Exploratory data analysis of genomic data from breast cancer studies

Urminder Singh
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Introduction

Our data consists of gene expression data from various samples collected to study breast cancer (Breast Invasive Carcinoma). These samples were collected from a number of different individuals. Then, the samples' RNA were extracted sequenced to get RNA-seq reads from the sample. Using the RNA-seq, the gene expression was estimated. However, we used the data from (Wang et. al. 2018) where the expression data was mapped and normalized again to remove batch effects, which made the dataset comparable across the samples. Apart from the gene expression data for each sample, we also have associated metadata for each sample. This metadata provides information about the individual, tissue, disease etc.associated with each sample.

Our dataset has expression estimates of around 20,000 genes across 1,092 samples. Out of these 1,092 samples, 982 had tumor and 110 were normal tissues without tumor. We also added the gene metadata to our dataset which describes the gene features and functions. This also includes mutation information of the genes. This information can provide crucial information about how different mutations in the genome can lead to cancer.

Overall, our dataset is GxS matrix where each row corresponds to a gene and each column corresponds to a RNA-seq sample. Additionally, we have metadata for all the rows and all the columns.

Data

We used data from TCGA. TCGA is a comprehensive repository of human cancer molecular and clinical data, TCGA database has collected clinical and molecular phenotypes of thousands of tumor patients across different tumor types. The TCGA dataset, contains:

- Clinical information about participants
- Metadata about the samples (e.g. the weight of a sample portion, etc.)
- Histopathology slide images from sample portions
- Molecular information derived from the samples (e.g. mRNA/miRNA expression, protein expression, copy number, etc.)

First, we collected data clinical data from TCGA for breast cancer (BRCA) studies. This data contains Then we downloaded the mutation information for the BRCA studies. This data Finally, I collected the gene expression data from (Wang et. al. 2018). This data contains the estimated expression values of genes in the BRCA study. This data was normalized and batch corrected so it is comparable across different TCGA samples.

Data downloading and processing

This section describes how the data was downloaded and pre-processed.

Downloading clinical data from TCGA

We used the TCGABiolinks package to download TCGA data. The function $GDCquery_clinic$ is used to download the data. We wrote additional functions to clean the data and arrange it into a dataframe.

```
colsToKeep<-c("clinical.submitter_id","clinical.classification_of_tumor",</pre>
              "clinical.primary_diagnosis", "clinical.tumor_stage",
              "clinical.age at diagnosis", "clinical.vital status",
              "clinical.days_to_death", "clinical.tissue_or_organ_of_origin",
              "clinical.days_to_birth", "clinical.site_of_resection_or_biopsy",
              "clinical.days_to_last_follow_up", "clinical.cigarettes_per_day",
              "clinical.weight", "clinical.alcohol_history", "clinical.bmi",
              "clinical.years_smoked", "clinical.height", "clinical.gender",
              "clinical.year_of_birth", "clinical.race", "clinical.ethnicity",
              "clinical.year_of_death", "clinical.bcr_patient_barcode",
              "clinical.disease", "submitter_id", "sample_type", "tissue_type",
              "portions.submitter_id", "portions.analytes.analyte_type",
              "portions.analytes.submitter_id", "portions.analytes.analyte_type_id",
              "portions.analytes.aliquots.analyte_type", "portions.analytes.aliquots.submitter_id")
#Function takes a df and expands it by unlisting elements at a column
expand<-function(df,colName){
  res<-data.frame()
  #for each row
  for(i in 1: dim(df)[1]){
    thisRow<-df[i, ! (colnames(df) %in% c(colName))]</pre>
    tempdf<-as.data.frame(df[i, c(colName)])</pre>
    #if list is empty skip that row
    if(dim(tempdf)[1]<1){</pre>
      next
    }
    #change colnames so they are unique
    colnames(tempdf)<-paste(paste(colName,".",sep = ""),colnames(tempdf),sep = "")</pre>
    #print(paste(i,colnames(tempdf)))
    newRow<-cbind(thisRow,tempdf,row.names = NULL)</pre>
    res<-bind rows(res,newRow)</pre>
  #print(res)
 return(res)
}
getjoinedBiospcClinc<-function(projName){</pre>
  print(paste("Downloading",projName))
  clinicalBRCA <- GDCquery_clinic(project = projName, type = "clinical")</pre>
  biospecimenBRCA <- GDCquery_clinic(project = projName, type = "Biospecimen")
  #rename all cols from clinical table with suffix clinical
  colnames(clinicalBRCA)<- paste0("clinical.",colnames(clinicalBRCA))</pre>
  #expand biospecimen data in the order portions, portions.analytes, portions.analytes.aliquots
  toUnpack<-c("portions", "portions.analytes", "portions.analytes.aliquots")
  for(s in toUnpack){
    biospecimenBRCA<-expand(biospecimenBRCA,s)
```

```
#add patient barcode to biospecimen data
 biospecimenBRCA<- biospecimenBRCA %>% mutate(clinical.bcr_patient_barcode=substr(submitter_id,1,nchar
 #join clinical and biospecimen
 finalJoined <- join (clinical BRCA, biospecimen BRCA, by="clinical.bcr_patient_barcode")
 return(finalJoined)
}
projName<-"BRCA"
##Download only BRCA metadata
brcaDF<-getjoinedBiospcClinc(paste("TCGA",projName,sep = "-"))</pre>
## [1] "Downloading TCGA-BRCA"
brcaDF<-brcaDF[,colsToKeep]</pre>
#remove cols with all NA values
naCols<-colnames(brcaDF)[sapply(brcaDF, function(x)all(is.na(x)))]</pre>
brcaDF<-brcaDF[,!(colnames(brcaDF) %in% naCols)]</pre>
head(brcaDF[,1:5],3)
    clinical.submitter_id clinical.classification_of_tumor
## 1
             TCGA-3C-AAAU
                                             not reported
## 2
             TCGA-3C-AAAU
                                             not reported
## 3
             TCGA-3C-AAAU
                                             not reported
   clinical.primary_diagnosis clinical.tumor_stage
##
## 1
        Lobular carcinoma, NOS
                                           stage x
## 2
        Lobular carcinoma, NOS
                                           stage x
## 3
        Lobular carcinoma, NOS
                                            stage x
   clinical.age_at_diagnosis
## 1
                        20211
                        20211
## 2
## 3
                        20211
```

The clinical data for BRCA studies is stored in a dataframe brcaDF.

Downloading mutation data from TCGA

Next, we downloaded mutation data using TCGABiolinks. After getting the mutation data, we joined the clinical table with the mutation table to have clinical information for each mutation in the mutation table.

We now have a data frame of dimensions 93612, 145. This data frame contains clinical information and coresponding mutation information for each BRCA patient. We will use this data to explore various...

Downloading gene expression data

Next, we downloaded the gene expression data for the BRCA studies.

```
#download exp data from github for BRCA samples. reading files can take several minutes brcaexp_nontumor <- read_delim("https://raw.githubusercontent.com/mskcc/RNAseqDB/master/data/normalized
```

We now have two expression datasets for tumor and normal or non-tumor tissues from BRCA studies. These datasets contains the gene expression patterns over the tumor samples and normal samples. Each row corresponds to a gene and each column corresponds to a sample. The data dimentions for tumor expression data is 19738, 982 and for normal expression data is 19738, 110

Finally, our data consist of three data frames

- 1. brcaMAF_MD contains all the mutation information
- 2. brcaexp_nontumor contains the gene expression values over normal samples from TCGA
- 3. brcaexp_tumor contains the gene expression values over tumor samples from TCGA

Analysis

First, we looked at the mutation data to find which genes are highly mutated in BRCA and what types of different mutations are present in the data. We used the package *maftools* to do this analysis.

```
#function to take a list of maf files and summarize them
plotmafSummaryList <- function(mafList) {</pre>
11 <- mafList
#for each item in list do calculations
lnames <- names(11)</pre>
plotList <- list()</pre>
k <- 1
plist <- list()</pre>
for (i in lnames) {
print((i))
print(dim(l1[[i]]))
if (dim(l1[[i]])[1] < 1) {</pre>
next
#plot summary and save to pdf
maf <- read.maf(l1[[i]], isTCGA = T)</pre>
plotmafSummary(
maf = maf,
rmOutlier = TRUE,
addStat = 'median',
dashboard = T,
titvRaw = FALSE,
showBarcodes = F,
top = 10
)
```

```
mtext(
paste("Mutation summary by", i),
outer = T,
cex = NA,
line = -1.0,
side = 3
)
}
}
#split maf file into categories by a given variable in the clinical metadata file
splitMafby <- function(clinicalData, by, mafData) {</pre>
#get tumor samps fo different values of by
myList <- list()</pre>
uniqVals <- clinicalData %>% select(by) %>% unique
for (i in 1:nrow(uniqVals)) {
#i<-1
s <- as.character(uniqVals[i, 1])</pre>
print(s)
tList <-
clinicalData %>% filter(clinicalData[, by] == s) %>% select(Tumor_Sample_Barcode) %>%
thisData <-
mafData %>% filter(Tumor Sample Barcode %in% tList$Tumor Sample Barcode)
myList[[s]] <- thisData</pre>
}
return(myList)
}
plotGeneVarFreq <- function(mafList) {</pre>
#for each item in list
lnames <- names(mafList)</pre>
plotList <- list()</pre>
k < -1
plist <- list()</pre>
getPalette = colorRampPalette(brewer.pal(9, "Paired"))
# assign colors manually to be consistent with missing data
variants <- c(</pre>
"Missense_Mutation",
"Silent",
"3'UTR",
"Nonsense Mutation",
"5'Flank",
"Intron",
"Splice_Region",
"RNA",
"5'UTR",
"Splice_Site",
"In_Frame_Del",
"Frame_Shift_Ins",
"Frame_Shift_Del",
```

```
"In_Frame_Ins",
"3'Flank",
"Nonstop Mutation",
"Translation_Start_Site",
"IGR"
colourCount = length(unique(variants))
palette <- getPalette(colourCount)</pre>
names(palette) <- variants</pre>
for (i in lnames) {
print((i))
print(dim(l1[[i]]))
if (dim(l1[[i]])[1] < 1) {</pre>
next
}
#find top mutated genes
thisMaf <- mafList[[i]]</pre>
topGenes <-
thisMaf %>% filter(Variant_Classification != "Silent") %>% select(Hugo_Symbol) %>% group_by(Hugo_Symbol
maftest <- thisMaf %>% filter(Hugo_Symbol %in% topGenes$Hugo_Symbol & Variant_Classification != "Silent
p <-
ggplot(data = maftest, aes(x = Hugo_Symbol, fill = Variant_Classification)) +
geom_bar(stat = "count") + scale_x_discrete(limits = rev(topGenes$Hugo_Symbol)) +
coord_flip() +
theme(legend.position = "right") +
theme(
axis.text.x = element_text(size = 11, face = "bold"),
axis.text.y = element_text(size = 11, face = "bold"),
panel.grid.major = element_blank(),
panel.grid.minor = element_blank(),
panel.border = element_blank(),
panel.background = element_blank(),
axis.title = element_text(size = 12, face = "bold")
ylab("") + xlab("") + scale_fill_manual(values = palette) + ggtitle(i)
plist[[k]] <- p
k = k + 1
}
#arrange plots on a grid
do.call("grid_arrange_shared_legend", c(plist))
}
#plot on a grid with single legend
#function reference https://stackoverflow.com/questions/13649473/add-a-common-legend-for-combined-ggplo
grid_arrange_shared_legend <- function(...) {</pre>
```

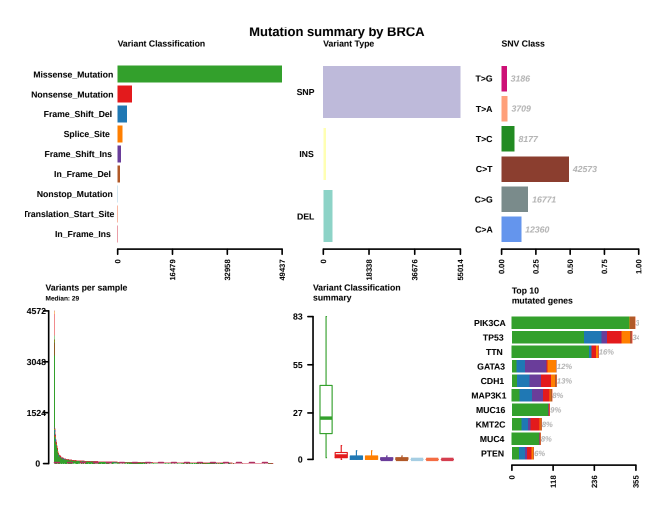


Figure 1: plotting example

```
plots <- list(...)</pre>
g <-
ggplotGrob(plots[[1]] + theme(legend.position = "bottom"))$grobs
legend <- g[[which(sapply(g, function(x)</pre>
x$name) == "guide-box")]]
lheight <- sum(legend$height)</pre>
grid.arrange(
do.call(arrangeGrob, lapply(plots, function(x)
x + theme(legend.position = "none"))),
legend,
ncol = 1,
heights = unit.c(unit(1, "npc") - lheight, lheight)
)
}
#plot summary for full dataset
tempList<-list()</pre>
tempList[["BRCA"]]<-brcaMAF</pre>
plotmafSummaryList(tempList)
```

We found that

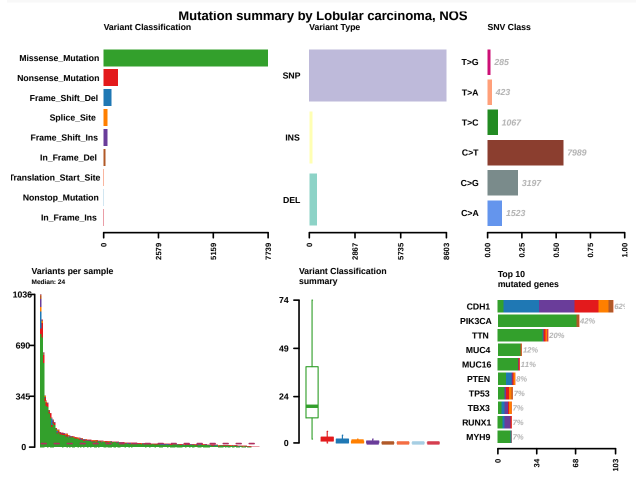
Next we looked at how the gene mutations differ for different categories e.g gender, race cancer type etc. First, we looked at which genes were mutated for samples from different race.

```
#split the maf data by clinical.race and plot summary for each category
11<-splitMafby(brcaDF,"clinical.race",brcaMAF)
plotGeneVarFreq(11)</pre>
```

We found that

Then, we looked at how genes are mutated for top two cancer types i.e. ductal carcinoma and lobular carcinoma.

#split the maf data by clinical.race and plot summary for each category
filterList<- brcaDF %>% select(clinical.primary_diagnosis) %>% count %>% arrange(desc(freq)) %>% top_n(
12<-splitMafby(brcaDF%>%filter(clinical.primary_diagnosis %in% filterList\$clinical.primary_diagnosis),"
plotmafSummaryList(12)



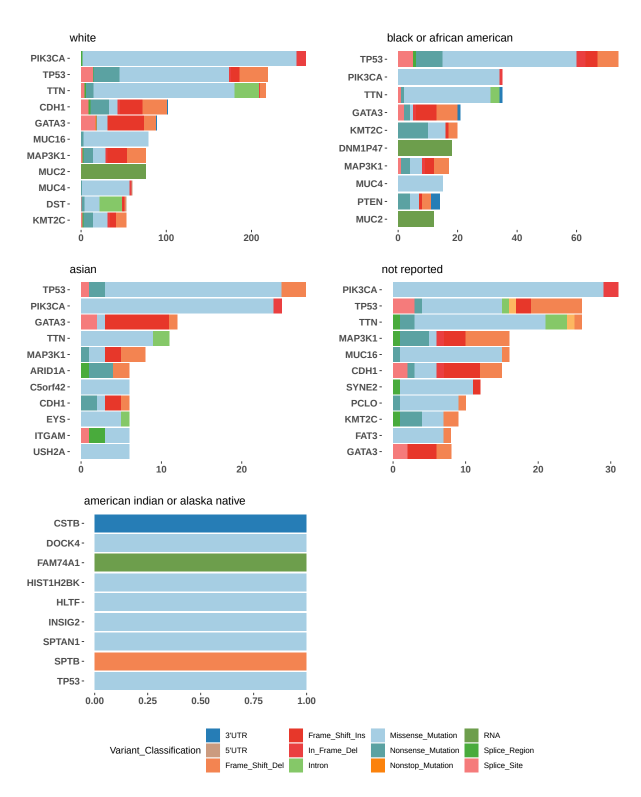
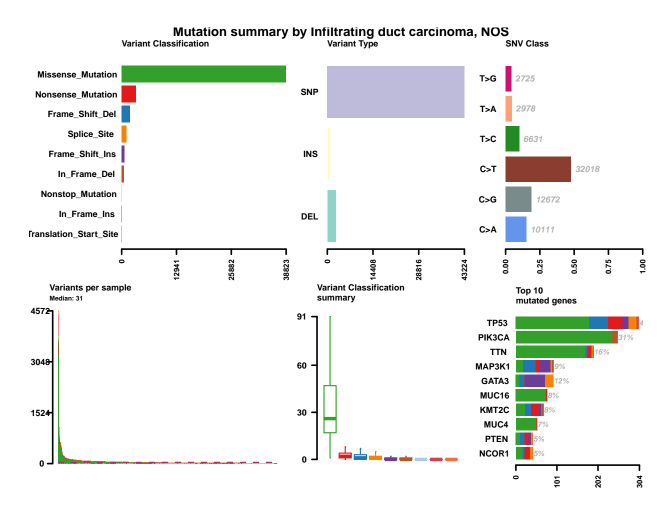
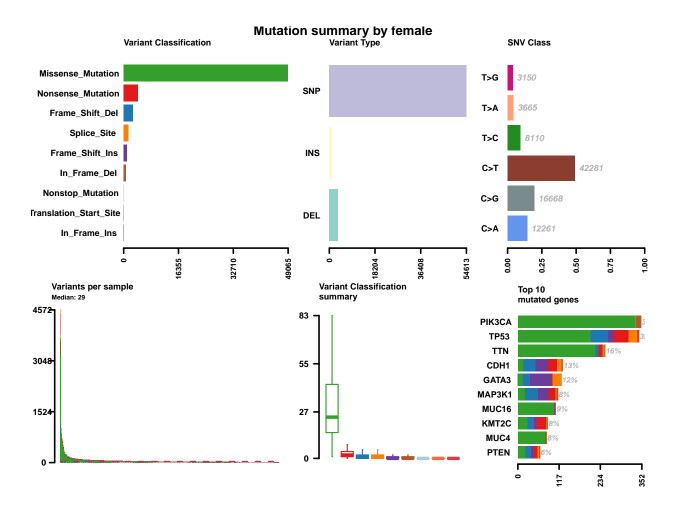


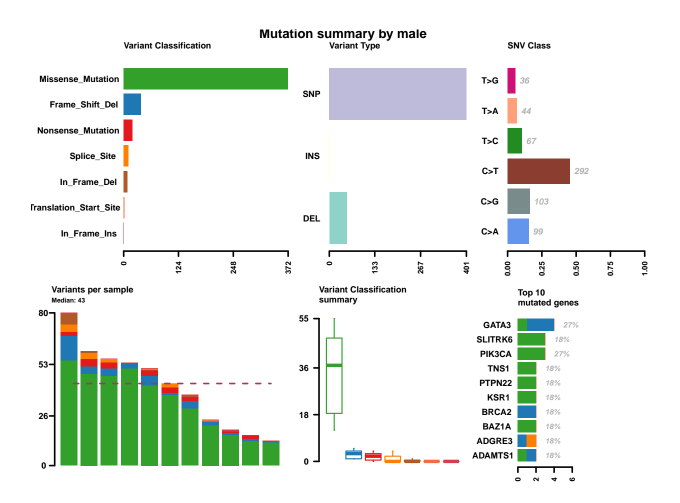
Figure 2: plotting examplerace



Finally, we looked at if there is a difference in the set of genes mutated by gender.

#split the maf data by clinical.race and plot summary for each category
13<-splitMafby(brcaDF, "clinical.gender", brcaMAF)
plotmafSummaryList(13)</pre>





From the analysis above we noticed that BRCA1 and BRCA2 are not in the set of top mutated genes. These genes are known to be associated with cancer and a mutation in these genes can increase the risk of breast and ovarian cancer in women. We then compared how many mutations were associated with BRCA1 and BRCA2.

```
brcaMAF_MD_geneGroupBRCA<-brcaMAF_MD%>%select(Hugo_Symbol)%>%mutate(geneName=ifelse(Hugo_Symbol=="BRCA1")
p1<-ggplot(data=brcaMAF_MD_geneGroupBRCA, aes(x=geneName,y=logFreq,fill=geneName))+geom_bar(stat = "ide:
    theme(axis.text.x = element_text(size = 15,face = "bold"),axis.text.y = element_text(size = 10,face =
        panel.grid.minor = element_blank(),
        panel.border = element_blank(),
        panel.background = element_blank(),
        axis.title=element_text(size=12,face="bold"))+ scale_fill_b</pre>
p2<-ggplot(data=brcaMAF_MD%>%filter(Hugo_Symbol %in% c("BRCA1","BRCA2")),aes(x=Hugo_Symbol,fill=Variant
    theme(axis.text.x = element_text(size = 15,face = "bold"),axis.text.y = element_text(size = 10,face =
        panel.grid.minor = element_blank(),
        panel.border = element_blank(),
        panel.border = element_blank(),
        panel.background = element_blank(), axis.title=element_text(size=12,face="bold"))+ scale_fill_b
grid.arrange(p1, p2, nrow = 1)
```

From figure we see that ...

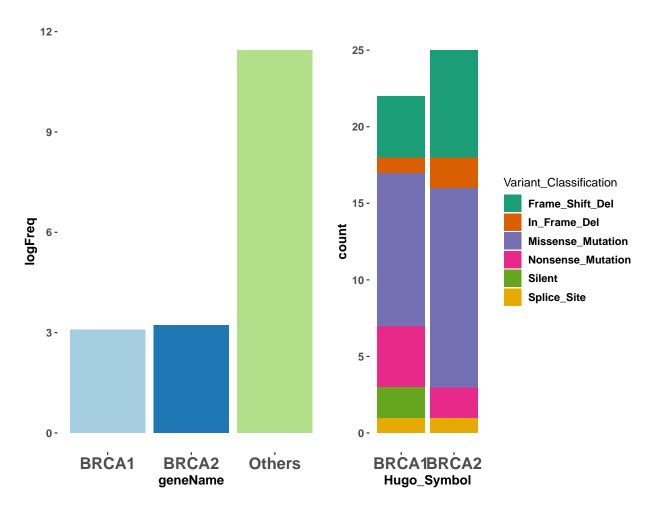
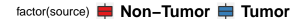


Figure 3: plotting example BRCA $\,$



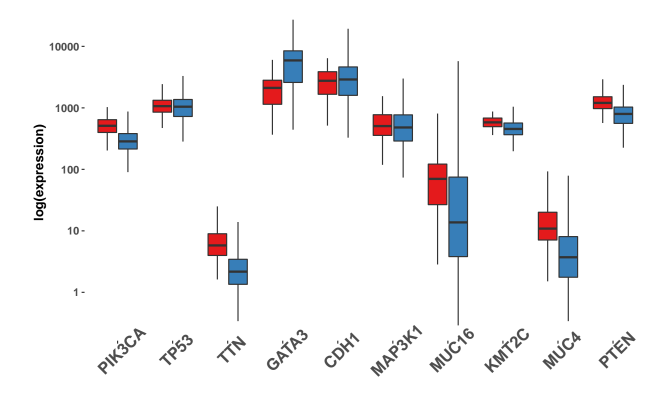


Figure 4: plotting express

After looking at various mutated genes, we looked at the expression pattern of these genes for tumor and normal samples.

```
#plot box plots of top mutated genes
topGenes<-c("PIK3CA", "TP53", "TTN", "GATA3", "CDH1", "MAP3K1", "MUC16", "KMT2C", "MUC4", "PTEN")
nt<-as.data.frame(t(brcaexp_nontumor[topGenes,]))</pre>
colnames(nt)<-topGenes</pre>
#tumordata
tdata<-as.data.frame(t(brcaexp_tumor[topGenes,]))</pre>
colnames(tdata)<-topGenes</pre>
nt<-nt%>%mutate(source="Non-Tumor")
tdata<-tdata%>%mutate(source="Tumor")
topCombined<-bind_rows(nt,tdata)</pre>
ggplot(melt(topCombined), aes(x=factor(variable),y=value,fill=factor(source)))+geom_boxplot(outlier.col
  theme(axis.text.x = element_text(angle=45, size = 15, face = "bold"), axis.text.y = element_text(size =
        panel.grid.minor = element_blank(),
        panel.border = element_blank(),
        panel.background = element_blank(), axis.title=element_text(size=12,face="bold"))+
  scale_fill_discrete(name = "Gene")+ylab("log(expression)")+xlab("")+scale_fill_brewer(palette="Set1")
```

From fig we found that the expression of

After that we compiled a list of genes are known to be differentially expressed in cancer.

```
#ref for data https://www.ncbi.nlm.nih.gov/pmc/articles/PMC5470989/
upGenes<-c("E2F1", "EZH2", "FOXM1", "MYBL2", "PLK1")
dwnGenes<-c("SCARA5", "MYOM1", "NKAPL", "PEG3", "USP2")
ntUP<-as.data.frame(t(brcaexp_nontumor[upGenes,]))</pre>
colnames(ntUP)<-upGenes</pre>
ntUP<-ntUP%>%mutate(source="Non-Tumor")
tdataUP<-as.data.frame(t(brcaexp_tumor[upGenes,]))</pre>
colnames(tdataUP)<-upGenes</pre>
tdataUP<-tdataUP%>%mutate(source="Tumor")
diffExpgenes<-bind_rows(ntUP,tdataUP)</pre>
dodge <- position_dodge(width = 0.5)</pre>
p1<-ggplot(melt(diffExpgenes), aes(x=factor(variable),y=value,fill=factor(source)))+geom_violin(position)
  theme(axis.text.x = element_text(angle=45, size = 15, face = "bold"), axis.text.y = element_text(size =
        panel.grid.minor = element_blank(),
        panel.border = element_blank(),
        panel.background = element_blank(), axis.title=element_text(size=12,face="bold"))+ scale_fill_d
#for down regulated in cancer
ntDWN<-as.data.frame(t(brcaexp_nontumor[dwnGenes,]))</pre>
colnames(ntDWN)<-dwnGenes</pre>
ntDWN<-ntDWN%>%mutate(source="Non-Tumor")
tdataDWN<-as.data.frame(t(brcaexp tumor[dwnGenes,]))</pre>
colnames(tdataDWN)<-dwnGenes</pre>
tdataDWN<-tdataDWN%>%mutate(source="Tumor")
diffExpgenesDWN<-bind_rows(ntDWN,tdataDWN)</pre>
p2<-ggplot(melt(diffExpgenesDWN), aes(x=factor(variable),y=value,fill=factor(source)))+geom_violin(posi
  theme(axis.text.x = element_text(angle=45, size = 15, face = "bold"), axis.text.y = element_text(size =
        panel.grid.minor = element_blank(),
        panel.border = element_blank(),
        panel.background = element_blank(), axis.title=element_text(size=12,face="bold"))+ scale_fill_d
grid.arrange(p1, p2, nrow = 1)
```

Comparing expression of mutated BRCA genes with other cancer types

Finally, we wanted to see expression of genes in different cancer types.

```
#download more expression data
#read tumor data for brca, liver, stomach, colon, Lung
brca <- brcaexp_tumor

coad<-read_delim("https://raw.githubusercontent.com/mskcc/RNAseqDB/master/data/normalized/coad-rsem-fpkccoadRownames<-coad$Hugo_Symbol
coad<-coad[,3:dim(coad)[2]]
rownames(coad)<-coadRownames</pre>
```

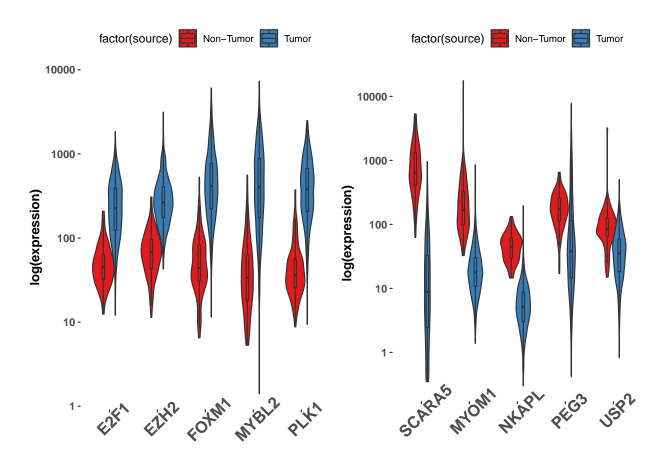


Figure 5: plotting express

```
lihc<-read_delim("https://raw.githubusercontent.com/mskcc/RNAseqDB/master/data/normalized/lihc-rsem-fpk
lihcRownames<-lihc$Hugo_Symbol
lihc<-lihc[,3:dim(lihc)[2]]</pre>
rownames(lihc)<-lihcRownames
luad<-read_delim("https://raw.githubusercontent.com/mskcc/RNAseqDB/master/data/normalized/luad-rsem-fpk</pre>
luadRownames<-luad$Hugo_Symbol</pre>
luad<-luad[,3:dim(luad)[2]]</pre>
rownames(luad) <- luadRownames
stad<-read_delim("https://raw.githubusercontent.com/mskcc/RNAseqDB/master/data/normalized/stad-rsem-fpk
stadRownames<-stad$Hugo_Symbol
stad<-stad[,3:dim(stad)[2]]</pre>
rownames(stad) <- stadRownames
topGenes<-c("PIK3CA", "TP53", "GATA3", "CDH1", "MAP3K1", "PTEN")
expVals<-c()
temp<-t(brca[topGenes,])</pre>
colnames(temp)<-topGenes</pre>
meanbrca<-colMeans(temp)</pre>
temp<-t(coad[topGenes,])</pre>
colnames(temp)<-topGenes</pre>
meancoad<-colMeans(temp)</pre>
temp<-t(lihc[topGenes,])</pre>
colnames(temp)<-topGenes</pre>
meanlihc<-colMeans(temp)</pre>
temp<-t(luad[topGenes,])</pre>
colnames(temp)<-topGenes</pre>
meanluad <- colMeans (temp)
temp<-t(stad[topGenes,])</pre>
colnames(temp)<-topGenes</pre>
meanstad<-colMeans(temp)</pre>
organs<-c("breast","colon","liver","lung","stomach")</pre>
type<-c( "other", "digestion", "digestion", "respiratory", "digestion")</pre>
colour<-c("#41ab5d","orange","orange","steelblue","orange")</pre>
plots<-list()</pre>
for(i in 1:6){
  vals<-c(meanbrca[i],meancoad[i],meanlihc[i],meanluad[i],meanstad[i])</pre>
  gganatogramData <-data.frame(organ=organs, type=type, colour=colour, value=as.numeric(vals), stringsAsFact
  p<-gganatogram(data=gganatogramData, fillOutline='#a6bddb', organism='human', sex='female', fill="val</pre>
    theme_void()+ scale_fill_gradient(low = "yellow", high = "red",name= paste(topGenes[i],"(fpkm)"))+
  plots[[i]]<-p
n <- length(plots)</pre>
```

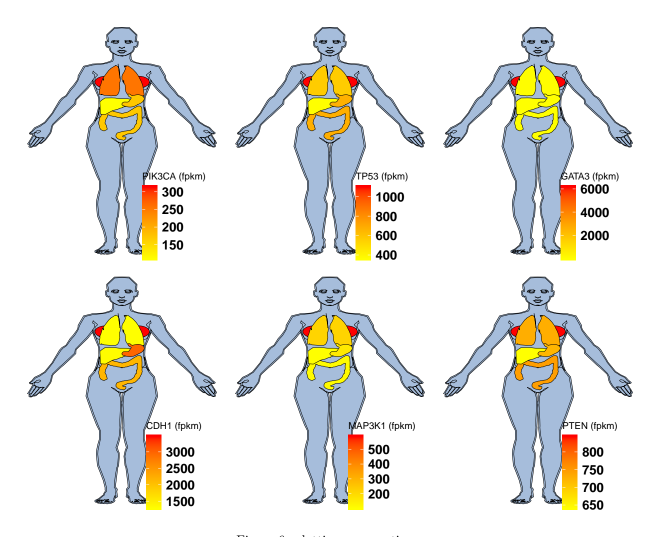


Figure 6: plotting express tissue

```
nCol <- floor(sqrt(n))
do.call("grid.arrange", c(plots, ncol=3))</pre>
```

The plots in fig shows that

Conclusion

System information

```
sessionInfo()
```

```
## R version 3.5.1 (2018-07-02)
## Platform: x86_64-w64-mingw32/x64 (64-bit)
## Running under: Windows 10 x64 (build 17134)
##
## Matrix products: default
```

```
##
## locale:
## [1] LC_COLLATE=English_United States.1252
## [2] LC_CTYPE=English_United States.1252
## [3] LC_MONETARY=English_United States.1252
## [4] LC NUMERIC=C
## [5] LC_TIME=English_United States.1252
##
## attached base packages:
## [1] grid
                 parallel stats
                                     graphics grDevices utils
                                                                    datasets
## [8] methods
                 base
## other attached packages:
## [1] bindrcpp_0.2.2
                            RColorBrewer_1.1-2
                                                gridExtra_2.3
## [4] viridis_0.5.1
                            viridisLite_0.3.0
                                                gganatogram_1.1.1
   [7] ggpolypath_0.1.0
                            ggplot2_3.1.0
                                                readr_1.2.1
                                                BiocGenerics_0.28.0
## [10] maftools_1.8.0
                            Biobase_2.42.0
## [13] plyr_1.8.4
                            data.table 1.11.8
                                                DT 0.5
## [16] dplyr_0.7.8
                            TCGAbiolinks_2.10.0
## loaded via a namespace (and not attached):
     [1] backports_1.1.2
                                     circlize_0.4.5
##
     [3] aroma.light_3.12.0
                                     NMF 0.21.0
                                     ConsensusClusterPlus 1.46.0
##
     [5] selectr 0.4-1
##
     [7] lazyeval_0.2.1
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