Exploratory Analysis

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Libraries

Reading in Data

```
setwd("~/harvard-summer-biostats")
illinois <- read_csv("data/illinois.csv") #read in data</pre>
## Warning: Missing column names filled in: 'X1' [1]
## Parsed with column specification:
## cols(
##
     X1 = col_double(),
##
     state = col_character(),
     site = col_character(),
##
    disposal.area = col_character(),
     type = col_character(),
##
##
     well.id = col_character(),
##
     gradient = col_character(),
     samp.date = col_character(),
##
     contaminant = col character(),
##
     measurement.unit = col_character(),
##
##
     concentration = col_double()
## )
```

Checking unique/distinct levels

```
#check how many observations there
nrow(illinois)

## [1] 38792

#checking distincts sites
unique(illinois$site)

## [1] "Baldwin Energy Complex"
## [2] "Coffeen Power Station"
```

```
## [3] "Dallman Power Generating Station"
## [4] "Duck Creek Power Station"
## [5] "Edwards Power Station"
## [6] "Havana Power Station"
   [7] "Hennepin Power Station"
## [8] "Joliet #29 Generating Station"
## [9] "Joliet #9 Generating Station"
## [10] "Joppa Power Station"
## [11] "Kincaid Power Station"
## [12] "Newton Power Station"
## [13] "Powerton Generating Station"
## [14] "Prairie State Generating Company, LLC"
## [15] "SIPC Marion Power Plant"
## [16] "Waukegan Station"
## [17] "Will County"
## [18] "Wood River Power Station"
#checking distinct gradient
unique(illinois$gradient)
## [1] "Downgradient" "Upgradient"
                                     "Unknown"
#checking distinct gradient
unique(illinois$measurement.unit)
## [1] "mg/l" "su"
                       "pCi/l" "ug/l"
#checking disposal area
unique(illinois$disposal.area)
##
  [1] "Baldwin Bottom Ash Pond"
  [2] "Baldwin Fly Ash Pond System"
## [3] "Baldwin Bottom Ash Pond, Baldwin Fly Ash Pond System"
   [4] "Coffeen Ash Pond No. 2"
## [5] "Coffeen Ash Pond No. 1"
  [6] "Coffeen GMF Recycle Pond"
  [7] "Coffeen GMF Gypsum Stack Pond"
##
## [8] "Coffeen GMF Gypsum Stack Pond, Coffeen Landfill"
## [9] "Coffeen Ash Pond No. 1, Coffeen Ash Pond No. 2"
## [10] "Coffeen Landfill"
## [11] "Coffeen Ash Pond No. 2, Coffeen GMF Recycle Pond"
## [12] "Dallman Ash Pond, Lakeside Ash Pond"
## [13] "Duck Creek Bottom Ash Basin"
## [14] "Duck Creek Landfill"
## [15] "Duck Creek GMF Pond"
## [16] "Edwards Ash Pond"
## [17] "Havana East Ash Pond (Cells 1, 2, 3, and 4)"
## [18] "Hennepin Ash Pond No. 2, Hennepin East Ash Pond, Hennepin Landfill"
## [19] "Henepin Old West Ash Pond (Pond No. 1 and Pond No. 3) and Hennepin Old West Polishing Pond"
## [20] "Hennepin Ash Pond No. 2"
## [21] "Hennepin East Ash Pond"
## [22] "Hennepin Landfill"
```

```
## [23] "Ash Pond 2"
## [24] "Lincoln Stone Quarry"
## [25] "Joppa Landfill"
## [26] "Joppa East Ash Pond"
## [27] "Kincaid Ash Pond"
## [28] "Newton Landfill 2"
## [29] "Newton Primary Ash Pond"
## [30] "Ash By-pass Basin, Ash Surge Basin, Former Ash Basin"
## [31] "Near Field Landfill"
## [32] "Settling Pond"
## [33] "West Ash Pond, East Ash Pond"
## [34] "Ash Pond 2 South, Ash Pond 3 South"
## [35] "Wood River West Ash Ponds 1, 2E, 2W"
## [36] "Wood River Primary East Ash Pond"
#checking type (L = land\ fuel, SI = surface\ impoundement, M = mixed/multiunit)
#there will be multiple different dumpsites at a powerplant site. some sites are wet (ponds of water wh
unique(illinois$type)
## [1] "SI" "M" "L"
illinois %>%
  group_by(type) %>%
  summarize(n())
## # A tibble: 3 x 2
   type `n()`
    <chr> <int>
## 1 L
            8869
## 2 M
             875
## 3 SI
           29048
illinois %>%
  group_by(measurement.unit) %>%
  summarize(n())
## # A tibble: 4 x 2
    measurement.unit `n()`
##
   <chr>
                      <int>
## 1 mg/l
                      32526
## 2 pCi/l
                      1721
## 3 su
                       1964
## 4 ug/l
                       2581
#checking wells
illinois %>%
  group_by(well.id) %>%
 summarize(n())
## # A tibble: 198 x 2
## well.id `n()`
```

```
##
      <chr>
              <int>
##
    1 03R
                175
##
   2 05DR
                175
##
   3 05R
                175
##
   4 08D
                175
##
   5 12
                175
   6 13
##
                175
  7 18D
##
                175
##
   8 18S
                172
## 9 2
                174
## 10 21
                350
## # ... with 188 more rows
```

Tony: Molybdenum to Thallium

use illinois and newyork

lower detection limit (if a chemical is really low, it can't differentiate beyond a certain low level, it might be the values with all teh same values)

all of the contaminants are harmful, some chemicals are more useful for detecting coal contamination (most important/indicative of coal contamination is boron (occurs at really high levels in coal)),

looking at the report given to us might be good about the different chemicals in coal (if we are curious)

total dissolved solids are probably not important, imprevise measurement of how much "crap is in the water" - doesnt represent any toxic chemicals

Wrangling

```
illinois1 <- illinois %>%
  select(site, disposal.area, type, well.id, gradient, contaminant,
         measurement.unit, concentration) %>%
  mutate(well.id_contaminant = paste0(well.id, "_", contaminant)) %>% #for future use
  rename(c("disposal_area" = "disposal.area", "well_id" = "well.id",
           "unit" = "measurement.unit"))
#fixing 'contaminant' string by removing everything after the comma
illinois1$contaminant=gsub(", total", "", illinois1$contaminant)
#testing
avg_contaminant <- illinois1 %>%
  group_by(well_id, contaminant) %>%
  summarise_each(funs(mean)) %>%
  select(1,2,8) #selecting only numeric columns
#temporarily uniting columns for joining in next step
temp <- avg_contaminant %>%
  unite("well.id_contaminant", well_id, contaminant)
#joining orig dataframe and avg_contaminant dataframe
combined <- left_join(temp, illinois1, by = "well.id_contaminant") %>%
  distinct(well.id_contaminant, .keep_all = TRUE) %>%
  separate(well.id_contaminant, c('well_id', 'contaminant'), sep="_") %>%
```

```
#spreading to wide data frame format to add missing info
combined2 <- combined %>% #collapse empty rows
    spread(contaminant, concentration.x) %>%
    group_by(well_id) %>%
    summarise_each(funs(first(.[!is.na(.)]))) %>%
    select(-c(unit))

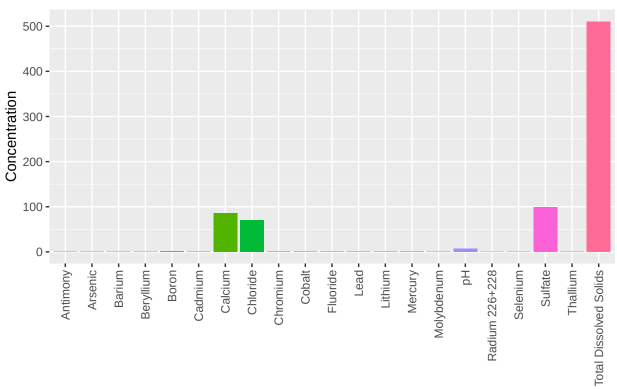
#gathering back to long data frame format
combined3 <- combined2 %>%
    gather("contaminant", "concentration", 6:26)
```

Summary Statistics

```
illinois %>%
  group_by(contaminant) %>%
  summarize(mean_concentration = mean(concentration))
## # A tibble: 21 x 2
##
     contaminant mean_concentration
##
     <chr>
                                  <dbl>
## 1 Antimony, total
                                 0.626
## 2 Arsenic, total
                                2.56
## 3 Barium, total
                               39.8
## 4 Beryllium, total
                                0.141
## 5 Boron, total
                              125.
## 6 Cadmium, total
                                 0.141
## 7 Calcium, total
                             7460.
## 8 Chloride
                                79.1
## 9 Chromium, total
                                 0.969
## 10 Cobalt, total
                                 0.761
## # ... with 11 more rows
```

Visualizations

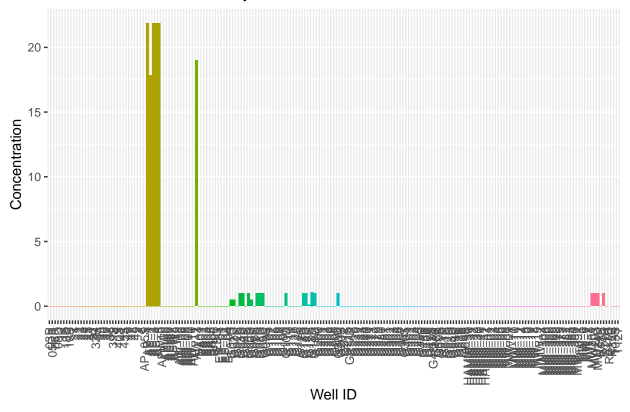
Concentration of Contaminants in Well 03R



Contaminant

Warning: Removed 1 rows containing missing values (position_stack).

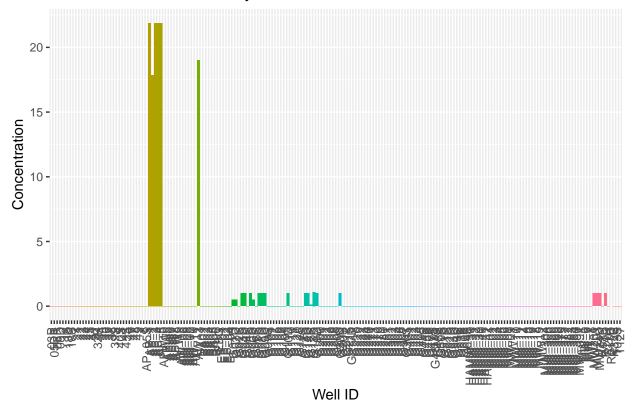
Concentration of Antimony across Wells



The histogram is incompehensible, there are way too many wells in Illinois. Unsure, but having to make all these plots constantly seems unreasonable for all of these potential wells and contaminants to look through, maybe choose only those wells with concentration values greater than some certain value? (0.001 maybe?)

Warning: Removed 1 rows containing missing values (position_stack).

Concentration of Antimony across Wells



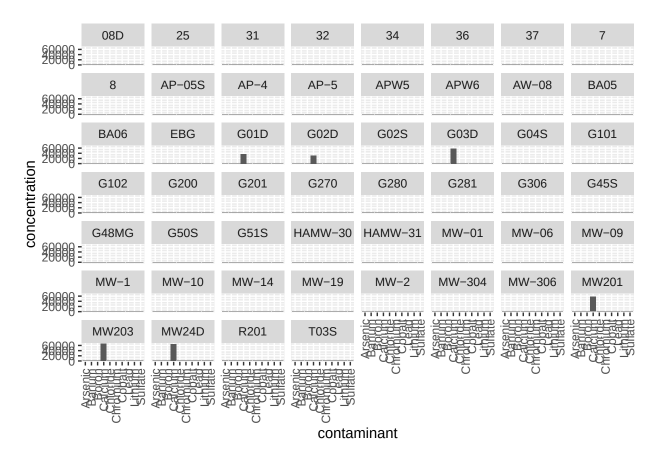
levels(as.factor(combined3\$well_id))

##	[1]	"03R"	"05DR"	"05R"	"08D"	"12"	"13"	"18D"
##	[8]	"18S"	"2"	"21"	"22"	"23"	"24"	"25"
##	[15]	"31"	"32"	"32R"	"34"	"35"	"36"	"37"
##	[22]	"38"	"395"	"4"	"40S"	"41"	"45S"	"46"
##	[29]	"47"	"48"	"49"	"7"	"8"	"AP-05S"	"AP-1"
##	[36]	"AP-2"	"AP-3"	"AP-4"	"AP-5"	"APW10"	"APW5"	"APW6"
##	[43]	"APW7"	"APW8"	"APW9"	"AW-05"	"AW-06"	"AW-08"	"AW-09"
##	[50]	"AW-10"	"AW-11"	"AW-3"	"BA01"	"BA02"	"BA03"	"BA04"
##	[57]	"BA05"	"BA06"	"EBG"	"EP-01"	"EP-02"	"EP-03"	"EP-04"
##	[64]	"G01D"	"G02D"	"G02S"	"G03D"	"G04D"	"G04S"	"G05D"
##	[71]	"G06D"	"G06S"	"G07D"	"G08D"	"G09D"	"G09S"	"G101"
##	[78]	"G102"	"G105"	"G106"	"G107"	"G109"	"G10D"	"G110"
##	[85]	"G111"	"G120"	"G125"	"G12S"	"G13D"	"G15D"	"G15S"
##	[92]	"G17D"	"G19D"	"G200"	"G201"	"G202"	"G203"	"G206"
##	[99]	"G208"	"G209"	"G20D"	"G20S"	"G212"	"G215"	"G217D"
##	[106]	"G218"	"G220"	"G222"	"G223"	"G224"	"G270"	"G271"
##	[113]	"G273"	"G276"	"G279"	"G280"	"G281"	"G301"	"G302"
##	[120]	"G303"	"G304"	"G306"	"G307"	"G30S"	"G401"	"G402"
##	[127]	"G403"	"G404"	"G405"	"G44S"	"G45S"	"G46S"	"G47S"
##	[134]	"G48MG"	"G48S"	"G50S"	"G51D"	"G51S"	"G52D"	"G53D"
##	[141]	"G54D"	"G54S"	"G57S"	"G60S"	"G64S"	"HAMW-30"	"HAMW-31"
##	[148]	"HAMW-32"	"HAMW-39"	"HAMW-40"	"HAMW-41"	"HAMW-42"	"MW-01"	"MW-02"
##	[155]	"MW-03"	"MW-04"	"MW-05"	"MW-06"	"MW-08"	"MW-09"	"MW-1"

```
## [162] "MW-10"
                    "MW-11"
                              "MW-12"
                                        "MW-14"
                                                   "MW-15"
                                                             "MW-16"
                                                                        "MW-17"
                                        "MW-304"
                                                             "MW-356"
## [169] "MW-18"
                   "MW-19"
                              "MW-2"
                                                   "MW-306"
                                                                       "MW-366"
                   "MW-370"
## [176] "MW-369"
                              "MW-375"
                                        "MW-377"
                                                   "MW-382"
                                                             "MW-383"
                                                                       "MW-384"
## [183] "MW-390"
                                                   "MW-7"
                   "MW-391"
                              "MW-5"
                                        "MW-6"
                                                             "8-WM"
                                                                        "MW201"
## [190] "MW203"
                    "MW24D"
                              "R08S"
                                        "R11D"
                                                   "R201"
                                                             "R217D"
                                                                        "R32S"
## [197] "T03S"
                   "T127"
#Comparing Upgradient Wells
#qet the first 10 highest avg mg/l (not ph or total sulfate)
#not pH, total dissolve solids, or radium b/c different units
dont_use <- c("pH", "Radium 226+228", "Total Dissolved Solids")</pre>
#pull out the names (from top 10 high to low avg concentrations) of contaminants
chemicals <- combined3 %>%
    filter(contaminant != "pH") %>% #take our nonuseful contaminants
  filter(contaminant != "Radium 226+228") %>%
  filter(contaminant != "Total Dissolved Solids") %>%
    group_by(contaminant) %>%
    summarise(avg_conc = mean(concentration, na.rm = TRUE)) %>%
    arrange(desc(avg_conc)) %>%
  slice(1:10) %>% #there only seem to be 5 chemicals of nonzero(basically) values
    pull(contaminant)
chemicals
                                          "Chloride" "Barium"
    [1] "Calcium"
                   "Sulfate"
                               "Boron"
                                                                 "Arsenic"
    [7] "Chromium" "Lithium"
                               "Lead"
                                          "Cobalt"
combined3 %>%
    group by(contaminant) %>%
    summarise(avg_conc = mean(concentration))
## # A tibble: 21 x 2
```

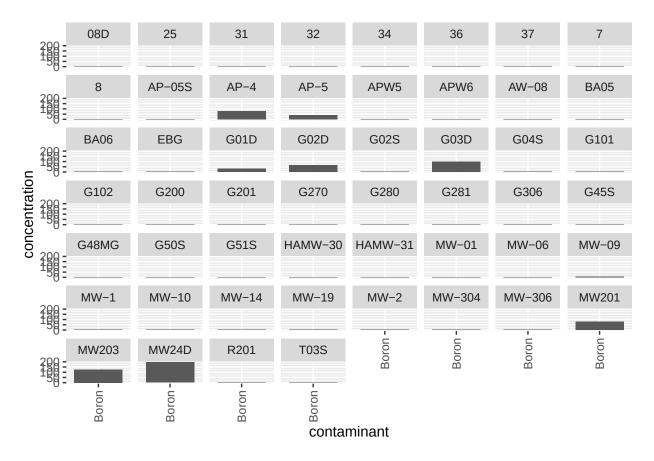
```
##
      contaminant avg conc
##
      <chr>
                     <dbl>
##
  1 Antimony
                      NA
##
  2 Arsenic
                      NA
##
   3 Barium
                      NA
## 4 Beryllium
                      NA
## 5 Boron
                     145.
## 6 Cadmium
                      NA
## 7 Calcium
                    8142.
## 8 Chloride
                      75.3
## 9 Chromium
                      NA
## 10 Cobalt
## # ... with 11 more rows
```

Is doing the mean of concentrations amongst all wells OK to do? Some wells like arsenic have wells with 0.001 (sign that measurement was insignificant/below threshold), and taking the avg sometimes makes it return NA as the mean (I THINK??? what does NA mean...?) ASK LULI (i think some of the chemicals in the paper had thresholds of DANGER that were in mg and others in microg, how would we account for this...?)



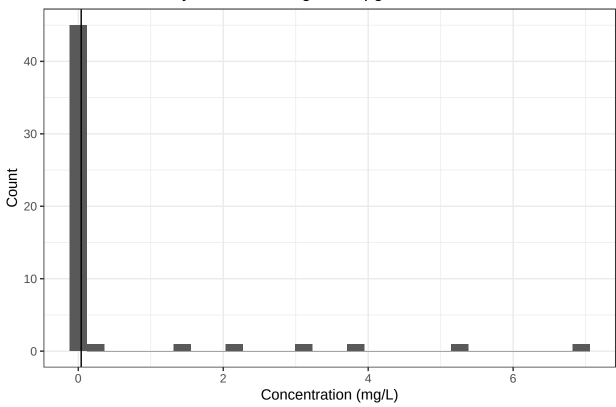
The concentration of calcium is so high in some of the wells, it makes it difficult to look at the other contaminants' concentration across all wells, perhaps we should make different facetted plots for all contaminants.

```
#Boron
ggplot(data = combined3 %>%
    filter(gradient == "Upgradient") %>%
    filter(contaminant %in% chemicals) %>%
    filter(contaminant == "Boron"),
    aes(x = contaminant, y = concentration)) +
    geom_bar(stat = "Identity") +
    facet_wrap(well_id ~ .) +
    theme(axis.text.x = element_text(angle = 90, vjust = 0.5, hjust=1))
```

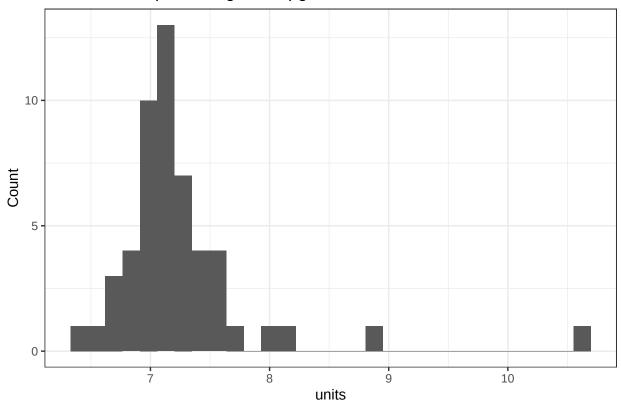


Let's make a histogram to show distribution of certain contaminants:

Distribution of Molybdenum amongst all Upgradient Wells

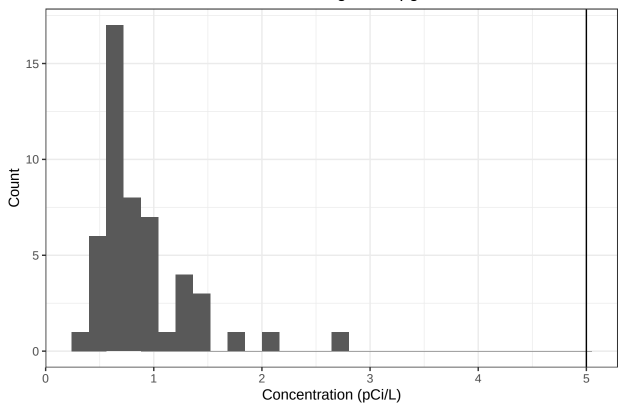


Distribution of pH amongst all Upgradient Wells

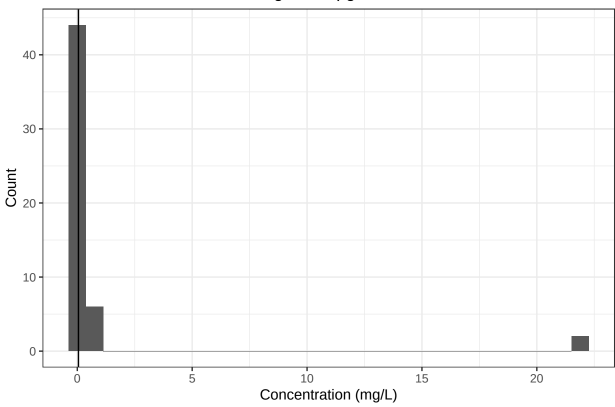


- ## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
- ## Warning: Removed 2 rows containing non-finite values (stat_bin).

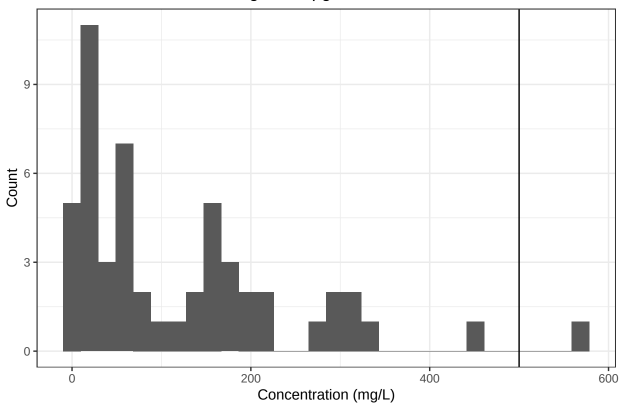
Distribution of Radium 226+228 amongst all Upgradient Wells



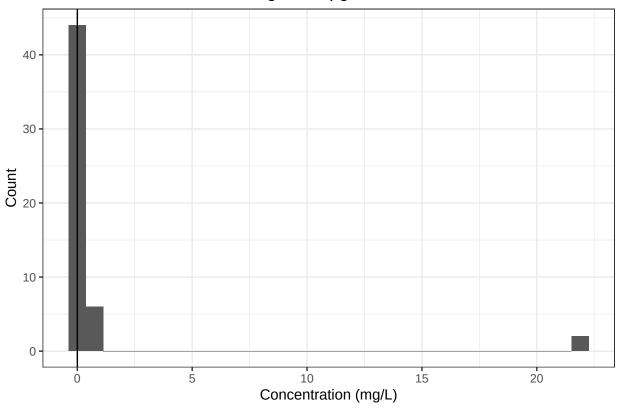
Distribution of Selenium amongst all Upgradient Wells



Distribution of Sulfate amongst all Upgradient Wells



Distribution of Thallium amongst all Upgradient Wells



We can see that there are some wells with concentrations that lie far beyond the threshold value (obtained from the PDF), and it would be nice to identify them. We will pull out the observations with concentrations past the threshold and place them within a new dataset.

```
#creating vector of contaminants
contaminant <- unique(combined3$contaminant)</pre>
#creating vector of threshold values for each contaminant
threshold <- c(6/1000, 10/1000, 2, 4/1000, 3, 5/1000, NA, NA, 100/1000,
                6/1000, NA, 15/1000, 40/1000, 2/1000, 40/1000, NA, 5,
               50/1000, 500, 2/1000, NA)
contam_t <- cbind(contaminant, threshold) %>%
  na.omit()
#creating function to obtain all observations with values above threshold
getOverThreshold <- function(df){</pre>
  datalist = list()
  for(i in 1:nrow(contam_t)){ #for each contaminant i
    df1 <- filter(df, gradient == "Upgradient")</pre>
    df2 <- filter(df1, contaminant == contam_t[i])</pre>
    data <- filter(df2, concentration > contam_t[nrow(contam_t) + i])
    datalist[[i]] <- data</pre>
  }
  toReturn <- do.call(rbind, datalist)</pre>
```

```
return(toReturn)
}

overthreshold_df <- getOverThreshold(combined3) #df with all observations over threshold value

#exporting data
write.csv(combined3,"data/wrangled_illinois.csv", row.names = FALSE)</pre>
```