Presentation Project Head and Neck Cancer

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Predicting Cancer-Specific Vulnerability via DAISY System

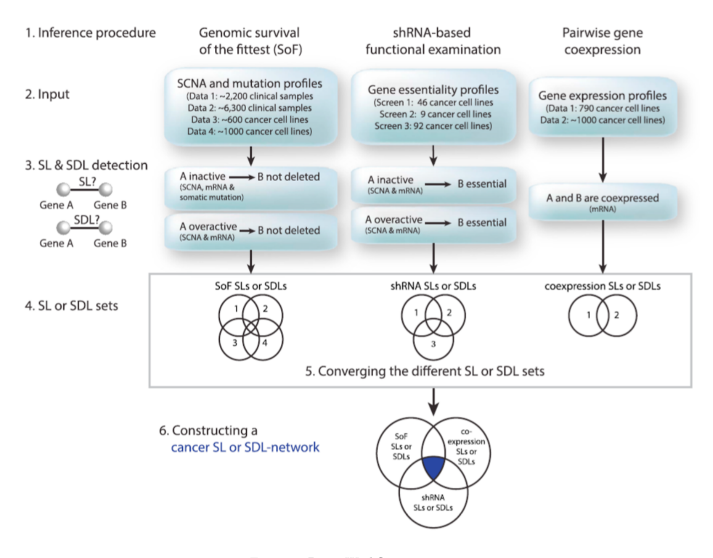


Figure 1: Daisy-Workflow

Milestones

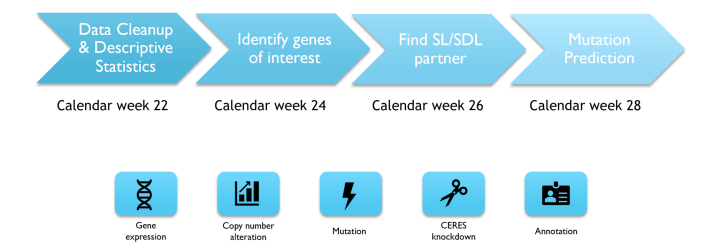


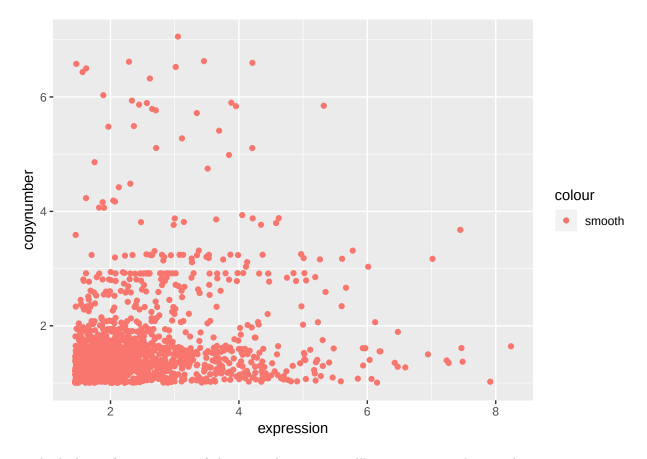
Figure 2: Milestones of our project

Identify genes of interest

k-means clustering of CNA and expression data with normalized values (-reference) for SDL GOI

To find potential genes for SDL pair detection, clustering through k-means was executed.

```
# Intersect set of expression and CNA
km.genes = Reduce(intersect, list(rownames(expression.HNC), rownames(copynumber.HNC)))
# Create data rame with expression and CNA values
km.expression = expr.HNC.norm[km.genes,]
km.copynumber = cna.HNC.norm[km.genes,]
km.data = lapply(1:length(ID), function(x) {
  out = data.frame(genes = km.genes,
                   expression = km.expression[,x],
                   copynumber = km.copynumber[,x])
  return (out)
})
# Combine rows for k mean clustering
comb.ex.cna = do.call(rbind, km.data)
comb.ex.cna = comb.ex.cna[which(comb.ex.cna$copynumber >= 1 &
                                  comb.ex.cna$expression >= quantile(comb.ex.cna$expression,0.95)),]
# Plotting of normalized expression valuess against CNA data for a short overview.
ggplot(comb.ex.cna[, 2:3],
       aes(x = expression, y = copynumber, color = "smooth")) +
  geom_point()
```



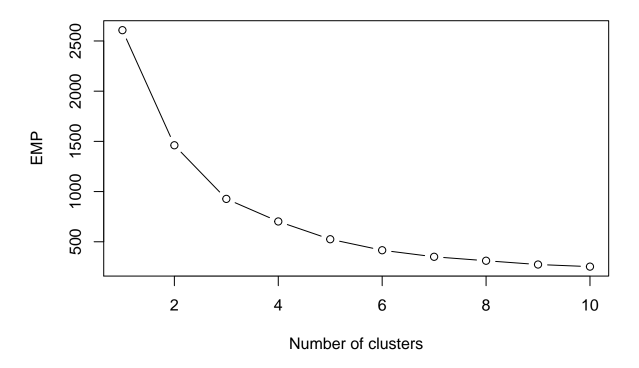
To check the perfect ammount of clusters in k-means, an ellbow test was implemented.

```
# Ellbow method to find optimal ammount of clusters
emp = vector() # Create an empty vector
# Calculate the sum within squared values for 10 clusters
for (i in 1:10) emp[i] = sum(kmeans(comb.ex.cna[2:3], i)$withinss)
emp # Look at the sum within squares for 10 clusters
```

```
[1] 2607.5505 1461.2845 927.0961 702.1999 525.1874 415.6683 350.4419 [8] 311.1543 273.4154 252.4910
```

```
# Plotting of Ellbow method
plot(1:10,
    emp,
    type = 'b',
    main = paste('The Elbow Method'),
    xlab = 'Number of clusters',
    ylab = 'EMP')
```

The Elbow Method

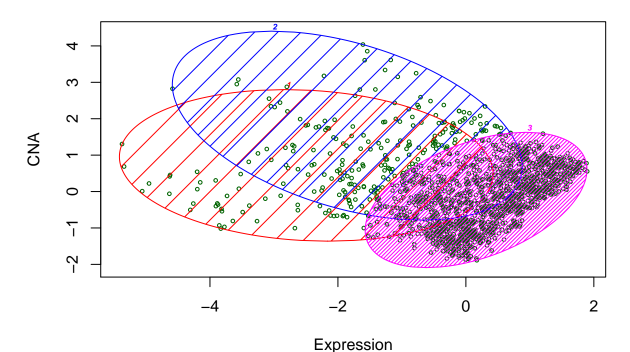


Result: The ellbow test shows that 3 clusters are sufficient.

K-means were executed and plotted. Afterwards the potential SDL candidates were identified through higher expression values.

```
# Fitting\ k-means to the dataset - elbow method (k = 3)
kmeans = kmeans(x = comb.ex.cna[,2:3], centers = 3,iter.max = 1000, nstart = 1)
y_kmeans = kmeans$cluster
# Visualising the clusters
clusplot(comb.ex.cna,
         y_kmeans,
         lines = 0,
         shade = TRUE,
         color = TRUE,
         labels = 4,
         plotchar = FALSE,
         span = TRUE,
         stand = FALSE,
         cex = 0.5,
         main = paste('Clusters of Genes'),
         xlab = 'Expression',
         ylab = 'CNA')
```

Clusters of Genes



These two components explain 72.39 % of the point variability.

The Plot "Cluster of genes" features the 3 clusters of values from expression and CNA data. Due to little variance between the data pairs, the intersection set is comparatively big.

In order to define SDL candidates, the cluster with highest expression values was chosen and filtered again according to expression numbers.

```
comb.ex.cna = cbind(comb.ex.cna, y_kmeans)# Combine kmeans with genes
# Sort comb.ex.cna data decresing for selecting adequate number of cluster
comb.ex.cna = comb.ex.cna[order(comb.ex.cna$expression,decreasing =TRUE),]

# Filtering of the most expressed and alterated genes
goi.sdl = comb.ex.cna[which(comb.ex.cna$y_kmeans == comb.ex.cna[1,4]),]

# Sort dataframe
goi.sdl = goi.sdl[order(goi.sdl$expression,decreasing=TRUE),]
kable(goi.sdl[1:5,])
```

	genes	expression	copynumber	y_kmeans
428012	DSG3	8.237872	1.641760	2
41915	FXYD3	7.914724	1.022479	2
33343	CST6	7.484452	1.372464	2
185157	F3	7.466245	1.610442	2
354894	FOXE1	7.447917	3.676721	2

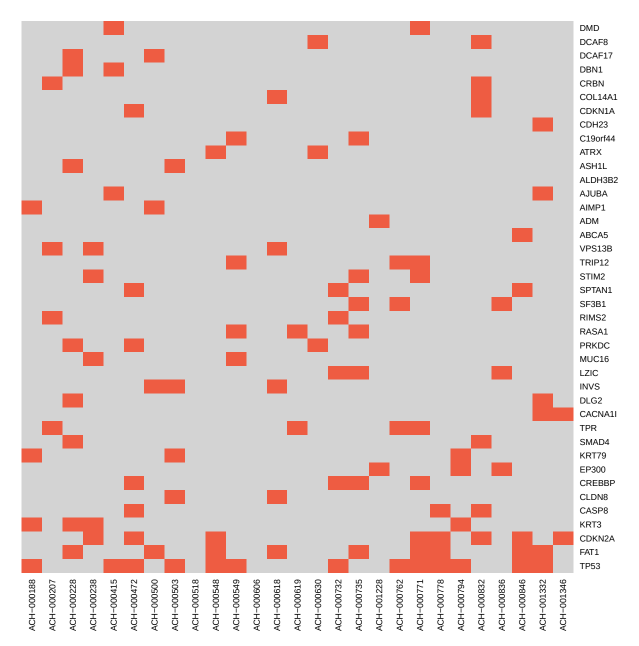
```
goi.SDL = as.character(goi.sdl$genes[1:5])
```

Investigate high frequencies of deleterious mutations SL GOI

Create a dataframe with data for the heatmap.

```
# All mutation data of HNC samples in one dataframe
mutation.HNC.df = do.call(rbind, mutation.HNC)
# Dataframe with all mutated genes in HNC samples just once
heatmap.genes = as.data.frame(table(mutation.HNC.df$Hugo_Symbol))
heatmap.data = as.data.frame(sapply(1:(length(ID)), function(a) {
  # Per HNC sample: every gene which is mutated gets a 1, otherwise a 0
  out = ifelse(heatmap.genes$Var1 %in% mutation.HNC[[a]]$Hugo_Symbol, 1, 0)
  return(out)
}))
# Define row and column names
rownames(heatmap.data) = heatmap.genes$Var1
colnames(heatmap.data) = ID
# Order from high to low frequency
heatmap.data$sum = apply(heatmap.data, 1, sum)
heatmap.data = heatmap.data[order(-heatmap.data$sum), 1:27]
# Heatmap of the most frequently deleterious mutated genes in HNC samples
heatmap(data.matrix(heatmap.data)[1:40,],
       Rowv=NA,
        Colv=NA,
        col = c("lightgrey", "tomato2"),
        scale="column",
       cexRow = 0.8,
        cexCol = 0.8,
       main = "Most frequently deleterious mutated genes in HNC samples")
```

Most frequently deleterious mutated genes in HNC samples



Result: TP53, FAT1, CDKN2A, KRT3 and CASP8 are the 5 most frequently deleterious mutated genes in our HNC samples.

According to Leemans et al. (2018) these are frequent mutations in HNC:

Cellular process	Gene	Protein	Type of gene	Mutation frequency (%)	CNA frequency (%)
Cell cycle	CDKN2A	p16 ^{INK4A}	Tumour suppressor	22	32
	TP53	p53	Tumour suppressor	72	1.4
	CCND1	G1-S-specific cyclin D1	Oncogene	0.6	25
Growth signals	EGFR	EGFR	Oncogene	4	11
Survival	PIK3CA	Catalytic p110α subunit of class 1 PI3Ks	Oncogene	18	21
	PTEN	PTEN	Tumour suppressor	3	4
WNT signalling	FAT1	Protocadherin FAT1	Tumour suppressor	23	8
	AJUBA	LIM domain-containing protein AJUBA	Tumour suppressor	7*	1
	NOTCH1	NOTCH1	Tumour suppressor	18	4
Epigenetic	KMT2D	Histone-lysine N-methyltransferase KMT2D	Tumour suppressor	16	0.4
regulation	NSD1	Histone-lysine N-methyltransferase NSD1	Tumour suppressor	12*	0.8

Data from REF. 3. Mutation data were taken from The Cancer Genome Atlas (TCGA) (n = 504) using the cBioPortal. CNA, copy number alteration; EGFR, epidermal growth factor receptor. *Putative passenger mutation that requires further functional studies.

Figure 3: Genes with frequent and highly significant somatic genetic changes in HNSCC

Compare our SL GOI with genes from literature.

```
# create dataframe with goi from literature
goi.lit = data.frame(Gene = c("CDKN2A", "TP53", "CCND1", "EGFR", "PIK3CA", "PTEN", "FAT1",
                               "AJUBA", "NOTCH1", "KMT2D", "NSD1"),
                      "Mutation.frequency" = c(22, 72, 0.6, 4, 18, 3, 23, 7, 18, 16, 12),
                      "CNA.frequency" = c(32, 1.4, 25, 11, 21, 4, 8, 1, 4, 0.4, 0.8))
{\it \# Search \ for \ the \ position \ of \ goi.lit \ in \ mutated \ genes \ of \ HNC \ samples}
heatmap_position = sapply(1:nrow(goi.lit), function(a) {
  position = which(rownames(heatmap.data) == goi.lit$Gene[a])
  # If gene is not mutated in HNC samples its position is replaced with NA
  if(length(position) == 0) {position = NA}
  return(position)
})
goi.lit$"Heatmap.position" = heatmap_position # Add heatmap position to goi.lit
rownames(goi.lit) = goi.lit$Gene
goi.lit$Gene = NULL
goi.lit = goi.lit[order(goi.lit$"Heatmap.position"),]
kable(goi.lit) # Show a table of goi.lit
```

	Mutation.frequency	CNA.frequency	Heatmap.position
TP53	72.0	1.4	1
FAT1	23.0	8.0	2
CDKN2A	22.0	32.0	3
AJUBA	7.0	1.0	28
KMT2D	16.0	0.4	66
NOTCH1	18.0	4.0	82
NSD1	12.0	0.8	715
CCND1	0.6	25.0	NA
EGFR	4.0	11.0	NA
PIK3CA	18.0	21.0	NA

	Mutation.frequency	CNA.frequency	Heatmap.position
PTEN	3.0	4.0	NA

Because our most frequent mutated genes aproximately match with frequent mutations from literature we choose the 5 genes with most frequent mutations in HNC samples.

```
goi.SL = rownames(heatmap.data)[1:5]
goi.SL
```

[1] "TP53" "FAT1" "CDKN2A" "KRT3" "CASP8"

Find SL/SDL partner

SL/SDL partner search following the daisy model

To examine gene pairs A and B which fulfill the criteria of SoF and functional examination we performe a Wilcoxon rank sum test on copynumber or CERES knockdown data. We have 5 inactive genes (goi.SL) and 5 overactive genes (goi.SDL) which are defined as genes A. The Wilcoxon rank sum test now returns all genes B which pass the test in a significant manner (p<0.05).

```
'Daisy.Wilcox' = function(input_genes,input_data, inactive_overactive, less_greater){
 Daisy Wilcox = lapply(1:length(input genes), function(m) {
 goi = input_genes[m] # Set 1 of 5 goi
 # Depending on SL or SDL partner search
 # (1) gene B is overactive (SDL)
 if(inactive_overactive == 1) { # inactive_overactive == 1 chooses overactive
   mean.exp.goi = mean(t(expression[goi,]))
   exp.goi.norm = expression[goi,] - mean.exp.goi # Normalised expression values of goi
   mut_overa.ID = lapply(1:ncol(expression), function(a) {
     # Filter all IDs which are overactive: overexpressed and CNA > 0,3
     out = if(exp.goi.norm[goi, a] > 0 & copynumber[goi,a] > 0.3) {a} else {NA}
     return(out)
   })
 # (2) gene B is inactive (SL)
 } else { # inactive_overactive == 0 chooses inactive
   mut_overa.ID = lapply(1:length(mutation), function(a) {
     # Filter all IDs which have a mutation of goi
     out = ifelse(goi %in% mutation[[a]]$Hugo_Symbol, a, NA)
     return(out)
   })
 }
 # ID of samples which have NO inactivity/overactivity of goi
 not mut overa.ID = grep("NA", mut overa.ID)
 # ID of samples which HAVE inactivity/overactivity of goi
 mut_overa.ID = as.integer(mut_overa.ID[is.na(mut_overa.ID) == FALSE])
 p.value = sapply(1:nrow(input_data), function(b) {
   mut_overa.data = t(input_data[b, mut_overa.ID])
   not_mut_overa.data = t(input_data[b, not_mut_overa.ID])
   if (less_greater == 1) {
     # One sided (greater) Wilcoxon Sum Rank Test
     p = wilcox.test(mut_overa.data, not_mut_overa.data, alternative = "greater")$p.value
   } else {
     # One sided (greater) Wilcoxon Sum Rank Test
     p = wilcox.test(mut overa.data, not mut overa.data, alternative = "less")$p.value
   }
 })
 out = data.frame(genes = rownames(input_data), p_value = p.value)
 out = out[which(out$p_value < 0.05),]</pre>
 return(out)
 })
```

```
names(Daisy_Wilcox) = input_genes
return(Daisy_Wilcox)
}
```

To examine gene pairs A and B which fulfill the criteria of gene co expression we performe a Spearman correlation test on the expression data. We have 5 inactive genes (goi.SL) and 5 overactive genes (goi.SDL) which are defined as genes A The correlation test now returns all genes B that correlate to gene A in a significant manner (p < 0.05).

```
'Daisy.Spearman' = function(input_genes){
  # Build up a list with the 5 goi; lapply: goes through the code for each gene A
  Daisy_Spearman = lapply(1:length(input_genes), function(m) {
  goi = input_genes[m]
  # Creation of a list with the length of the expression data
  # Go through every gene and calculate cor.test
  p.value = sapply(1:nrow(expression), function(a){
   expression.goi =t(expression[goi,]) # Expressiondata of the goi (gene A)
   expression.GeneB =t(expression[a,]) # Expressiondata of all other genes (gene B)
    # One sided spearman correlation test
   p = cor.test(expression.goi,expression.GeneB, alternative = "greater",
                 method = "spearman", exact = FALSE)$p.value
  })
  # Return a dataframe with the columns genes(HugoSymbols) and p_values (p.values)
  out = data.frame(genes = rownames(expression), p_value = p.value)
  out = out[which(out$p_value < 0.05),] # Filter for significant p.values
  # Only keep genes wich are not goi (would be a corr of 1)
  out = out[which(out$genes != goi),]
  return(out)
  })
  names(Daisy_Spearman) = input_genes
  return(Daisy_Spearman)
}
```

Both functions return a list, which contains the five genes of interest, each containing a dataframe with all genes examined as potential SL/SDL partners plus their calculated p-values.

SL-parter

SoF

```
genes p_value
10 NAALAD2 0.01270171
88 NAALADL1 0.01787355
```

```
105 MCTS2P 0.01801226
109 SNORD119 0.03415298
120 SNORD111B 0.04370297
148 MIR875 0.02447163
```

Functional examination

```
genes p_value
10 AADAC 0.008677123
21 AAR2 0.016419751
70 ABCG1 0.010374932
100 ABL2 0.042951952
149 ACN9 0.006751253
211 ACTR1B 0.010514096
```

Gene coexpression

```
SL.coexpression = Daisy.Spearman(goi.SL)
# Example of SL coexpression for first GOI (TP53)
head(SL.coexpression[[1]])
```

```
genes p_value
4 SCYL3 7.196232e-06
5 C1orf112 2.516883e-04
10 NFYA 2.641665e-09
13 LAS1L 3.844908e-02
17 ANKIB1 1.345544e-02
24 CD99 8.535496e-03
```

SDL-parter

SoF

Functional examination

```
SDL.coexpression = Daisy.Spearman(goi.SDL)
```

Converging the different SL and SDL sets

Define function to converge SL/SDL sets.

```
'partner' = function(input_genes, SoF, functional.examination, coexpression){
 partner = lapply(1:length(input_genes), function(a) {
   dat_picker.SoF = SoF[[a]]
   dat_picker.func.exam = functional.examination[[a]]
   dat_picker.coexpression = coexpression[[a]]
   # Intersect of genes from all three tests
   partnergenes = Reduce(intersect, list(dat_picker.SoF$genes,
                                          dat_picker.func.exam$genes,
                                          dat_picker.coexpression$genes))
   out = data.frame(
     genes = partnergenes,
     # Column with reduced genes
     SoF = dat_picker.SoF[which(dat_picker.SoF$genes %in% partnergenes), "p_value"],
     functional_examination = dat_picker.func.exam[which(dat_picker.func.exam$genes
                                                          %in% partnergenes), "p_value"],
     coexpression = dat_picker.coexpression[which(dat_picker.coexpression$genes
                                                   %in% partnergenes), "p_value"]
   ) # Column with p value from coexpression
   return(out)
 names(partner) = input_genes
 return(partner)
```

Converge different sets.

SL.genes	SL.number.of.partner	SDL.genes	SDL.number.of.partner
TP53	21	DSG3	96
FAT1	54	FXYD3	78
CDKN2A	60	CST6	232
KRT3	124	F3	78
CASP8	22	FOXE1	136

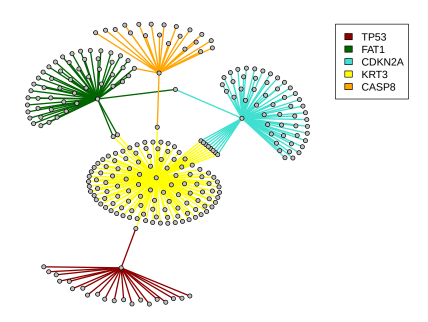
Plotting networks of the Interactions

Creation of a function for transforming given data into a format iGraph can work with and plotting of the network.

```
'network' = function(X.partner,width,node.color,frame.color,node.size,Titel,label,GOI){
 # Input: SL/SDL.partner, number, string, string, number, string, NA or names(plot)
 # Define colours
 color.vector = c("darkred", "darkgreen", "turquoise", "yellow", "orange")
 # Transform data into a dataframe with 3 colums for each GOI
 edgelist = lapply(1:length(X.partner), function(a){
   dat_picker = X.partner[[a]]
   edge.list = data.frame(Gene.A = names(X.partner)[a],
                          Gene.B = dat_picker$genes,
                           colours = color.vector[a])
 })
 edge.list = do.call(rbind, edgelist) # Combine all goi-dataframes to one.
 # Change column colours from factor to chracter for plotting function
 edge.list$colours = as.character(edge.list$colours)
 # Tranform the dataframe into an iGraph object
 net = graph_from_data_frame(edge.list, directed = T)
 layout1 = layout_with_kk(net) # Given layout
 # Plotting parameters are defined. vertex= Genes edges = interactions
 plot(net, layout = layout1, rescale = T,
      vlim = c(-0.9,1),
      xlim = c(-0.1, -0.1),
      edge.color = edge.list$colours,
      edge.width = width,
      edge.arrow.mode = 0,
      vertex.color = node.color,
      vertex.frame.color = frame.color,
      vertex.size = node.size,
      vertex.label.font = 2,
      vertex.label.color = "black",
      vertex.label = label,
      margin = 0.1)
 # Add legend to the graph
 legend (x = 1.2,
         y = 1, GOI,
         fill = c("darkred", "darkgreen", "turquoise", "yellow", "orange"))
 # Add titel to the graph
 title(Titel,cex.main=3,col.main="black ")
```

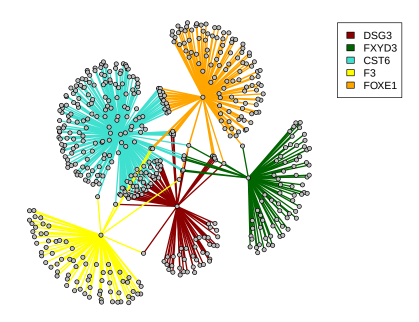
network(SL.partner,2,"grey","black",3,"Network of SL interaction", NA, goi.SL)

Network of SL interaction



```
network(SDL.partner,2,"grey","black",3,"Network of SDL interaction", NA, goi.SDL )
```

Network of SDL interaction



Because there are to many SL/SDL partner to plot a network we reduce SL/SDL partner to the most significants.

To do so, we reduce the number of partner genes for better plotting without removing genes which are connected to more than one GOI.

```
'reduce.partner' = function(list){
    # Go through every element of SL/SDL partner
    reduced_partner = lapply(1:length(list), function(a){
        dat_picker = list[[a]]

# Create an extra column with the mean p value of all 3 tests
        dat_picker$order = apply(dat_picker[, 2:4], 1, mean)
    # Order the partner genes after this mean value from most to least significant p value
        ordered_data = dat_picker[order(dat_picker$order), ][, 1:4]

gene_intersect = lapply(1:length(list), function(b) {
        # Search for genes that are partner for more than one GOI
        as.list(Reduce(intersect, list(dat_picker$genes), list[[b]]$genes))
    })
    gene_intersect[a] = NULL # Remove intersect of the GOI partner genes with itself
    keep_genes = c(gene_intersect[[1]],
```

```
gene_intersect[[2]],
                   gene_intersect[[3]],
                   gene_intersect[[4]]) # Combine all the genes in a list
   position = sapply(1:length(keep_genes), function(c) {
      which(as.character(ordered_data$genes) == keep_genes[c])
    # Partner genes that are already in the first 20 genes don't need to be chosen
   position = position[which(position > 20)]
   if (length(position) > 3) {
      # If there are still more than 3 extra genes only choose the first 3
     position = position[1:3]
   out = ordered_data[c(1:20, position), ]
   return(out) # Return reduced data
  })
  names(reduced_partner) = names(list)
  return(reduced_partner)
}
```

Reduce SL.parter/SDL.partner to the most significant

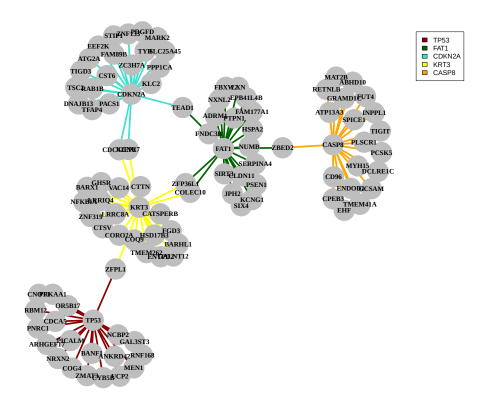
SL.genes	SL.partner	SDL.genes	SDL.partner
TP53	20	DSG3	23
FAT1	21	FXYD3	23
CDKN2A	22	CST6	23
KRT3	23	F3	23
CASP8	20	FOXE1	23

Now the number of SL/SDL partner is much better to plot.

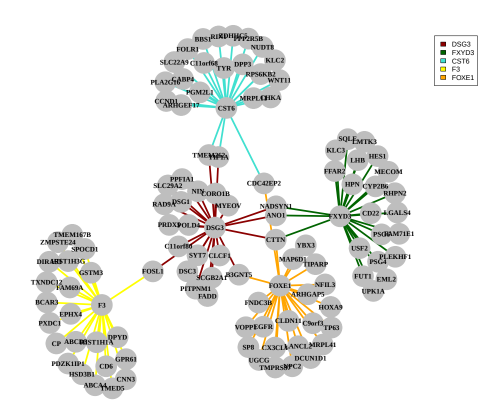
Plotting SL/SDL partner network with reduced number of partner

SL interaction plot with reduced partners

Network of SL interaction (reduced)



Network of SDL interaction (reduced)



Mutation prediction via logistic regression

Prepare data for regression model

```
# Reduce all genes to those of which we have expression, copynumber and knockdown data.
regression.genes = Reduce(intersect,
                          list(rownames(expression.HNC),
                               rownames(copynumber.HNC),
                               rownames(kd.ceres.HNC)
                          ))
# Create a dataframe with expression, CNA, knockdown and mutation data from a HNC sample.
regression.data = lapply(1:length(ID), function(a){
  regression.mutation = sapply(1:length(regression.genes), function(b){
    ifelse(regression.genes[b] %in% mutation.HNC[[a]]$Hugo_Symbol, TRUE, FALSE)
  })
  regression.data = data.frame("genes" = paste(regression.genes, ID[a], sep = "_"),
                               "expression" = expression.HNC[regression.genes, a],
                               "copynumber" = copynumber.HNC[regression.genes, a],
                               "kockdown" = kd.ceres.HNC[regression.genes, a],
                               "mutation" = regression.mutation)
})
# Bind these 27 dataframes to one dataframe
regression.data = do.call(rbind, regression.data)
dim(regression.data)
[1] 458190
                5
head(regression.data)
```

```
genes expression copynumber
                                             kockdown mutation
                                                         FALSE
1
     DPM1_ACH-000188 5.3593103
                                    0.1574 -0.42737583
    SCYL3_ACH-000188 1.2570106
                                    0.1441 -0.01577348
                                                         FALSE
3 Clorf112_ACH-000188 3.1093606
                                    0.1441 -0.03149540
                                                         FALSE
4
      FGR_ACH-000188 0.1634987
                                   0.2019 -0.10972076
                                                         FALSE
                                                         FALSE
5
      CFH_ACH-000188 4.0285692
                                   -0.0637 0.03189585
                                   0.1872 0.02271554
    FUCA2_ACH-000188 4.6536333
                                                         FALSE
```

Reformat the data because strings can not be an input for a machine learning model

```
regression.data$mutation = factor(regression.data$mutation, levels = c("FALSE", "TRUE"))
# Reformat the rownames
rownames(regression.data) = regression.data$genes
# Get rid of the gene names becaue they are now the rownames
regression.data = regression.data[,2:ncol(regression.data)]
head(regression.data)
```

```
expression copynumber
                                            kockdown mutation
DPM1 ACH-000188
                    5.3593103
                                  0.1574 -0.42737583
                                                       FALSE
SCYL3_ACH-000188
                    1.2570106
                                  0.1441 -0.01577348
                                                       FALSE
C1orf112_ACH-000188 3.1093606
                                  0.1441 -0.03149540
                                                       FALSE
FGR_ACH-000188
                    0.1634987
                                  0.2019 -0.10972076
                                                       FALSE
CFH_ACH-000188
                                                       FALSE
                    4.0285692
                                 -0.0637 0.03189585
FUCA2_ACH-000188
                    4.6536333
                                  0.1872 0.02271554
                                                       FALSE
```

Split the data into train and test-data

```
# 75% of data is used to train, 25% to test the regression model
inTrain = createDataPartition(y = regression.data$mutation, p = .75, list = FALSE)
# Only get training data
train.set = regression.data[inTrain,]
head(train.set)
```

```
expression copynumber
                                            kockdown mutation
DPM1_ACH-000188
                    5.3593103
                                  0.1574 -0.42737583
                                                        FALSE
SCYL3_ACH-000188
                    1.2570106
                                                        FALSE
                                  0.1441 -0.01577348
Clorf112_ACH-000188 3.1093606
                                  0.1441 -0.03149540
                                                        FALSE
FGR_ACH-000188
                    0.1634987
                                  0.2019 -0.10972076
                                                        FALSE
                                 -0.0637 0.03189585
CFH_ACH-000188
                    4.0285692
                                                        FALSE
FUCA2_ACH-000188
                    4.6536333
                                  0.1872 0.02271554
                                                        FALSE
```

```
# Only get testing data
test.set = regression.data[-inTrain,]
head(test.set)
```

```
expression copynumber
                                          kockdown mutation
GCLC ACH-000188
                 4.14974712
                                0.1549 0.06840387
                                                     FALSE
                                0.2083 0.02314953
STPG1_ACH-000188
                 3.09254574
                                                      FALSE
ENPP4_ACH-000188 0.01435529
                               0.1549 0.09424808
                                                     FALSE
BAD_ACH-000188
                 6.37868501
                               1.1598 -0.08587470
                                                      FALSE
LAP3_ACH-000188
                 5.09634620
                               -0.7520 -0.18606702
                                                      FALSE
HS3ST1_ACH-000188 2.21723072
                               -0.7520 0.01858038
                                                      FALSE
```

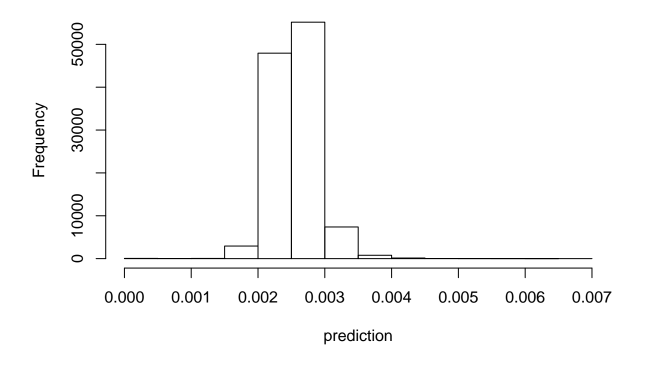
Train the model

```
regression.model = glm(mutation ~ ., data = train.set, family = "binomial")
summary(regression.model)
Call:
glm(formula = mutation ~ ., family = "binomial", data = train.set)
Deviance Residuals:
   Min
             1Q
                Median
                              3Q
                                     Max
-0.1289 -0.0739 -0.0712 -0.0689
                                   3.5945
Coefficients:
           Estimate Std. Error z value Pr(>|z|)
(Intercept) -6.01402
                      0.05078 -118.429 < 2e-16 ***
                      0.01486 1.732 0.08326 .
expression 0.02574
copynumber 0.22315
                      0.08116
                                 2.750 0.00597 **
kockdown
           0.22519
                      0.12317 1.828 0.06752 .
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 12249 on 343642 degrees of freedom
Residual deviance: 12235 on 343639 degrees of freedom
AIC: 12243
Number of Fisher Scoring iterations: 9
```

Evaluate the precision and the recall of the model

```
prediction = predict(regression.model, newdata = test.set, type="response")
hist(prediction) # Look at the predictions
```

Histogram of prediction



Devide prediction in threshholds

```
threshhold_ls = seq(min(prediction), max(prediction), by = 0.001) # Make a threshold list
length(threshhold_ls)
```

[1] 7

```
threshold_list = lapply(1:length(threshhold_ls), function(a){
    # Check for every threshhold:
    table(prediction > threshhold_ls[a], # Returns TRUE or FALSE
        test.set$mutation) # Comparing if return fits actual bimodal value
    # Counting True Negative, False Negative, False Positive, True Positive
})

threshold_list[[1]]
```

```
FALSE TRUE
FALSE 1 0
TRUE 114253 293
```

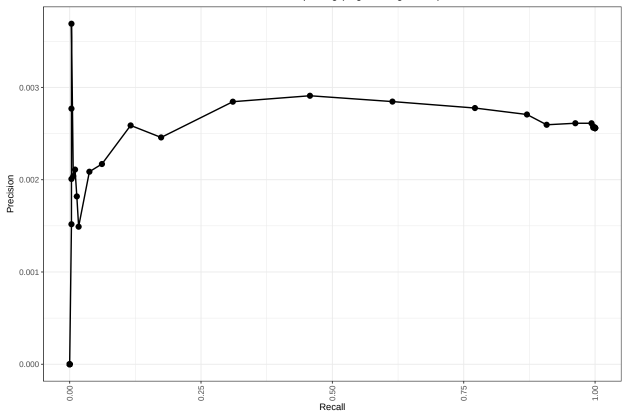
Calculate the recall and precision values

```
# Make a dataframe out of the list of lists
results_df = lapply(1:length(threshold_list), function(a) {
  threshold = threshold_list[[a]]
  threshold_value = threshhold_ls[[a]]
  df_precision_recall = as.data.frame(list(threshold["FALSE", "FALSE"],
                                      threshold["FALSE", "TRUE"],
                                      threshold["TRUE", "FALSE"],
                                      threshold["TRUE", "TRUE"]))
  df_precision_recall$threshold = threshold_value
  colnames(df_precision_recall) = c("TN", "FN", "FP", "TP", "Thresh")
  return(df_precision_recall)
})
# Bind all the lists together
results_df = do.call(rbind, results_df)
# Calculate the recall; recall = True positive / all positive events
recall = apply(results_df, 1, function(x) { x[["TP"]] / (x[["TP"]]+x[["FN"]]) })
# Calculate the precision; precision = True positive / all positive detected events
precision = apply(results_df, 1, function(x){x[["TP"]] / (x[["TP"]]+x[["FP"]])})
results_df$recall = recall # Bind recall to the data
results_df$precision = precision # Bind precision to the data
head(results_df) # Look at the data
```

```
TN FN FP TP Thresh recall precision
1 1 0 114253 293 1.635389e-06 1 0.002557924
2 41 0 114213 293 1.016354e-04 1 0.002558818
3 43 0 114211 293 2.016354e-04 1 0.002558863
4 46 0 114208 293 3.016354e-04 1 0.002558930
5 50 0 114204 293 4.016354e-04 1 0.002559019
6 54 0 114200 293 5.016354e-04 1 0.002559108
```

Plot the data

Precision/Recall plotting (Logistic Regression)



Looking on the precision it can be said that our logistic regression model for prediction of deleterious mutation in head and neck cancer scores very poorly. To be just in about 0,3% of all cases precise with the prediction is an unusuable model and has plenty of room for improvement. Especially regarding to declining precision towards a recall of 1.00, which is a true mutation and a result on which basis further decisions would be drawn.

References

Articles

- 1.) Jerby-Arnon, L., et al. (2014). "Predicting Cancer-Specific Vulnerability via Data-Driven Detection of Synthetic Lethality." Cell 158(5): 1199-1209.
- 2.) Ashworth, A., et al. (2011). "Genetic Interactions in Cancer Progression and Treatment." Cell 145(1): 30-38.
- 3.) O'Neil, N. J., et al. (2017). "Synthetic lethality and cancer." Nature Reviews Genetics 18: 613.

Figures

Figure 1 - Jerby-Arnon, L., et al. (2014). "Predicting Cancer-Specific Vulnerability via Data-Driven Detection of Synthetic Lethality." Cell 158(5): 1199-1209.

Figure 3 - Leemans, C. R. et al. (2018). The molecular landscape of head and neck cancer. Nat. Rev. Cancer 18, 269-282.