A sampling of bioinformatics using R

Abhijit Dasgupta, PhD

Spring 2019

Expectations

1. We will explore

- the basics of using Bioconductor packages for bioinformatics
- visualizations useful for bioinformatics
- ways to annotate our graphics

2. We will not explore

- how to do bioinformatics using R
- bioinformatic workflows
- data cleaning, modeling, analytics

This area is too broad and interests too varied to cover this with any sort of breadth or depth.

```
fix_names <- function(d) d %>% set_names(make.names(names(.)))
clinical <- rio::import('data/BreastCancer_Clinical.xlsx') %>% fix_names()
proteome <- rio::import('data/BreastCancer_Expression.xlsx') %>% fix_names()
final_data <- clinical %>%
    inner_join(proteome, by = c('Complete.TCGA.ID' = 'TCGA_ID')) %>%
    filter(Gender == 'FEMALE') %>%
    select(Complete.TCGA.ID, Age.at.Initial.Pathologic.Diagnosis, ER.Status, starts_with("NP"))
head(final_data)
```

```
Complete. TCGA. ID Age. at. Initial. Pathologic. Diagnosis ER. Status
#>
         TCGA-A2-A0CM
                                                     40 Negative
#> 1
#>
         TCGA-BH-A180
                                                     56 Negative
        TCGA-A7-A0CE
                                                     57 Negative
#>
#> 4
        TCGA-D8-A142
                                                     74 Negative
        TCGA-AO-A0J6
                                                        Negative
#>
         TCGA-A2-A0YM
                                                     67 Negative
#> 6
      NP_958782 NP_958785 NP_958786 NP_000436 NP_958781 NP_958780
   1 0.6834035 0.6944241 0.6980976 0.6870771 0.6870771 0.6980976
   2 0.1953407 0.2154129 0.2154129 0.2053768 0.2154129 0.2154129
   3 -1.1231731 -1.1231731 -1.1168605 -1.1294857 -1.1294857 -1.1200168
   4 0.5385958 0.5422105 0.5422105 0.5349810 0.5422105 0.5422105
   5 0.8311317 0.8565398 0.8565398 0.8367780 0.8650092 0.8565398
   6 0.6558497 0.6581426 0.6558497
                                     0.6558497 0.6512639 0.6581426
      NP 958783 NP 958784 NP 112598 NP 001611
   1 0.6980976 0.6980976 -2.6521501 -0.9843733
   2 0.2154129 0.2154129 -1.0357599 -0.5172257
   3 -1.1231731 -1.1231731 2.2445844 -2.5750648
   4 0.5422105 0.5422105 -0.1482049 0.2674902
```

. is the placeholder for what's specified inside the vars().

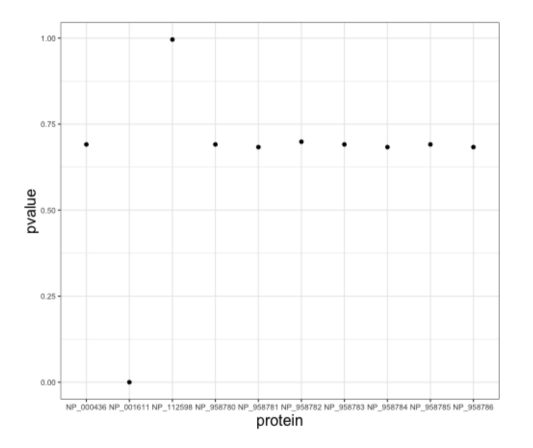
This isn't in the right format for me to plot

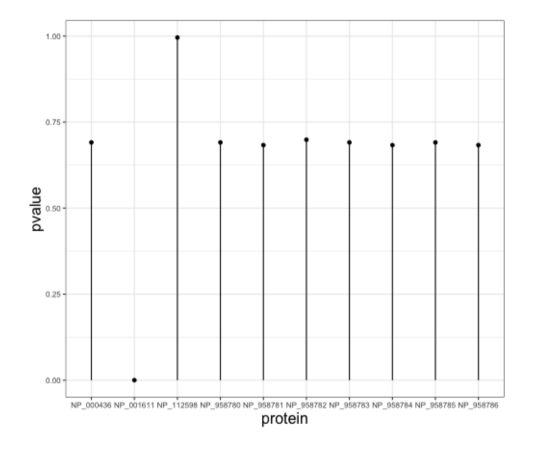
```
results %>% tidyr::gather(protein, pvalue)

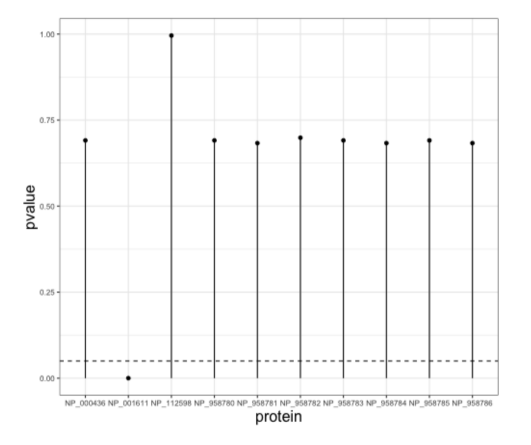
#> protein pvalue
#> 1 NP_958782 0.6988415415
#> 2 NP_958785 0.6910103484
#> 3 NP_958786 0.6832120796
#> 4 NP_000436 0.6910103484
#> 5 NP_958781 0.6832120796
#> 6 NP_958780 0.6910103484
#> 7 NP_958783 0.6910103484
#> 8 NP_958784 0.6832120796
#> 9 NP_112598 0.9957713985
#> 10 NP_001611 0.0001218627
```

```
theme_439 <- theme_bw() +
    theme(axis.title = element_text(size=16),
        axis.text = element_text(size=8))

results %>% tidyr::gather(protein, pvalue) %>%
    ggplot(aes(x = protein, y = pvalue)) +
        geom_point() +
    theme_439
```



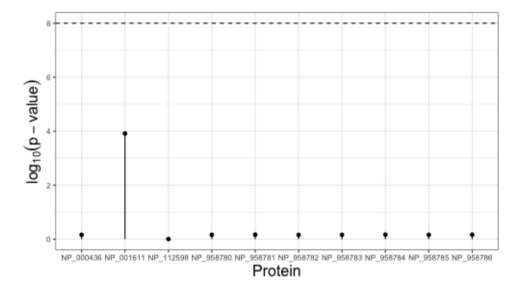




Manhattan plot

A Manhattan plot is used to visualize a set of p-values from unit-based tests

It plots the negative log p-value at each unit



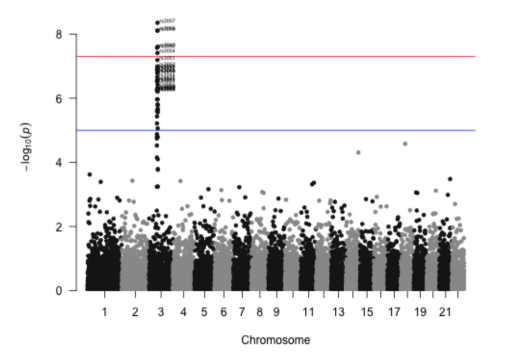
Manhattan plot

There is a specialized package for doing Manhattan plots and quantile plots for GWAS data

This package is meant to work with PLINK output, but the function is generic

```
library(qqman)
manhattan(gwasResults)
```

Manhattan plot



Data prep

```
head(gwasResults)
```

```
#> SNP CHR BP P

#> 1 rs1 1 1 0.9148060

#> 2 rs2 1 2 0.9370754

#> 3 rs3 1 3 0.2861395

#> 4 rs4 1 4 0.8304476

#> 5 rs5 1 5 0.6417455

#> 6 rs6 1 6 0.5190959
```

Data prep

```
plt_x_position <- gwasResults %>%
    group_by(CHR) %>%
    summarize(chr_len = max(BP)) %>%
    mutate(tot = cumsum(chr_len) - chr_len)
head(plt_x_position)
```

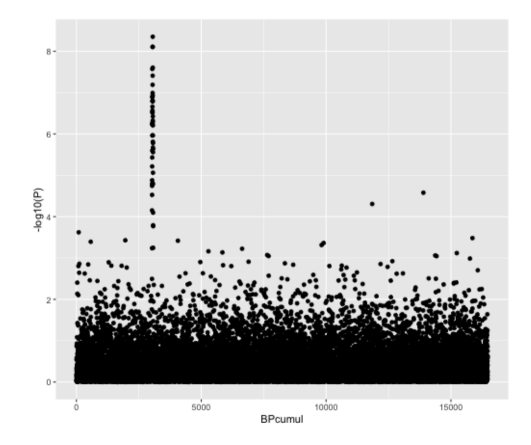
```
#> # A tibble: 6 x 3
#> CHR chr_len tot
#> <int> <int> <int>
#> 1     1     1500     0
#> 2     2     1191     1500
#> 3     3     1040     2691
#> 4     4     945     3731
#> 5     5     877     4676
#> 6     6     825     5553
```

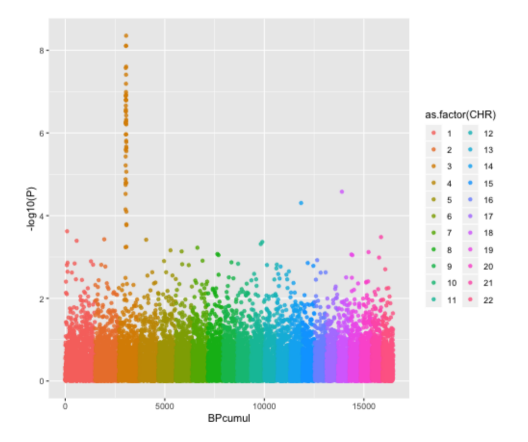
Data prep

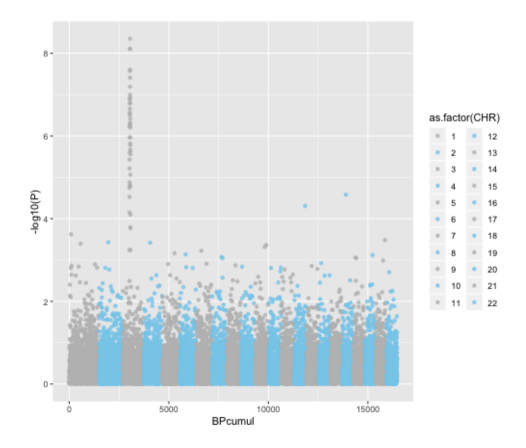
```
#>
              SNP CHR BP
                                      tot BPcumul
    16465 rs16465
                   22 530 0.5643702 15935
                                            16465
                                            16466
    16466 rs16466
                   22 531 0.1382863 15935
                                            16467
    16467 rs16467
                  22 532 0.3936999 15935
                                            16468
    16468 rs16468
                  22 533 0.1778749 15935
                                            16469
   16469 rs16469 22 534 0.2393020 15935
    16470 rs16470 22 535 0.2630441 15935
                                            16470
```

Data for plotting x-axis labels

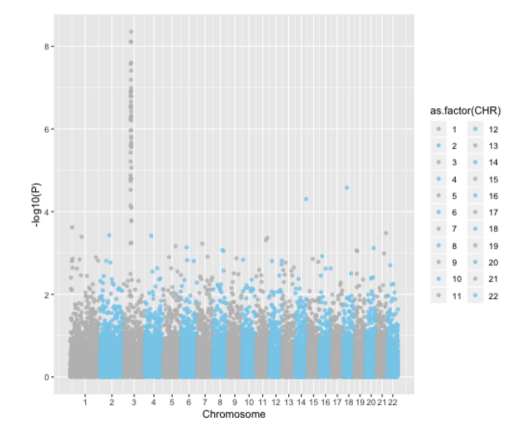
```
axisdf <- plt_dat %>%
    group_by(CHR) %>%
    summarize(center = (max(BPcumul)+min(BPcumul))/2)
axisdf
```

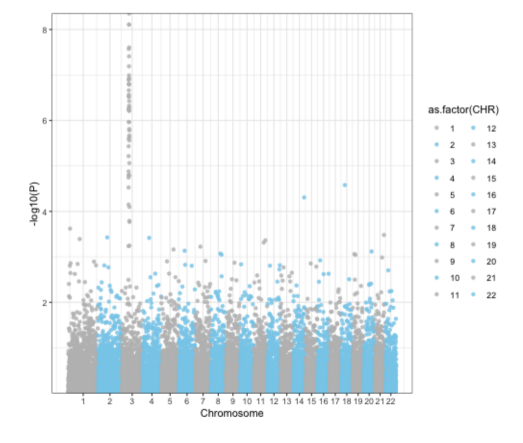


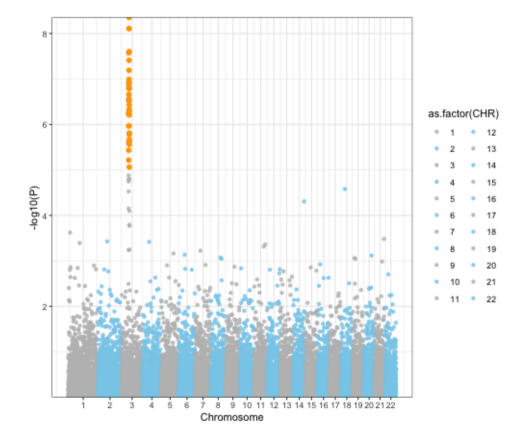




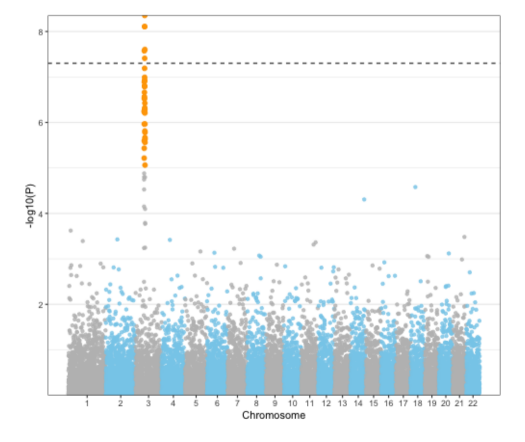
```
ggplot(plt_dat,
    aes(x = BPcumul, y = -log10(P)))+
    geom_point(aes(color = as.factor(CHR)),
        alpha = 0.8, size=1.3) +
    scale_color_manual(
        values = rep(c('grey', 'skyblue'), 22))+
    scale_x_continuous(
        name = 'Chromosome',
        breaks = axisdf$center,
        labels = axisdf$CHR
    )
```





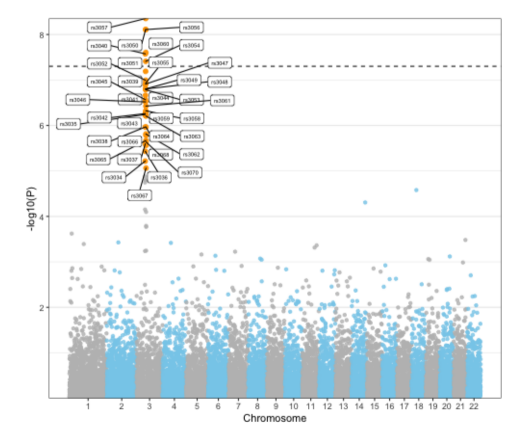


```
(plt_manhattan <- ggplot(plt_dat,</pre>
       aes(x = BPcumul, y = -log10(P))+
    geom_point(aes(color = as.factor(CHR)),
               alpha = 0.8, size=1.3) +
    scale_color_manual(
        values = rep(c('grey', 'skyblue'), 22))+
    scale_x_continuous(
        name = 'Chromosome',
        breaks = axisdf$center,
        labels = axisdf$CHR
    scale_y\_continuous(expand = c(0,0)) +
    theme_bw() +
    geom_point(data = plt_dat %>% filter(P < 1e-5),</pre>
               color = 'orange',
               size=2)+
    geom_hline(yintercept = -log10(5e-8), linetype=2)
    theme(legend.position='none',
          panel.grid.major.x=element_blank(),
          panel.grid.minor.x = element_blank())
```

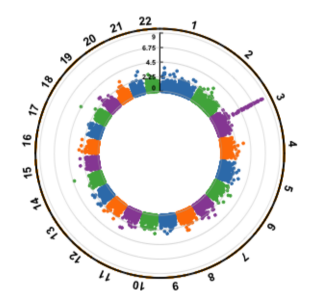


Annotation

```
library(ggrepel)
plt_manhattan +
    geom_label_repel(
        data = plt_dat %>% filter(P < 1e-5),
        aes(label = SNP),
        size = 2)</pre>
```



Circular Manhattan plots



Heatmaps

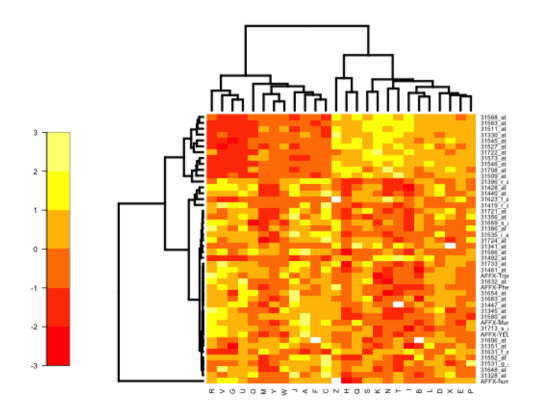
Let us count the ways

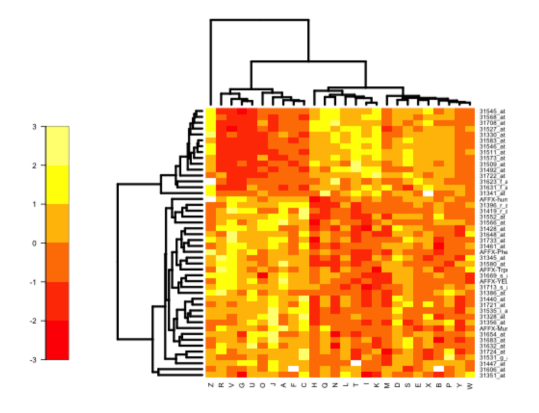
There are several ways of doing heatmaps in R:

- http://sebastianraschka.com/Articles/heatmaps_in_r.html
- https://plot.ly/r/heatmaps/
- http://moderndata.plot.ly/interactive-heat-maps-for-r/
- http://www.siliconcreek.net/r/simple-heatmap-in-r-with-ggplot2
- https://rud.is/b/2016/02/14/making-faceted-heatmaps-with-ggplot2/

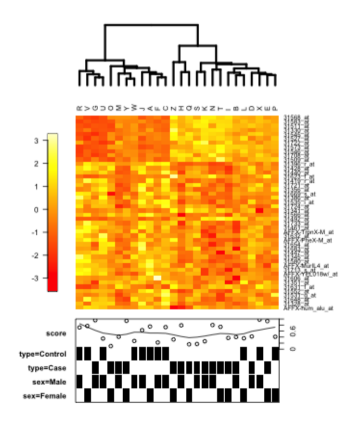
Some example data

```
library(Biobase)
#data(sample.ExpressionSet)
exdat <- readRDS('data/exprset.rds')</pre>
library(limma)
design1 <- model.matrix(~type, data=pData(exdat))</pre>
lm1 <- lmFit(exprs(exdat), design1)</pre>
lm1 <- eBayes(lm1) # compute linear model for each probeset</pre>
geneID <- rownames(topTable(lm1, coef = 2, number = 100,</pre>
                            adjust.method = 'none',
                            p.value = 0.05)
exdat2 <- exdat[geneID,] # Keep features with p-values < 0.05
head(exdat2)
#> ExpressionSet (storageMode: lockedEnvironment)
#> assayData: 1 features, 26 samples
#> element names: exprs, se.exprs
#> protocolData: none
#> phenoData
      sampleNames: A B ... Z (26 total)
#> varLabels: sex type score
#> varMetadata: labelDescription
#> featureData: none
#> experimentData: use 'experimentData(object)'
#> Annotation: hgu95av2
```

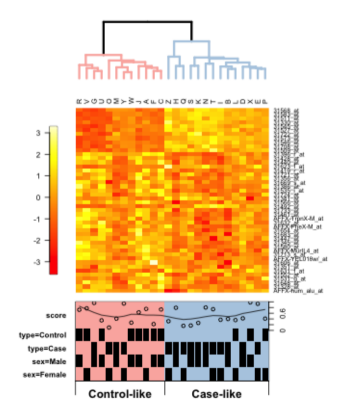




Adding annotations



Adding annotations



Playing with Seurat

Example data

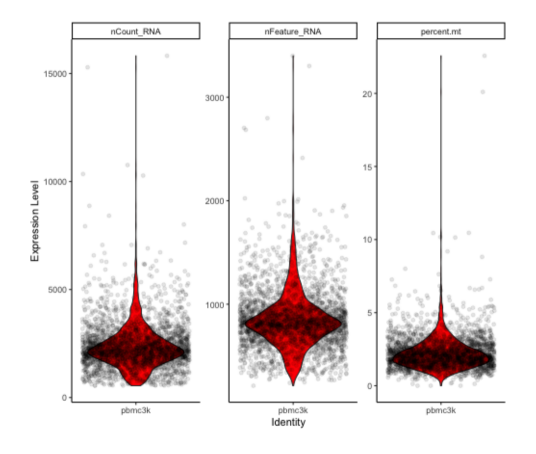
```
library(Seurat)
# pbmc.data <- Read10X(data.dir='data/hg19/')</pre>
# pbmc <- CreateSeuratObject(counts = pbmc.data, project='pbmc3k', min.cells=3, min.features=200)</pre>
pbmc <- readRDS('data/pbmc.rds')</pre>
pbmc
#> An object of class Seurat
#> 13714 features across 2700 samples within 1 assay
#> Active assay: RNA (13714 features)
names(pbmc)
#> [1] "RNA"
slotNames(pbmc)
     [1] "assays"
                                        "active.assay" "active.ident"
                         "meta.data"
    [5] "graphs"
                         "neighbors"
                                        "reductions" "project.name"
    [9] "misc"
                                        "commands"
                                                        "tools"
#>
                         "version"
```

Adding QC metrics and plotting

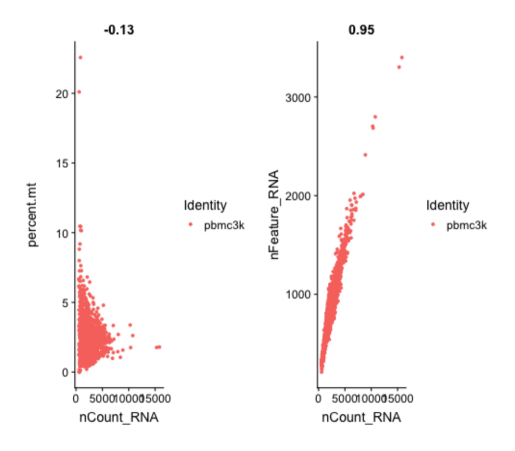
We'll calculate mitochondrial QC metrics (percentage counts originating from mitochondrial genes)

```
pbmc[['percent.mt']] <- PercentageFeatureSet(pbmc, pattern = '^MT-')</pre>
head(pbmc@meta.data)
#>
                   orig.ident nCount_RNA nFeature_RNA percent.mt
   AAACATACAACCAC
                       pbmc3k
                                    2419
                                                  779 3.0177759
   AAACATTGAGCTAC
                       pbmc3k
                                   4903
                                                 1352 3.7935958
   AAACATTGATCAGC
                       pbmc3k
                                   3147
                                                 1129 0.8897363
   AAACCGTGCTTCCG
                       pbmc3k
                                    2639
                                                      1.7430845
                       pbmc3k
   AAACCGTGTATGCG
                                    980
                                                       1.2244898
   AAACGCACTGGTAC
                       pbmc3k
                                                      1.6643551
                                    2163
```

Visualizing metrics

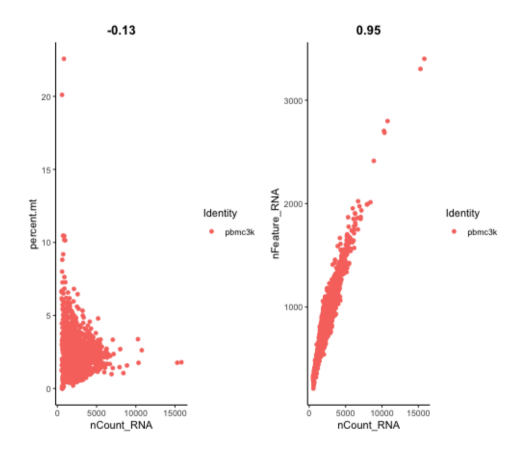


Visualizing feature-feature relationships



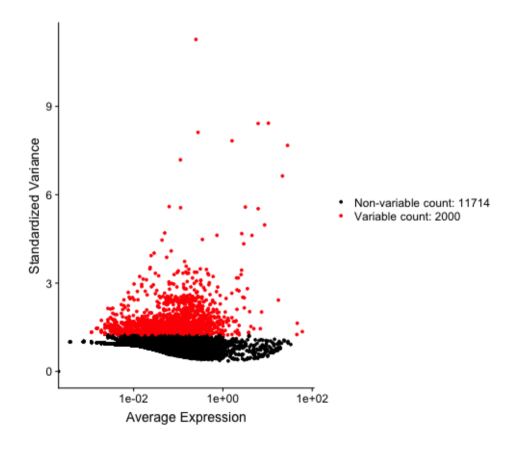
Visualizing feature-feature relationships

```
cormatrix <- cor(pbmc@meta.data %>% select(-orig.iden
plt1 <-
  ggplot(pbmc@meta.data,
         aes(x = nCount_RNA,
             v = percent.mt
             group = orig.ident.
             color = orig.ident)) +
  geom_point() +
    theme classic() +
    labs(color = 'Identity',
         title=as.character(round(cormatrix['nCount_R
  theme(plot.title = element_text(face = 'bold', hjus
plt2 <-
  ggplot(pbmc@meta.data,
         aes(x = nCount_RNA,
             y = nFeature_RNA,
             group = orig.ident,
             color = orig.ident)) +
  geom_point() +
 theme_classic() +
 labs(color = 'Identity',
       title=as.character(round(cormatrix['nCount_RNA
  theme(plot.title = element_text(face = 'bold', hjus
ggpubr::ggarrange(plt1, plt2, nrow = 1, ncol=2)
```



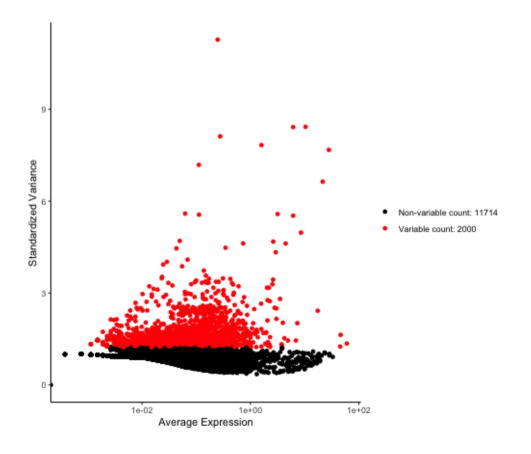
Feature selection

```
pbmc <- subset(x = pbmc,</pre>
    subset = nFeature_RNA > 200 & nFeature_RNA < 2500</pre>
pbmc <- NormalizeData(object = pbmc,</pre>
                       normalization.method = "LogNorm
                       scale.factor = 10000)
# This is stored in pbmc[['RNA']]@meta.features
pbmc <- FindVariableFeatures(object = pbmc,</pre>
                               selection.method = "vst"
                               nfeatures = 2000)
# Identify the 10 most highly variable genes
top10 <- head(x = VariableFeatures(object = pbmc), 10
# plot variable features with and without labels
plot1 <- VariableFeaturePlot(object = pbmc)</pre>
plot1
```

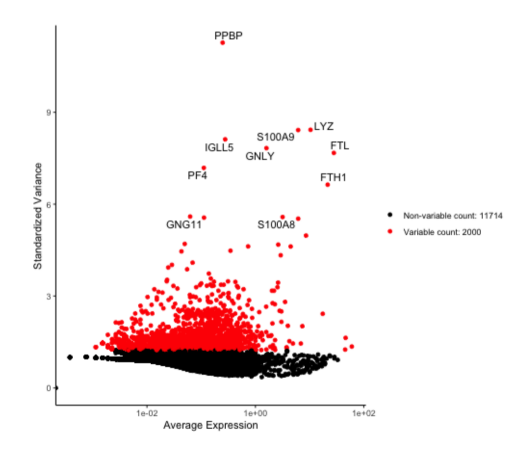


Feature selection

```
plt_data <- pbmc[['RNA']]@meta.features %>%
    rownames_to_column(var='id')
topvars <- pbmc[['RNA']]@var.features
plt_data <- plt_data %>%
    mutate(indic = ifelse(id %in% topvars,
                           'Variable count'.
                           'Non-variable count'))
bl <- plt_data %>%
    count(indic) %>%
    glue::glue_data("{indic}: {n}")
names(bl) <- c('Non-variable count', 'Variable count')</pre>
plt_data <- plt_data %>%
 mutate(indic = bl[indic])
plt11 <- ggplot(plt_data,</pre>
                aes(x = mean,
                    y = variance.standardized,
                    color = indic)) +
 geom_point() +
  scale_x_log10() +
  scale_color_manual(values = c('black', 'red')) +
 labs(x = 'Average Expression', y = 'Standardized Va
 theme_classic()
plt11
```



Feature selection



There's a lot more

We'll stop our sampling here.

- Many Bioconductor packages do use ggplot, however some use base graphics
 - Faster
- Key is to find where the data is stored, and use that to create visualizations
- Bioconductor tends to create
 - One monolithic object
 - Containing different information in slots
 - combined by lists
- slotNames and names are your friends