqB11I.blm dbl>,
       eiqB11T.blm <dbl>, eiqB11F.pos <dbl>, eiqB11I.pos <dbl>,
       eiqB11T.pos <dbl>
kmeans summary standardised
```

A tibble: 3 x 12

```
n mean_wild_ab mean_social_rich eigB11F.pre eigB11I.pre
##
       <dbl> <int>
                           <dbl>
                                            <dbl>
                                                         <dbl>
                                                                     <dbl>
                                                         -0.25
                            1.26
                                             0.42
                                                                      0.02
## 1
           1
                14
## 2
                 4
                            1.44
                                             0.28
                                                          0.9
                                                                      0.02
           2
## 3
           5
                 1
                            0.74
                                             0.14
                                                         -0.15
                                                                     -0.34
## # ... with 6 more variables: eigB11F.blm <dbl>, eigB11I.blm <dbl>,
       eigB11T.blm <dbl>, eigB11F.pos <dbl>, eigB11I.pos <dbl>,
       eigB11T.pos <dbl>
maximum final
## # A tibble: 8 x 12
##
                 n mean_wild_ab mean_social_rich eigB11F.pre eigB11I.pre
     cluster
##
       <int> <int>
                          <dbl>
                                            <dbl>
                                                         <dbl>
                                                                     <dbl>
## 1
           1
                 4
                           1.48
                                             0.36
                                                         157.
                                                                      0
## 2
           2
                 2
                                             0.34
                                                          29.9
                                                                      0
                           1.85
## 3
           3
                 2
                          1.58
                                             0.3
                                                          15.3
                                                                      3.37
## 4
           4
                 1
                          1.06
                                             0.27
                                                         164.
                                                                      2.31
## 5
           5
                 2
                                             0.27
                           0.98
                                                         112.
                                                                      Ω
## 6
           6
                 5
                           1.12
                                             0.5
                                                         101.
                                                                      0.19
## 7
           7
                           1.58
                                             0.48
                                                         108.
                                                                      Ω
                 1
           8
                 2
                           0.580
                                             0.33
                                                         195.
                                                                      0.19
## # ... with 6 more variables: eigB11F.blm <dbl>, eigB11I.blm <dbl>,
       eiqB11T.blm <dbl>, eiqB11F.pos <dbl>, eiqB11I.pos <dbl>,
       eiqB11T.pos <dbl>
euclidean final
## # A tibble: 4 x 12
                 n mean_wild_ab mean_social_rich eiqB11F.pre eiqB11I.pre
     cluster
##
       <int> <int>
                           <dbl>
                                            <dbl>
                                                         <dbl>
                                                                     <dbl>
## 1
                 6
                            1.18
                                             0.35
                                                         170.
                                                                      0.06
           1
## 2
           2
                 4
                            1.72
                                             0.32
                                                          22.6
                                                                      1.68
## 3
                            1.01
                                             0.27
           3
                 3
                                                         130.
                                                                      0.77
           4
                            1.2
                                             0.5
                                                         102.
                                                                      0.16
## # ... with 6 more variables: eiqB11F.blm <dbl>, eiqB11I.blm <dbl>,
       eigB11T.blm <dbl>, eigB11F.pos <dbl>, eigB11I.pos <dbl>,
       eiqB11T.pos <dbl>
agglom_summary
## # A tibble: 7 x 12
##
                 n mean_wild_ab mean_social_rich eigB11F.pre eigB11I.pre
     cluster
       <dbl> <int>
                           <dbl>
                                            <dbl>
                                                         <dbl>
                                                                     <dbl>
                                            0.47
                                                        127.
## 1
           1
                 9
                            1.06
                                                                      0.15
## 2
           2
                 3
                            1.56
                                            0.28
                                                        162.
                                                                      Ω
## 3
           3
                 1
                            1.06
                                            0.27
                                                        164.
                                                                      2.31
                            0.98
## 4
           4
                 2
                                            0.27
                                                        112.
## 5
           5
                 2
                            2.05
                                            0.27
                                                         31.5
                                                                      3.37
                                                          9.38
## 6
           6
                 1
                            1.59
                                            0.290
                                                                      0
## 7
           7
                 1
                            1.17
                                            0.45
                                                         18.0
## # ... with 6 more variables: eiqB11F.blm <dbl>, eiqB11I.blm <dbl>,
```

kmeans_summary

eiqB11T.pos <dbl>

eiqB11T.blm <dbl>, eiqB11F.pos <dbl>, eiqB11I.pos <dbl>,

```
##
     cluster
                 n mean_wild_ab mean_social_rich eiqB11F.pre eiqB11I.pre
       <dbl> <int>
                                                                     <dbl>
##
                           <dbl>
                                            <dbl>
                                                         <dbl>
           1
                            1.06
                                            0.27
                                                        164.
                                                                      2.31
## 1
                 1
           2
                                            0.38
## 2
                 4
                            1.15
                                                        110.
                                                                      0.19
## 3
           3
                10
                            1.16
                                            0.41
                                                        133.
## 4
           5
                 3
                            1.76
                                            0.33
                                                         27.0
                                                                      2.25
                                                          9.38
## 5
           6
                            1.59
                                            0.290
                 1
## # ... with 6 more variables: eiqB11F.blm <dbl>, eiqB11I.blm <dbl>,
       eiqB11T.blm <dbl>, eiqB11F.pos <dbl>, eiqB11I.pos <dbl>,
## #
       eiqB11T.pos <dbl>
#Could do something with this and plots coloured by cluster or something like that perhaps?
temp_nat_adjusted %>%
  group_by(orchard) %>%
  summarise(mean_honey_ab = mean(apisAb),
            mean_wild_ab = mean(adjusted_bees),
            mean_social_rich = mean(adjusted_social))
## # A tibble: 19 x 4
##
      orchard mean_honey_ab mean_wild_ab mean_social_rich
##
      <fct>
                      <dbl>
                                    <dbl>
                                                     <dbl>
## 1 A
                       5.12
                                    1.22
                                                    0.595
## 2 B
                       8
                                    1.70
                                                   -0.207
## 3 C
                       5.75
                                    1.67
                                                    0.556
## 4 D
                                    1.32
                                                    0.493
                      18.8
## 5 E
                       7.25
                                    2.53
                                                     0.230
## 6 F
                      10.2
                                    1.59
                                                    0.295
## 7 G
                       3.25
                                    1.06
                                                    0.272
## 8 H.nooil
                       3.75
                                    1.09
                                                    0.574
## 9 I
                       1.75
                                    1.04
                                                     0.808
## 10 J
                       6.25
                                    1.58
                                                     0.485
## 11 K
                       6.25
                                    0.347
                                                     0.347
## 12 L
                                    0.806
                       6
                                                     0.321
## 13 M
                       2.75
                                    1.55
                                                     0.562
## 14 N
                       5.5
                                    1.17
                                                     0.449
                       8.75
## 15 O.nooil
                                    1.57
                                                     0.303
## 16 P
                       2
                                    0.735
                                                     0.140
## 17 Q
                      15
                                                    0.487
                                    1.07
## 18 R
                       4
                                    1.22
                                                    0.525
## 19 S
                       4.25
                                    0.869
                                                   -0.0364
save(data_2012, maximum_final, maximum_final_app, euclidean_final, agglom_summary, kmeans_summary,
               maximum_final_standardised, euclidean_final_standardised,
```

A tibble: 5 x 12

agglom_summary_standardised, kmeans_summary_standardised, clust_comps, standardised_clus