### Calculated Info

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This package contains a couple tools to create useful views of the data not by default in the dataframe. These mostly come in the form of the default case and wastewater dataframe augment function.

## Wastewater augmeanting function

The wastewater function does three things. first it replaces Concentrations below the level of detection (LOD) with LOD / 2 which is considered best practice. Second it calculates a geometric average of the two gene concentrations. Finally it gets a flow and population normalized loged version of the data which previous work has shown the best results.

```
library(DSIWastewater)
## Registered S3 method overwritten by 'quantmod':
##
    method
                       from
##
     as.zoo.data.frame zoo
data(WasteWater_data, package = "DSIWastewater")
head(buildWasteAnalysisDF(WasteWater_data))
## # A tibble: 6 x 9
##
                                         flow geoMean sars_cov2_adj_load_l~1
     site
            date
                         pop
                                N1
                                      N2
     <chr> <date>
                       <dbl> <dbl> <dbl> <dbl>
                                                  <dbl>
                                                                         <dbl> <int>
## 1 Algoma 2020-10-06 3171 10000
                                      NA 0.498
                                                    NA
                                                                        NA
                                                                                   34
## 2 Algoma 2020-10-13
                        3171 10000 16500 0.499
                                                12845.
                                                                         0.306
                                                                                   34
## 3 Algoma 2020-10-20
                                                                         0.212
                        3171 10000 16500 0.402 12845.
                                                                                   34
## 4 Algoma 2020-10-27
                        3171 10000 16500 0.67
                                                 12845.
                                                                         0.434
                                                                                   34
                                                                         0.297
## 5 Algoma 2020-11-03 3171 10000 16500 0.489
                                                12845.
                                                                                   34
## 6 Algoma 2020-11-10 3171 10000 16500 0.463 12845.
                                                                         0.273
## # i abbreviated name: 1: sars_cov2_adj_load_log10
```

# Case augmeanting function

The case function does two things. first it normalizes the data by population. Second it calculates a rolling sum and average of the data. This needs the population data contained in its own data frame.

```
data(Case_data, package = "DSIWastewater")
data("pop_data", package = "DSIWastewater")#need to fix pop_data
#head(buildCaseAnalysisDF(left_join(Case_data, pop_data, by = "site")))
```

These functions are slightly out of date and not used in many new analysis projects

#TODO work on comparing the output of these methods