## MA615 Unit 1 final

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### Introduction

This project uses row data "berries.csv" to produce a tidy data. Also, I extract subset of tidy data to do EDA and PCA. You will see notation after "##" and "#". "##" means primary item and "#" means branch item.

#### read the data

```
berries <- read_csv("berries.csv", col_names = TRUE)
```

```
## Parsed with column specification:
## .default = col_character(),
  Year = col double(),
##
    `Week Ending` = col_logical(),
    `Ag District` = col_logical(),
##
    `Ag District Code` = col_logical(),
##
##
   County = col logical(),
    `County ANSI` = col_logical(),
##
    `Zip Code` = col_logical(),
##
## Region = col_logical(),
    Watershed = col logical(),
##
##
   `CV (%)` = col logical()
## )
```

```
## See spec(...) for full column specifications.
```

```
Data = berries
```

## Data cleaning

```
##Remove single value

Data %<>% select(-c("Program", "Geo Level", "Ag District", "Week Ending", 8:15, 21))

#Remove NA, D and process (Z)

Data <- filter(Data, Value!='(D)', Value!='(NA)')

Data$Value[which(Data$Value=='(Z)')] <- 0
```

```
##Split Data Item
new <- Data
# new <- separate(new, col = "Data Item", into = c("species", "definition", "unit"), sep = ", ", remov
e = TRUE)
# new <- separate(new, col = "Data Item", into = c("q", "w", "e", "r"), sep = ", ", remove = TRUE)
# new$unit <- tail(strsplit(new$`Data Item`, split=", ")[[1]], 1)</pre>
```

Because Data Item is mess, I split it into several columns and change every wrong item into a right value. Finally, I extract right values from each columns and delete redundant columns.

```
#1. Unit
nr <- nrow(new)
for (i in 1:nr) {
  new$unit[i] <- tail(strsplit(new$`Data Item`, split=", ")[[i]],1)
## Warning: Unknown or uninitialised column: `unit`.
new1 <- new
new2 <- new1
#Replace untidy data
new2$unit[which(new2$unit=="STRAWBERRIES - ACRES HARVESTED"|
                  new2$unit==" WILD - ACRES HARVESTED"|
                  new2$unit==" RED - ACRES HARVESTED"|
                  new2$unit==" TAME - ACRES HARVESTED"|
                  new2$unit=="RASPBERRIES - ACRES HARVESTED"|
                  new2$unit==" BLACK - ACRES HARVESTED")] <- "ACRES HARVESTED"</pre>
new2$unit[which(new2$unit=="STRAWBERRIES - ACRES PLANTED")] <- "ACRES PLANTED"</pre>
unique (new2$unit)
## [1] " MEASURED IN $ / LB"
                                   " MEASURED IN $ / CWT"
## [3] "ACRES HARVESTED"
                                   " MEASURED IN LB"
## [5] " MEASURED IN LB / ACRE" " MEASURED IN $"
## [7] " AVG"
                                  "ACRES PLANTED"
## [9] " MEASURED IN CWT"
                                   " MEASURED IN CWT / ACRE"
## [11] " MEASURED IN $ / TON"
                                   " MEASURED IN TONS"
#2. Type
new3 <- new2
new3 %<>% separate(`Data Item`, c("B", "type", "meas", "what"), sep = ",", remove = FALSE)
## Warning: Expected 4 pieces. Missing pieces filled with `NA` in 1819 rows [1, 4,
## 5, 6, 7, 8, 11, 14, 17, 20, 21, 26, 27, 28, 29, 30, 33, 34, 35, 36, ...].
new3 %<>% select(-B)
new3 %<>% separate(type,c("b1", "type", "b2", "lab1", "lab2"), " ")
## Warning: Expected 5 pieces. Additional pieces discarded in 130 rows [4, 5, 7,
## 26, 28, 33, 46, 129, 533, 1767, 1822, 1823, 1824, 1828, 1829, 1844, 1846, 1847,
## 1853, 1856, ...].
## Warning: Expected 5 pieces. Missing pieces filled with `NA` in 5594 rows [2, 3,
```

## 9, 10, 12, 13, 15, 16, 18, 19, 22, 23, 24, 25, 31, 32, 35, 36, 37, 38, ...].

```
new3[is.na(new3)] <- " " ## OK now Data Item has been split into parts

# unique(new3$type)

#I found there is something redundant about new3$type. There will be 3 types in the final data:
tame, wild and bearing. So I remove others.

new3$type[which(new3$type=="MEASURED"

# new3$type=="PROCESSING"/

# new3$type=="NOT"/

# new3$type=="NOT"/

# new3$type=="BLACK"/

# new3$type=="BLACK"/

# new3$type=="RED"

)] <- " "

new3$type[which(new3$type=="NOT")] <- "NOT SOLD"

new3$type[which(new3$type=="FRESH")] <- "FRESH MARKET"
```

```
#3. Production
#The true values are hided in columns "lab1, lab2, meas, what"
new4 <- new3
new4 %<>% select(-c(`State ANSI`, b1, b2))
#settle these 4 columns and paste into 1 column
# unique(new4$1ab1)
new4\$1ab1[which(new4\$1ab1=="\$"]
                new4$1ab1=="-"|
                new4$1ab1=="ACRES"
                new4$1ab1=="LB"|
                new4$1ab1=="CWT")] <- " "
# unique(new4$1ab2)
new4\$1ab2[which(new4\$1ab2=="/"|
                new4$1ab2=="HARVESTED")] <- " "
# unique(new4$meas)
new4$meas[which(new4$meas==" MEASURED IN $ / LB"|
                new4$meas==" MEASURED IN LB / ACRE"|
                new4$meas==" MEASURED IN LB / ACRE / YEAR"|
                new4$meas==" MEASURED IN $"|
                new4$meas==" MEASURED IN $ / CWT"|
                new4$meas==" MEASURED IN NUMBER"|
                new4$meas==" MEASURED IN CWT"|
                new4$meas=="MEASURED IN LB"|
                new4$meas==" MEASURED IN LB / ACRE / APPLICATION"|
                new4$meas==" MEASURED IN PCT OF AREA BEARING"|
                new4$meas==" MEASURED IN $ / TON")] <- " "
new4$meas[which(new4$meas=="FRESH MARKET - PRICE RECEIVED"|
                new4$meas==" PROCESSING - PRICE RECEIVED" )] <- "PRICE RECEIVED"</pre>
new4$meas[which(new4$meas==" FRESH MARKET - PRODUCTION"|
                new4$meas==" NOT SOLD - PRODUCTION"|
                new4$meas==" PROCESSING - PRODUCTION"|
                new4$meas==" UTILIZED - PRODUCTION")] <- "PRODUCTION"</pre>
new4$meas[which(new4$meas==" UTILIZED - YIELD")] <- "YIELD"</pre>
# unique(new4$what)
new4 %<>% select(-what)
```

```
#combine these 3 columns
new4 %<>% mutate(production = str_trim(paste(lab1, lab2, meas)) )
#unique(new4$production)
new4$production[which(new4$production=="PRICE")] <- "PRICE RECEIVED"</pre>
new4 \% \% select (-c (lab1, lab2, meas))
#process column production
new4$production[c(4,5,7)] <- "PRICE RECEIVED"</pre>
##onto Domain
new5 <- new4
# new5$Domain %>% unique()
new5 %<>% separate(Domain, c("D_left", "D_right"), sep = ", ")
## Warning: Expected 2 pieces. Missing pieces filled with `NA` in 1791 rows [1, 2,
## 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, ...].
# new5$D left %>% unique()
# new5$D_right %>% unique()
new5[is.na(new5)] \leftarrow ""
# And now Domain Category
# new5$`Domain Category` %>% unique()
new5 %<>% separate(`Domain Category`, c("DC_left", "DC_right"), sep = ", ")
## Warning in gregexpr(pattern, x, perl = TRUE): PCRE error
## 'UTF-8 error: isolated byte with 0x80 bit set'
## for element 1922
## Warning in gregexpr(pattern, x, perl = TRUE): PCRE error
## 'UTF-8 error: isolated byte with 0x80 bit set'
## for element 1983
## Warning in gregexpr(pattern, x, perl = TRUE): PCRE error
  'UTF-8 error: isolated byte with 0x80 bit set'
## for element 2042
## Warning in gregexpr(pattern, x, perl = TRUE): PCRE error
## 'UTF-8 error: isolated byte with 0x80 bit set'
## for element 2107
## Warning in gregexpr(pattern, x, perl = TRUE): PCRE error
## 'UTF-8 error: isolated byte with 0x80 bit set'
## for element 2174
```

```
## Warning in gregexpr(pattern, x, perl = TRUE): PCRE error
  'UTF-8 error: isolated byte with 0x80 bit set'
## for element 3995
## Warning in gregexpr(pattern, x, per1 = TRUE): PCRE error
  'UTF-8 error: isolated byte with 0x80 bit set'
  for element 4052
## Warning in gregexpr(pattern, x, perl = TRUE): PCRE error
## 'UTF-8 error: isolated byte with 0x80 bit set'
## for element 4107
## Warning in gregexpr(pattern, x, perl = TRUE): PCRE error
## 'UTF-8 error: isolated byte with 0x80 bit set'
## for element 4169
## Warning in gregexpr(pattern, x, perl = TRUE): PCRE error
  'UTF-8 error: isolated byte with 0x80 bit set'
## for element 4233
## Warning: Expected 2 pieces. Additional pieces discarded in 20 rows [184, 259,
## 332, 414, 498, 1336, 1385, 1431, 1478, 1531, 1932, 1993, 2052, 2117, 2184, 4005,
## 4062, 4117, 4179, 4243].
## Warning: Expected 2 pieces. Missing pieces filled with `NA` in 1801 rows [1, 2,
## 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, ...].
# looks like DC left combines labels
head(new5$DC_left %>% unique(), n=20)
## [1] "NOT SPECIFIED"
## [2] "CHEMICAL"
## [3] "FERTILIZER: (NITROGEN)"
## [4] "FERTILIZER: (PHOSPHATE)"
## [5] "FERTILIZER: (POTASH)"
## [6] "FERTILIZER: (SULFUR)"
## [7] "CHEMICAL, INSECTICIDE: (CYFLUMETOFEN<U+00A0>= 138831)"
head(new5$DC_right %>% unique(), n=20)
```

```
##
   [1] NA
  [2] "FUNGICIDE: (BOSCALID = 128008)"
##
## [3] "FUNGICIDE: (CYPRODINIL = 288202)"
## [4] "FUNGICIDE: (FLUDIOXONIL = 71503)"
  [5] "FUNGICIDE: (MYCLOBUTANIL = 128857)"
## [6] "FUNGICIDE: (PYRACLOSTROBIN = 99100)"
## [7] "FUNGICIDE: (TOTAL)"
  [8] "HERBICIDE: (TOTAL)"
## [9] "INSECTICIDE: (ACEQUINOCYL = 6329)"
## [10] "INSECTICIDE: (BIFENAZATE = 586)"
## [11] "INSECTICIDE: (METHOXYFENOZIDE = 121027)"
## [12] "INSECTICIDE: (PYRETHRINS = 69001)"
## [13] "INSECTICIDE: (SPINETORAM = 110007)"
## [14] "INSECTICIDE: (SPINOSAD = 110003)"
## [15] "INSECTICIDE: (TOTAL)"
## [16] "INSECTICIDE: (ZETA-CYPERMETHRIN = 129064)"
## [17] "OTHER: (TOTAL)"
## [18] "INSECTICIDE: (BT KURSTAKI ABTS-351 = 6522)"
## [19] "FUNGICIDE: (AZOXYSTROBIN = 128810)"
## [20] "FUNGICIDE: (BLAD = 30006)"
## work on DC left first
new5 %<>% separate(DC_left, c("DC_left_1", "DC_left_r"), sep = ": ")
## Warning in gregexpr(pattern, x, per1 = TRUE): PCRE error
## 'UTF-8 error: isolated byte with 0x80 bit set'
## for element 1922
## Warning in gregexpr(pattern, x, per1 = TRUE): PCRE error
## 'UTF-8 error: isolated byte with 0x80 bit set'
## for element 1983
## Warning in gregexpr(pattern, x, per1 = TRUE): PCRE error
## 'UTF-8 error: isolated byte with 0x80 bit set'
## for element 2042
## Warning in gregexpr(pattern, x, perl = TRUE): PCRE error
## 'UTF-8 error: isolated byte with 0x80 bit set'
## for element 2107
## Warning in gregexpr(pattern, x, per1 = TRUE): PCRE error
  'UTF-8 error: isolated byte with 0x80 bit set'
## for element 2174
## Warning in gregexpr(pattern, x, per1 = TRUE): PCRE error
## 'UTF-8 error: isolated byte with 0x80 bit set'
## for element 3995
```

```
## Warning in gregexpr(pattern, x, perl = TRUE): PCRE error
  'UTF-8 error: isolated byte with 0x80 bit set'
## for element 4052
## Warning in gregexpr(pattern, x, per1 = TRUE): PCRE error
  'UTF-8 error: isolated byte with 0x80 bit set'
  for element 4107
## Warning in gregexpr(pattern, x, perl = TRUE): PCRE error
## 'UTF-8 error: isolated byte with 0x80 bit set'
## for element 4169
## Warning in gregexpr(pattern, x, perl = TRUE): PCRE error
## 'UTF-8 error: isolated byte with 0x80 bit set'
## for element 4233
## Warning: Expected 2 pieces. Missing pieces filled with `NA` in 5781 rows [1, 2,
## 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, ...].
# new5$DC_left_1 %>% unique()
# new5$DC_left_r %>% unique()
## now work on DC_right
head(new5$DC_right %>% unique(), n=20)
## [1] NA
## [2] "FUNGICIDE: (BOSCALID = 128008)"
  [3] "FUNGICIDE: (CYPRODINIL = 288202)"
  [4] "FUNGICIDE: (FLUDIOXONIL = 71503)"
   [5] "FUNGICIDE: (MYCLOBUTANIL = 128857)"
  [6] "FUNGICIDE: (PYRACLOSTROBIN = 99100)"
   [7] "FUNGICIDE: (TOTAL)"
  [8] "HERBICIDE: (TOTAL)"
  [9] "INSECTICIDE: (ACEQUINOCYL = 6329)"
## [10] "INSECTICIDE: (BIFENAZATE = 586)"
## [11] "INSECTICIDE: (METHOXYFENOZIDE = 121027)"
## [12] "INSECTICIDE: (PYRETHRINS = 69001)"
## [13] "INSECTICIDE: (SPINETORAM = 110007)"
## [14] "INSECTICIDE: (SPINOSAD = 110003)"
## [15] "INSECTICIDE: (TOTAL)"
## [16] "INSECTICIDE: (ZETA-CYPERMETHRIN = 129064)"
## [17] "OTHER: (TOTAL)"
## [18] "INSECTICIDE: (BT KURSTAKI ABTS-351 = 6522)"
## [19] "FUNGICIDE: (AZOXYSTROBIN = 128810)"
## [20] "FUNGICIDE: (BLAD = 30006)"
```

```
new5 %<>% separate(DC_right, c("DC_right_1", "DC_right_r"), sep = ": ")

new5[is.na(new5)] <- " "

# OK now we need to eliminate the redundancy
# fine and remove redundant columns

# remove column new5$DC_left_1
new5 %<>% select(-DC_left_1)

# remove column DC_right_1
new5 %<>% select(-DC_right_1)

# remove "Chemical" and joint the columns
new5 %<>% mutate(D_left = "CHEMICAL", D_left = "")
new5 %<>% mutate(Chemical=paste(D_left, D_right))

new5 %<>% select(-c(D_left, D_right))

# Final tidy data
new5$DC_left_r %>% unique() # rename chemical_family
```

```
## [1] " " " (NITROGEN)" "(PHOSPHATE)" "(POTASH)" "(SULFUR)"
```

```
new5 %<>% rename( Chem_family = DC_left_r, Materials = DC_right_r)
new5 %<>% mutate(Chemical = str_trim(paste(Chem_family, Chemical)))
new5 %<>% select(-c(`Data Item`, Chem_family))
new5 %<>% rename( Type = type, Unit = unit, Production = production)
new5 %<>% select(Year, Period, State, Commodity, Type, Production, Chemical, Materials, Unit, Value)
Tidyberry <- new5
# write. table(Tidyberry, "D:/MSSP/Rdata/615/Berry/Tidyberry.csv", col. names = TRUE, row. name = FAL
SE, sep = ",")
#This Tidyberry contains all berries.</pre>
```

## Filter some data from Tidyberry

The data is tidy now. The majority of production is for application, so I filter rows of data in some situation.

```
#I choose raspberries and period=YEAR
Rberry <- Tidyberry %>% filter((Commodity=="RASPBERRIES") & (Period=="YEAR"))
Rberry %<>% select(-c(Period, Commodity))
## look at chemicals being applied to food
unfood <- Rberry %>% filter(Production=="APPLICATIONS")
unfood %<>% filter(Value != "(NA)")
#unique(unfood$Unit)
#in this case, I choose unit=AVG.
unfood %<>% filter(Unit == "AVG")
unfood$Value <- as.numeric(unfood$Value)

unfood_1 <- unfood %>% select(Year, State, Chemical, Value)
unfood_1$Value <- as.numeric(unfood_1$Value)
unfood_1 %<>% pivot_wider(names_from = Chemical, values_from = Value)
```

```
## Warning: Values are not uniquely identified; output will contain list-cols.
## * Use `values_fn = list` to suppress this warning.
## * Use `values_fn = length` to identify where the duplicates arise
## * Use `values_fn = {summary_fun}` to summarise duplicates
```

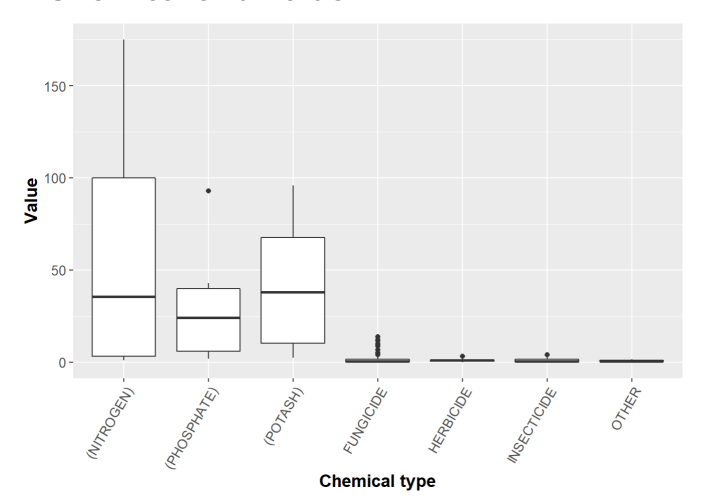
```
## Because of using the pivot_wider, some data was a list. Change to sum.
for (i in 1:6) {
  f <- as.numeric(unlist(unfood_1$INSECTICIDE[i]))</pre>
  unfood 1$INSECTICIDE[i] <- sum(f)</pre>
for (i in 1:6) {
  f <- as.numeric(unlist(unfood 1$FUNGICIDE[i]))</pre>
  unfood 1$FUNGICIDE[i] <- sum(f)</pre>
for (i in 1:6) {
  f <- as.numeric(unlist(unfood 1$HERBICIDE[i]))</pre>
  unfood_1$HERBICIDE[i] <- sum(f)
for (i in 1:6) {
  f <- as.numeric(unlist(unfood 1$OTHER[i]))</pre>
  unfood_1$OTHER[i] <- sum(f)
for (i in 1:6) {
  f <- as. numeric (unlist (unfood 1$`(NITROGEN)`[i]))
  unfood_1$`(NITROGEN)`[i] <- sum(f)</pre>
for (i in 1:6) {
  f <- as.numeric(unlist(unfood 1$`(PHOSPHATE)`[i]))</pre>
  unfood 1$`(PHOSPHATE)`[i] <- sum(f)
for (i in 1:6) {
  f <- as.numeric(unlist(unfood 1$`(POTASH)`[i]))</pre>
  unfood 1$`(POTASH)`[i] <- sum(f)
unfood_1$FUNGICIDE <- as.numeric(unfood_1$FUNGICIDE)</pre>
unfood_1$INSECTICIDE <- as.numeric(unfood_1$INSECTICIDE)</pre>
unfood_1$HERBICIDE <- as.numeric(unfood_1$HERBICIDE)</pre>
unfood_1$0THER <- as.numeric(unfood_1$0THER)</pre>
unfood 1$`(NITROGEN)` <- as.numeric(unfood 1$`(NITROGEN)`)</pre>
unfood 1$`(PHOSPHATE)` <- as.numeric(unfood 1$`(PHOSPHATE)`)</pre>
unfood_1$`(POTASH)` <- as.numeric(unfood_1$`(POTASH)`)</pre>
#kable(head(Rberry, n=10)) %>% kable_styling(fixed_thead = T, font_size = 10)
head (Rberry, n=10)
```

Y State <dbl><chr></chr></dbl>	Type <chr></chr>	Production <chr></chr>	Chemical <chr></chr>	Materials <chr></chr>	Unit <chr></chr>
2019 CALIFORN	IIA				ACRES HAF
2019 CALIFORN	IIA				MEASURED
2019 CALIFORN	IIA				MEASURED
2019 CALIFORN	IIA BEARIN	IGAPPLICATION	IS FUNGICIDI	E (BOSCALID = 128008)	MEASURED
2019 CALIFORN	IIA BEARIN	IGAPPLICATION	IS FUNGICIDI	E (CYPRODINIL = 288202)	MEASURED
2019 CALIFORN	IIA BEARIN	IGAPPLICATION	IS FUNGICIDI	E (FLUDIOXONIL = 71503)	MEASURED

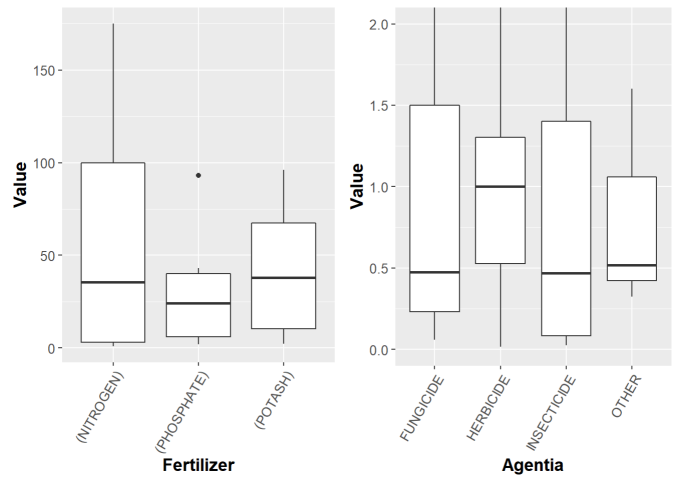
	State > <chr></chr>	Type <chr></chr>	Production <chr></chr>	Chemical <chr></chr>	Materials <chr></chr>	Unit <chr></chr>
2019	O CALIFORNIA	BEARING	SAPPLICATIONS	FUNGICIDE	(MYCLOBUTANIL = 128857)	MEASURED
2019	O CALIFORNIA	BEARING	SAPPLICATIONS	FUNGICIDE	(PYRACLOSTROBIN = 9910	0) MEASURED
2019	O CALIFORNIA	BEARING	SAPPLICATIONS	FUNGICIDE	(TOTAL)	MEASURED
2019	O CALIFORNIA	BEARING	SAPPLICATIONS	HERBICIDE	(TOTAL)	MEASURED
1-10	of 10 rows   1-7	of 8 colun	nns			
4						<b>&gt;</b>

## **EDA**

# 1. Chemical and Value



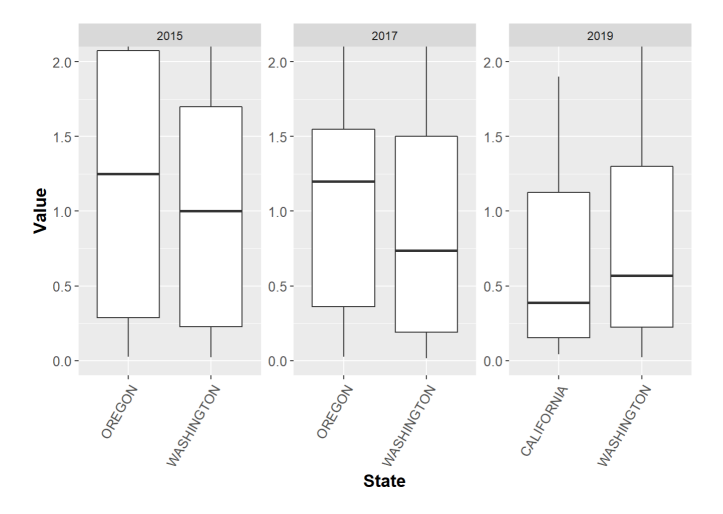
In this case, I separate the chemical into to part: agentia and fertilizer. I create an indicate to distinguish them. Two part are drawn separately, but I use grid.arrange function to combine them into same plot. It is clear that the value of raspberry using fertilizer('nitrogen', 'phosphate' and 'potash') is much higher than those using agentia('fungicide', 'herbicide', 'insecticide' and 'other')(because 'other' also have a small range so I put it in the agentia).



And also can see those outliers here.

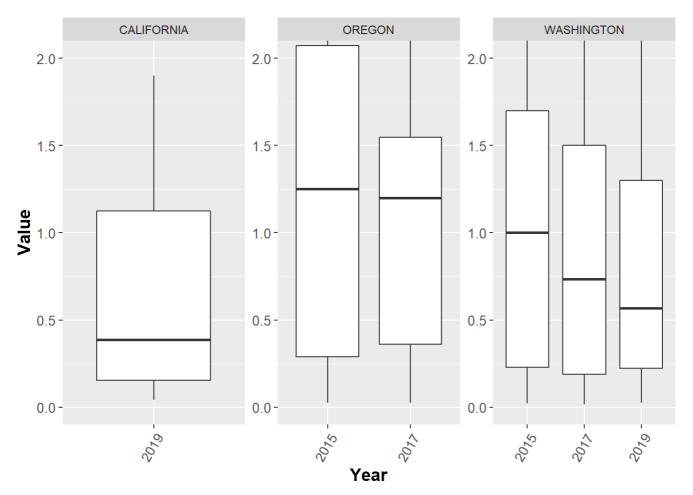
Y State <dbl><chr></chr></dbl>	Type <chr></chr>	Production <chr></chr>	Chemical <chr></chr>	Materials <chr></chr>	<b>U</b> <chr></chr>
2017 OREGON	BEARIN	GAPPLICATIONS	FUNGICIDE	(CALCIUM POLYSULFIDE = 76702)	AVG
2017 OREGON	BEARIN	GAPPLICATIONS	FUNGICIDE	(CALCIUM POLYSULFIDE = 76702)	AVG
2017 OREGON	BEARIN	GAPPLICATIONS	FUNGICIDE	(CAPTAN = 81301)	AVG
2017 WASHINGTO	)MBEARIN	GAPPLICATIONS	FUNGICIDE	(CALCIUM POLYSULFIDE = 76702)	AVG
2017 WASHINGTO	)MBEARIN	GAPPLICATIONS	FUNGICIDE	(CALCIUM POLYSULFIDE = 76702)	AVG
2017 WASHINGTO	)MBEARIN	GAPPLICATIONS	FUNGICIDE	(CAPTAN = 81301)	AVG
2017 WASHINGTO	)MBEARIN	GAPPLICATIONS	FUNGICIDE	(CAPTAN = 81301)	AVG
2015 OREGON	BEARIN	GAPPLICATIONS	FUNGICIDE	(CALCIUM POLYSULFIDE = 76702)	AVG
2015 OREGON	BEARIN	GAPPLICATIONS	FUNGICIDE	(CALCIUM POLYSULFIDE = 76702)	AVG
2015 WASHINGTO	)MBEARIN	GAPPLICATIONS	FUNGICIDE	(CALCIUM POLYSULFIDE = 76702)	AVG
1-10 of 10 rows   1-	8 of 10 col	umns			
4					•

# 2.state and value



# 3. Year and Value

The dataset only have one year of value in California so there is no much to discuss.



In 2015, the median of value of Oregon and Washington was close, around 1.1. In 2017, the median of value in Oregon is almost two times of that in Washington. In 2019, there was a small difference between the situation of California and Washington. From 2015 to 2017, the median of value of Oregon almost did not change, but the top value decreased. The median of value of Washington decreased by around 0.25. From 2015 to 2019, the median of value of Washington and range were continuously decreased.

## **PCA**

# Correlations between chemical type.

```
## Importance of components:

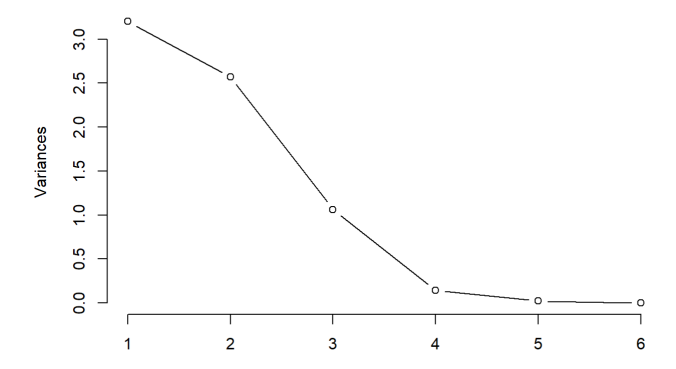
## PC1 PC2 PC3 PC4 PC5 PC6

## Standard deviation 1.7892 1.6033 1.0311 0.37724 0.15084 2.362e-16

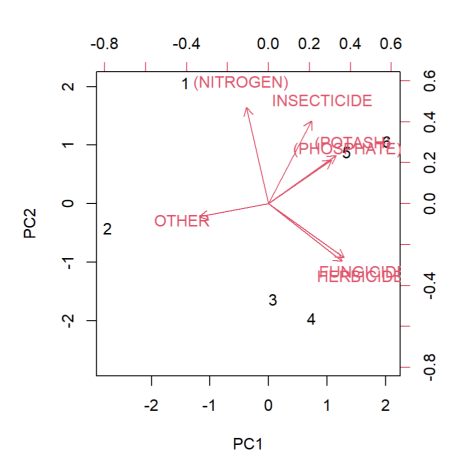
## Proportion of Variance 0.4573 0.3672 0.1519 0.02033 0.00325 0.000e+00

## Cumulative Proportion 0.4573 0.8245 0.9764 0.99675 1.00000 1.000e+00
```

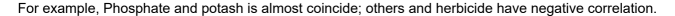


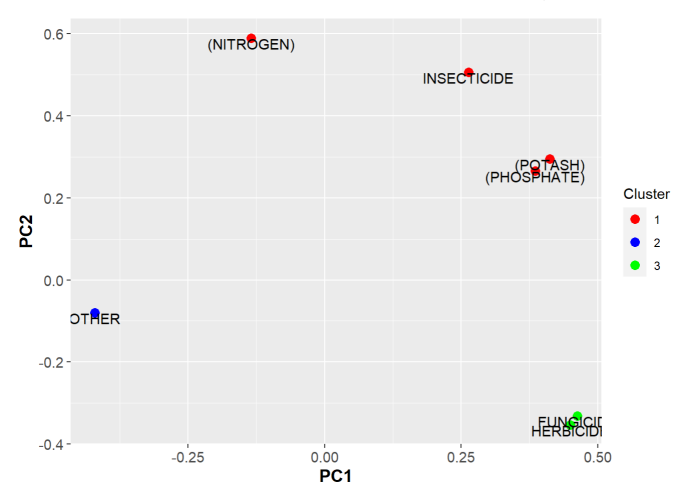


The PCA provides 6 components and 99% of the total variance is attributed to the first 4 components.



And in the biplot I can see the relationship between each variables. The size of the angle between vectors determines the correlation of the variables, which is the desired indicator to achieve the objective for this analysis. A small angle indicates a strong positive correlation, 90 degrees represents no correlation and 180 degrees represents a negative correlation.





In this plot, it is clear that chemical can be separate into to three part, just I mentioned before, apart from 'other', fertilizer('nitrogen', 'phosphate' and 'potash') gathered as cluster1, and agentia('fungicide', 'herbicide', 'insecticide') gathered as cluster3.

#### Conclusion

Honestly speaking, this project is a big and painful job for me as a rookie. I even did not know how to begin. But after discussing with others and having classes, I went into condition gradually. In this process, I found a big problem: I had some ideas to handle the problems but I can not put my theoretical ideas into practice. In balance, my R skill was too terrible to handle. So I tried to use internet and textbooks to improve my skill. This method was painful because I always made mistake and I even wanted to punch my laptop when R reported errors. But I do learn a lot about no matter in data cleaning, EDA, PCA or shiny.

### Citation

- [1] Exploratory data analysis into the relationship between different types of crime in London (https://towardsdatascience.com/exploratory-data-analysis-into-the-relationship-between-different-types-of-crime-in-london-20c328e193ff)
- [2] R for Data Science (https://r4ds.had.co.nz/)
- [3] dmorison/eda-relationships-between-crime-london (https://github.com/dmorison/eda-relationships-between-crime-london)
- [4] PCA.rmd (published%20in%20BB) [5] ag\_data(2).rmd (published%20in%20BB)
- [6] [many of my classmates](I discuss with them a lot. For example, Hao Shen and I discussed problem in data cleaning. He mentioned that we could use # to replace value of "Data Item". It's a very good way to separate Data Item by using sep="#". By contrast, my way was clumsy. Also, Zhe Yu gave me many advice about EDA and PCA.)