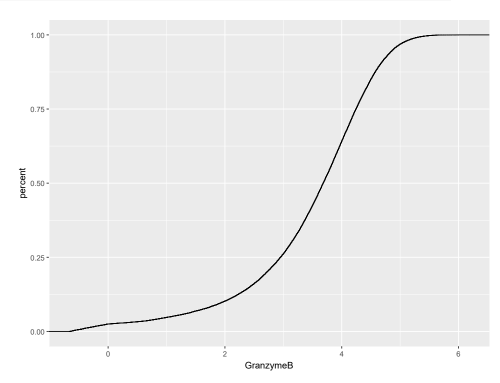
FA1

Baybayon, Darlyn Antoinette

```
library(ggplot2)
library(magrittr)
library(stringr)
library(gridExtra)
dataset <- read.csv("cytof_one_experiment.csv")
save(dataset, file="my_work_space.RData")</pre>
```

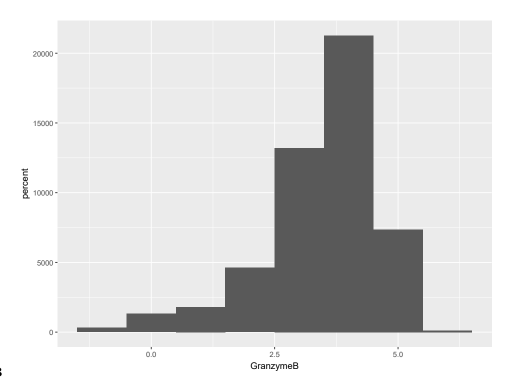
```
granzymeB <- dataset$GranzymeB
ggplot(dataset, aes(x = granzymeB)) + stat_ecdf() + xlab("GranzymeB") + ylab("percent")</pre>
```



ECDF plot for GranzymeB

This plot shows a slow increase at lower values, sharp rise in the middle. This means that most of the values in this column are concentrated in the middle and less extreme lower values.

```
ggplot(dataset, aes(x = granzymeB)) +geom_histogram(binwidth = 1) +
xlab("GranzymeB") + ylab("percent")
```

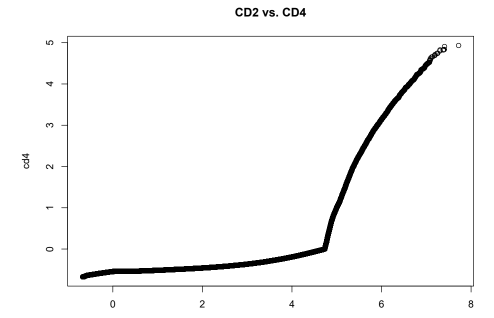


${\bf Histogram\ plot\ for\ GranzymeB}$

The histogram for granzymeB is skewed to the left which means that most of the values in this column are large and there are less lower values.

```
cd2 <- dataset$CD2
cd4 <- dataset$CD4

qqplot(cd2, cd4, main = "CD2 vs. CD4")</pre>
```



cd2

Q-Q plot for CD2 and CD4 $\,$

This plot deviates significantly from a straight diagonal line which implies that the quantiles do not have a similar distribution. Towards the right, CD4 values increase sharply compared to CD2 which suggests that it is more skewed to the right.