Data Visualizations and Analysis

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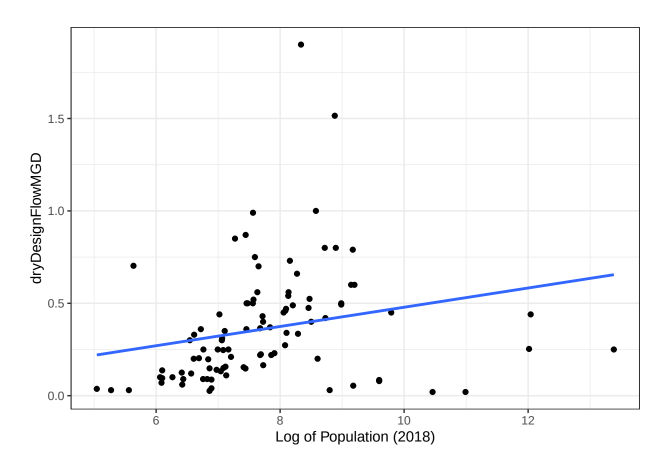
7/13/2020

```
ggplot(working_df, aes(x = log(pop_2018), y = dryDesignFlowMGD)) +
geom_point() +
geom_smooth(method = "lm", se = FALSE) +
theme_bw() +
labs(x = "Log of Population (2018)")
```

```
## 'geom_smooth()' using formula 'y ~ x'
```

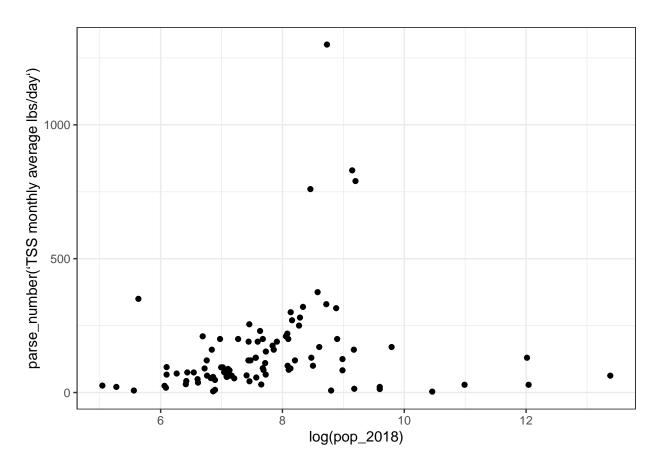
Warning: Removed 17 rows containing non-finite values (stat_smooth).

Warning: Removed 17 rows containing missing values (geom_point).



```
## Warning: 2 parsing failures.
## row col expected actual
## 13 -- a number na
## 45 -- a number na
## warning: 2 parsing failures.
## row col expected actual
## 13 -- a number na
## 45 -- a number na
```

Warning: Removed 19 rows containing missing values (geom_point).



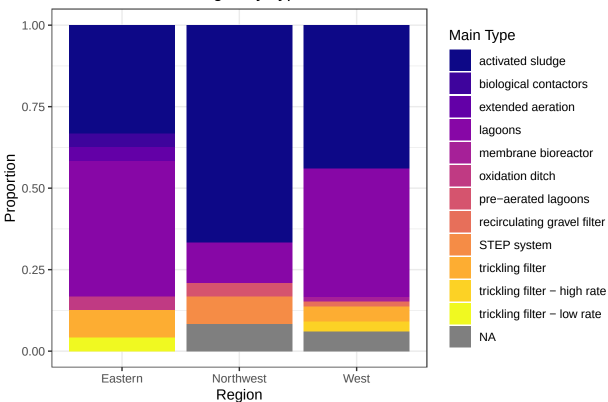
```
working_df %>%
  filter(type1 %in% c("lagoons", "activated sludge")) %>%
  group_by(type1) %>%
  summarize(median = median(pop_2018, na.rm = TRUE))
```

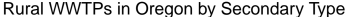
'summarise()' ungrouping output (override with '.groups' argument)

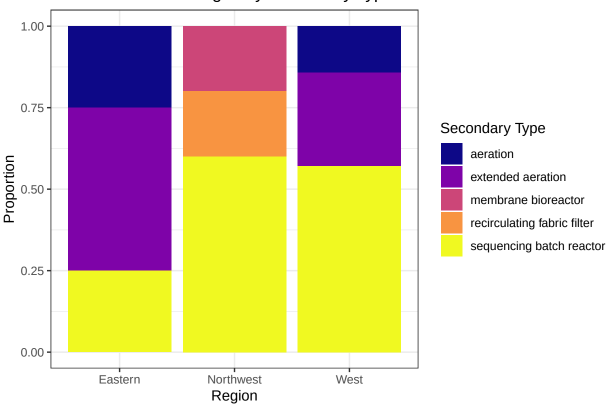
```
## # A tibble: 2 x 2
##
   type1
                     median
     <chr>
                      <dbl>
## 1 activated sludge 1962.
## 2 lagoons
                       1718.
library(viridis)
## Loading required package: viridisLite
library(plotly)
##
## Attaching package: 'plotly'
## The following object is masked from 'package:ggplot2':
##
##
       last_plot
## The following object is masked from 'package:stats':
##
##
       filter
## The following object is masked from 'package:graphics':
##
##
       layout
p <- working_df %>%
  ggplot(aes(x = Region.x,
            fill = type1)) +
  geom_bar(position = "fill") +
  scale_fill_viridis_d(option = "C", na.value = "grey50") +
  scale_x_discrete(labels=c("Eastern", "Northwest", "West")) +
  theme_bw() +
  labs(x = "Region",
      fill = "Main Type",
      y = "Proportion",
      title = "Rural WWTPs in Oregon by Type")
```

p

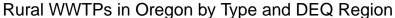
Rural WWTPs in Oregon by Type

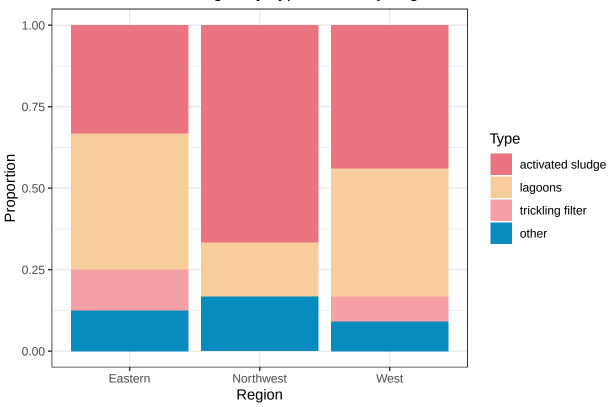






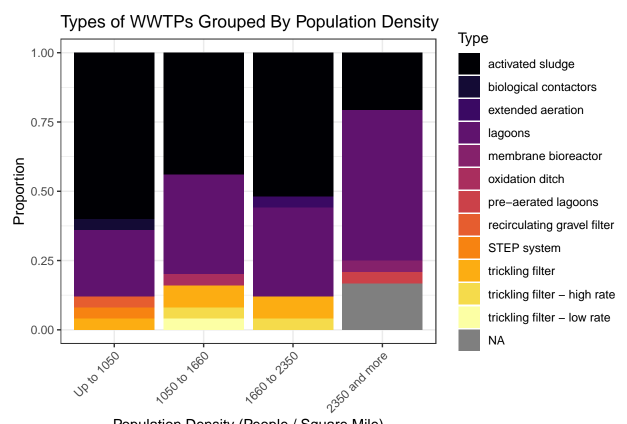
```
library(LaCroixColoR)
p4 <- working_df %>%
  mutate(
    type_plot = case_when(
      type1 %in% c("lagoons", "pre-aerated lagoons") ~ "lagoons",
      type1 %in% c("trickling filter", "trickling filter - high rate",
                   "trickling filter - low rate") ~ "trickling filter",
      type1 %in% c("activated sludge") ~ "activated sludge",
      type1 %in% c("extended aeration", "membrane bioreactor", "recirculating gravel filter", NA,
                   "STEP system", "oxidation ditch", "biological contactors"
    ) ~ "other"
  )) %>%
  ggplot(aes(x = Region.x,
             fill = factor(type_plot, levels = c("activated sludge", "lagoons", "trickling filter", "ot.
  geom_bar(position = "fill") +
  scale_fill_manual(values = lacroix_palette("Pamplemousse", type = "discrete")) +
  scale_x_discrete(labels=c("Eastern", "Northwest", "West")) +
  theme_bw() +
  labs(x = "Region",
       fill = "Type",
       y = "Proportion",
       title = "Rural WWTPs in Oregon by Type and DEQ Region")
  #theme(legend.position = "bottom")
p4
```



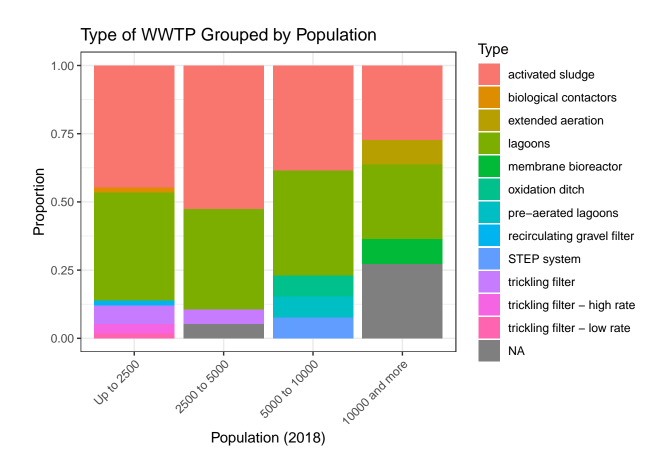


library(gtools)

Warning: package 'gtools' was built under R version 4.0.1



Population Density (People / Square Mile)



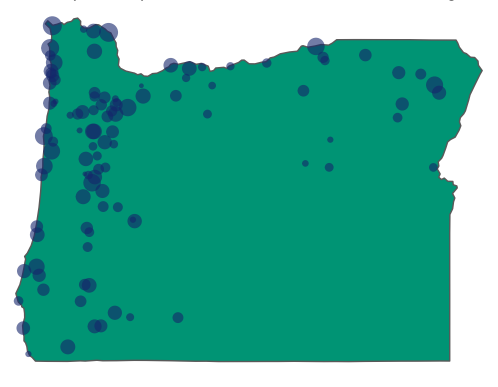
ggplotly(p1)

Maps:

library(sf)

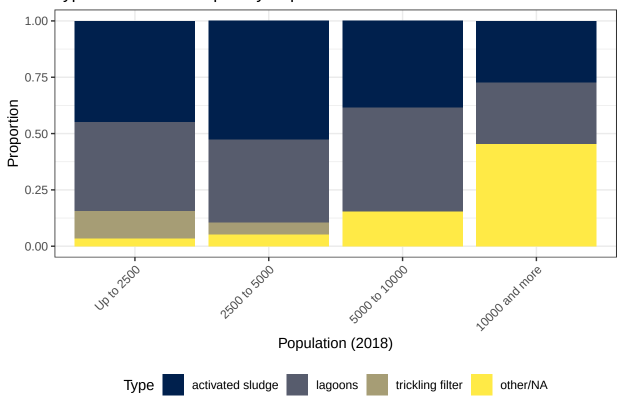
Linking to GEOS 3.7.2, GDAL 2.4.2, PROJ 5.2.0

Small (<1 MGD) Wastewater Treatment Facilities in Oregon



Dry Design Flow (MGD) 0.25 0.50 0.75 1.00

Type of WWTP Grouped by Population

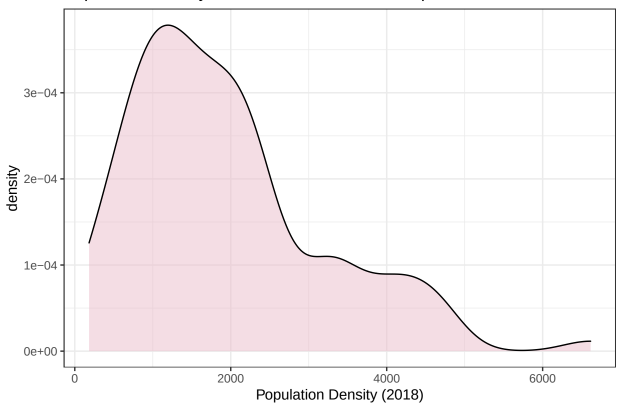


```
# do not run, does not work, crashes R
# try pop densities
# library(tidycensus)
# library(tigris)
#
# pop_block <- get_acs(
# geography = "block group",
# variables = "B01003_001",
# state = "OR",
# geometry = TRUE,
# key = "abac0e1ca2aa3d3ebb31d6d2fcdbaf52d3e25f7d"
# )
#</pre>
```

```
# area_2017 <-
# block_groups(year = 2017,
#
                 state = "OR",
#
                  class = "sf")
#
# area_2017 <- area_2017 %>%
# mutate(area = ALAND / 2589988) %>%
# select(area, geometry, GEOID)
#
# pop_block <- pop_block %>%
   select(estimate, geometry, GEOID)
\# class(pop\_block) \leftarrow "data.frame"
# class(area_2017) <- "data.frame"</pre>
# density <- left_join(pop_block, area_2017, by = c("GEOID" = "GEOID"))</pre>
# density <- density %>%
# mutate(pop_density = estimate/area)
# class(density) <- c("sf", "data.frame")</pre>
working_df %>%
```

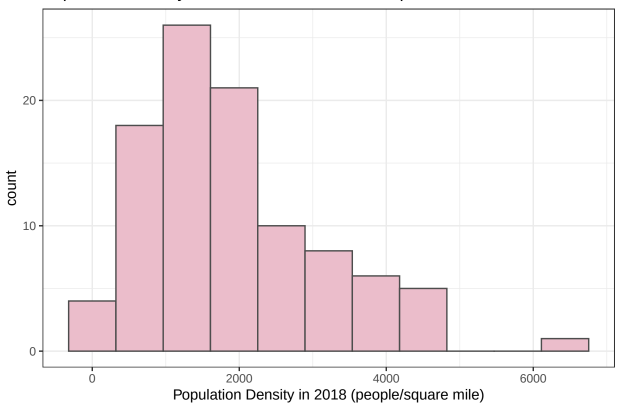
Warning: Removed 15 rows containing non-finite values (stat_density).

Population Density of Towns/Cities in Our Sample



Warning: Removed 15 rows containing non-finite values (stat_bin).

Population Density of Towns/Cities in Our Sample



```
working_df %>%
  filter(!is.na(basin)) %>%
  group_by(basin) %>%
  summarize(mgd = mean(dryDesignFlowMGD, na.rm = TRUE)) %>%
  ggplot(
    aes(x = reorder(basin, mgd), y = mgd)
) +
  geom_col(fill = "maroon") +
  coord_flip() +
  theme_bw() +
  labs(
    x = "Average Dry Design Flow (MGD)",
    y = "Basin",
    title = "Average Dry Design Flow, Grouped By Basin"
)
```

'summarise()' ungrouping output (override with '.groups' argument)



