### Direct inhibition of the NOTCH TF. Differenctial expression analysis

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# 1 Array expression profiling: direct inhibition of the NOTCH transcription complex

#### Goals

We will try to reproduce some of the differential expression results obtained by the paper *Direct inhibition of the NOTCH transcription complex*. In this paper, Moellering et al. try to design and synthesize a peptide able to inhibit NOTCH transcription factor for the treatment of individuals affected by Acute lymphoblastic leukemia (T-ALL).

NOTCH proteins regulate signaling pathways involved in cellular differentiation, proliferation and apoptosis. Overactive Notch signaling as been observed in numerous cancers and has been extensively studied in the context of T-ALL where more than 50% of pateints have mutant NOTCH1. Small molecule modulators of these proteins would be important for understanding the role of NOTCH proteins in malignant and normal biological processes.

The authors measure the global changes in gene expression upon treatment of the human T-ALL cell lines HPB-ALL and KOPT-K1 with either vehicle alone dimethylsulphoxide (DMSO) control or the designed peptite SAHM1, an alpha-helical hydrocarbon peptide derived from the MAML1 co-activator protein.

### Overall design

They triplicate cultures of KOPT-K1 or HPB-ALL cells that were treated with either DMSO or SAHM1 (20 uM) for 24 hours. Total RNA was extracted and hybridized to Affymetrix human U133 plus 2.0 microarrays (three arrays per treatment per cell line for a total of 12 arrays).

### 1.1 Pipeline

### 1.2 Python imports

```
Script 1.2.1 (python)

import rpy2.rinterface
wreload_ext rpy2.ipython
```

### 1.3 R imports

```
Script 1.3.1 (R)

1 %%R
2 ##1. Load libraries
3 library("affy")
4 library("limma")
5 library("genefilter")
6 library(simpleaffy)
7 library(hgu133plus2.db)
8 wd <- "/Users/nandoide/misc_work/Desktop/uni/TRREP"
9 setwd(wd)
```

### 1.4 Functions

```
Script 1.4.1 (R)
1 %%R
2
   import_CEL <- function(pattern) {</pre>
       # Import CEL files into affiBatch object
       files <- list.files(pattern = pattern)</pre>
       names <- gsub(".CEL.gz", "", files)</pre>
6
       abatch <- ReadAffy(filenames = files, compress = TRUE, sampleNames = names)</pre>
       return(abatch)
8
  }
9
10
  create_eset <- function(affyBatch) {</pre>
       # Generates object eset (class ExprSet),
12
       # expresso function provides intensities in log scale
13
       return(expresso(affyBatch,
14
                 bg.correct = TRUE,
15
                 bgcorrect.method="rma",
16
                 normalize = TRUE,
17
                 normalize.method="quantiles",
18
                 pmcorrect.method="pmonly",
19
                 summary.method="medianpolish",
20
                 verbose = TRUE))
21
22 }
23
  boxplots <- function(affyBatch, eset, title) {</pre>
24
       # Generate BOXPLOTS before and after normalization
25
       boxplot(affyBatch,
26
           main=paste0("Boxplot Before Normalization ", title),
27
           col = "lightgrey")
28
       df_eset <- as.data.frame(exprs(eset))</pre>
29
30
       boxplot(data.frame(df_eset),
31
           main=pasteO("Boxplot After Normalization (log scale) ", title), col = "white")
32
  }
33
34
   create_TopTable <- function(eset, control_samples=c(1,1,1,0,0,0)) {</pre>
35
       # Generate Toptable with limma
36
37
       # Data filtering using IQR.
38
       esetIQR <- varFilter(eset, var.func=IQR, var.cutoff=0.5, filterByQuantile=TRUE)</pre>
39
40
       # Differential expression analysis.######
41
       r_control_samples <- 1 - control_samples</pre>
42
       design <- cbind(DMSO=control_samples, SAHM1=r_control_samples)</pre>
43
44
       rownames(design) <- colnames(eset)</pre>
45
46
       #7. Contrasts matrix.
47
       cont.matrix <- makeContrasts(DMSO_SAHM1 = SAHM1 - DMSO, levels = design)</pre>
48
49
```

```
#8. Obtaining differentially expressed genes (DEGs)
50
       #Linear model and eBayes
51
       fit <- lmFit(esetIQR, design) ##qetting DEGs from IQR</pre>
52
       fit2 <- contrasts.fit(fit, cont.matrix)</pre>
53
       fit2 <- eBayes(fit2)</pre>
54
55
       #Table with DEGs results
56
       toptableIQR <- topTable(fit2, number=dim(exprs(esetIQR))[1], adjust.method="BH",</pre>
57

    sort.by="p")

       return(toptableIQR)
58
  }
59
60
   anotate_TopTable <- function(toptable) {</pre>
61
       # Obtain gene names from probe names and chip symbol dataset
62
       probenames_toptable <- as.character(rownames(toptable ))</pre>
63
       genesymbols_toptable <- as.character(mget(probenames_toptable, hgu133plus2SYMBOL))</pre>
64
       # Annotated gene table
65
       toptable_anot <- cbind(Symbol = genesymbols_toptable, toptable)</pre>
66
       return(toptable_anot)
67
  }
68
69
   generank_table <- function(toptable, rnk.file) {</pre>
70
       # Generate rank of table top 50 upregulared and top 50 downregulated from 250 better
       # adjustes p-values
72
       more_significant = toptable[order(toptable$adj.P.Val, decreasing = FALSE),][1:250,]
73
74
       up_50 = more_significant[which(toptable$logFC > 0), ] [1:50,] # up reg top 50
       down_50 = more_significant[which(toptable$logFC < 0), ] [1:50,] # down req top 50</pre>
75
76
77
       print("Down-regulated genes")
       print(down_50[order(down_50$logFC), c(1,2,6)])
78
79
       print("Up-regulated genes")
80
       print(up_50[order(up_50$logFC), c(1,2,6)])
81
82
       d \leftarrow rbind(down_50[order(down_50\$logFC), c(1,2,6)], up_50[order(up_50\$logFC), c(1,2,6)])
83
84
       #df <- data.frame(paste0(dSymbol,'|',row.names(d)),dlogFC)</pre>
85
       df <- data.frame(row.names(d), d$logFC)</pre>
86
       write.table(df,row.names=FALSE,col.names=FALSE,
87
                    quote=FALSE,sep="\t",file=paste0(rnk.file, ".rnk"))
88
89 }
```

### 1.5 Quality control

```
5 setwd(wd)
 affyBatch_MAS5 <- call.exprs(affyBatch,"mas5")</pre>
   qcs <- qc(affyBatch, affyBatch_MAS5)</pre>
 8 plot(qcs)
 9 qcs
An object of class "QCStats"
Slot "scale.factors":
 [1] 0.4624769 0.9923886 0.5749658 0.5299016 0.4725059 0.4445994 1.4066966
 [8] 1.2441075 1.3089160 1.9466663 2.1150710 2.3084671
Slot "target":
[1] 100
Slot "percent.present":
  HPB_DMSO_01.present
                        HPB_DMSO_02.present
                                               HPB_DMSO_03.present
             45.75583
                                                           44.97485
                                    41.61500
 HPB_SAHM1_01.present
                       HPB_SAHM1_02.present
                                              HPB_SAHM1_03.present
             45.39369
                                    45.37906
                                                           46.93004
 KOPT_DMSO_01.present
                       KOPT_DMSO_02.present
                                              KOPT_DMSO_03.present
             39.18793
                                    39.72016
                                                           40.17558
KOPT_SAHM1_01.present KOPT_SAHM1_02.present KOPT_SAHM1_03.present
             38.46182
                                    38.39049
                                                           38.57888
Slot "average.background":
  HPB_DMSO_01
                HPB_DMSO_02
                                            HPB_SAHM1_01 HPB_SAHM1_02
                               HPB_DMSO_03
     71.40948
                   51.82766
                                  67.15371
                                                73.25355
                                                               78.06755
HPB SAHM1 03 KOPT DMSO 01
                             KOPT_DMSO_02
                                            KOPT_DMSO_03 KOPT_SAHM1_01
     64.75102
                   58.67351
                                  58.61665
                                                56.23122
                                                               53.46125
KOPT_SAHM1_02 KOPT_SAHM1_03
     44.98804
                   45.99328
Slot "minimum.background":
                HPB_DMSO_02
  HPB_DMSO_01
                               HPB_DMSO_03 HPB_SAHM1_01 HPB_SAHM1_02
     67.78476
                   49.90441
                                  64.95388
                                                68.98329
                                                               74.59893
HPB_SAHM1_03 KOPT_DMSO_01
                             KOPT_DMSO_02 KOPT_DMSO_03 KOPT_SAHM1_01
                                  54.42714
                                                53.83356
                                                               50.28008
     61.72838
                   56.40174
KOPT_SAHM1_02 KOPT_SAHM1_03
     42.42914
                   43.87834
Slot "maximum.background":
  HPB_DMSO_01
                HPB_DMSO_02
                                            HPB_SAHM1_01 HPB_SAHM1_02
                               HPB_DMSO_03
     73.33486
                   52.82101
                                  69.57489
                                                75.70203
                                                               79.83810
HPB_SAHM1_03 KOPT_DMSO_01
                             KOPT_DMSO_02
                                            KOPT_DMSO_03 KOPT_SAHM1_01
     66.69588
                   60.24642
                                  60.27544
                                                57.78016
                                                               54.91569
KOPT_SAHM1_02 KOPT_SAHM1_03
     45.78716
                   46.82307
```

91 <sub>0</sub> +	"spikes"	
DIOL	phivep	٠

	${\tt AFFX-r2-Ec-bioB-3\_at}$	${\tt AFFX-r2-Ec-bioC-3\_at}$	AFFX-r2-Ec-bioD-3_at
HPB_DMSO_01	8.248482	9.704242	12.15884
HPB_DMSO_02	9.488241	11.021774	13.39970
HPB_DMSO_03	8.174298	9.589539	12.00550
HPB_SAHM1_01	8.381753	9.772156	12.21394
HPB_SAHM1_02	8.159541	9.712199	12.13458
HPB_SAHM1_03	7.898193	9.390437	11.90994
KOPT_DMSO_01	8.920814	8.512096	12.74309
KOPT_DMSO_02	9.007404	8.715061	12.82763
KOPT_DMSO_03	8.978651	8.593016	12.75616
KOPT_SAHM1_01	9.710773	9.355043	13.32068
KOPT_SAHM1_02	10.167828	9.646911	13.70326
KOPT_SAHM1_03	10.414865	9.875183	13.89010
	AFFX-r2-P1-cre-3_at		
HPB_DMSO_01	13.20270		
HPB_DMSO_02	14.44458		
HPB_DMSO_03	13.18265		
HPB_SAHM1_01	13.31337		
HPB_SAHM1_02	13.19213		
HPB_SAHM1_03	12.97417		
KOPT_DMSO_01	14.04051		
KOPT_DMSO_02	14.10125		
KOPT_DMSO_03	14.00066		
KOPT_SAHM1_01	14.63294		
KOPT_SAHM1_02	14.82687		
KOPT_SAHM1_03	15.01834		

Slot "qc.prob	es":	
	AFFX-HSAC07/X00351_3_at	AFFX-HSAC07/X00351_5_at
HPB_DMSO_01	12.73093	11.92493
HPB_DMSO_02	13.63221	12.27401
HPB_DMSO_03	12.77779	11.90235
HPB_SAHM1_01	12.79605	11.63738
HPB_SAHM1_02	12.66274	11.31439
HPB_SAHM1_03	12.56844	11.59679
KOPT_DMSO_01	13.53358	13.02196
KOPT_DMSO_02	13.49021	13.00410
KOPT_DMSO_03	13.50476	12.99276
KOPT_SAHM1_01	13.78372	13.04395
KOPT_SAHM1_02	13.85151	12.82967
KOPT_SAHM1_03	13.88275	12.84090
	AFFX-HSAC07/X00351_M_at	AFFX-HUMGAPDH/M33197_3_at
HPB_DMSO_01	12.19455	12.87855
HPB_DMSO_02	12.84310	13.87735
HPB_DMSO_03	12.24595	12.89411
HPB_SAHM1_01	12.17076	12.98237

HPB_SAHM1_02	11.91872	12.87606
HPB_SAHM1_03	11.97960	12.69856
KOPT_DMSO_01	13.21759	13.80409
KOPT_DMSO_02	13.20014	13.72550
KOPT_DMSO_03	13.16413	13.78359
KOPT_SAHM1_01	13.33575	14.13775
KOPT_SAHM1_02	13.31856	14.20714
KOPT_SAHM1_03	13.29794	14.34094
AFFX-HUN	MGAPDH/M33197_5_at AFF	X-HUMGAPDH/M33197_M_at
HPB_DMSO_01	12.86110	12.72697
HPB_DMSO_02	13.80521	13.68863
HPB_DMSO_03	12.86872	12.70080
HPB_SAHM1_01	12.84679	12.78306
HPB_SAHM1_02	12.80222	12.66854
HPB_SAHM1_03	12.67308	12.53523
KOPT_DMSO_01	13.68232	13.59723
KOPT_DMSO_02	13.74016	13.55294
KOPT_DMSO_03	13.71389	13.56799
KOPT_SAHM1_01	14.11439	13.93807
KOPT_SAHM1_02	14.20615	13.97154
KOPT_SAHM1_03	14.29181	14.05688
Slot "bioBCalls":		
HPB_DMSO_01.present	HPB_DMSO_02.present	HPB_DMSO_03.present
"P"	"P"	"P"
HPB_SAHM1_01.present	HPB_SAHM1_02.present	-
"P"	"P"	"P"
KOPT_DMSO_01.present	KOPT_DMSO_02.present	-
"P"	"P"	•
KOPT_SAHM1_01.present	KOPT_SAHM1_02.present	KOPT_SAHM1_03.present

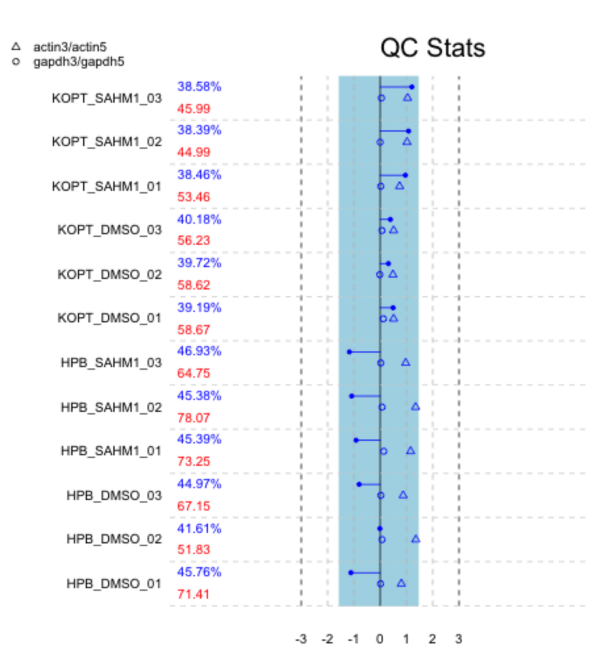
Slot "arraytype":

[1] "hgu133plus2cdf"

"P"

"P"

"P"



### 1.6 Load raw data

```
Script 1.6.1 (R)

1  %%R
2  setwd("GSE18198_data")
3  affyBatch_HPB = import_CEL("HPB*")
4  affyBatch_KOPT = import_CEL("KOPT*")
5  setwd(wd)
```

### 1.7 Create expression sets

```
Script 1.7.1 (R)

1  %%R
2  eset_HPB <- create_eset(affyBatch_HPB)
3  eset_KOPT <- create_eset(affyBatch_KOPT)
```

```
background correction: rma
normalization: quantiles
PM/MM correction : pmonly
expression values: medianpolish
background correcting...done.
normalizing...done.
54675 ids to be processed
|#########|
background correction: rma
normalization: quantiles
PM/MM correction : pmonly
expression values: medianpolish
background correcting...done.
normalizing...done.
54675 ids to be processed
| #####################
```

```
Script 1.7.2 (R)

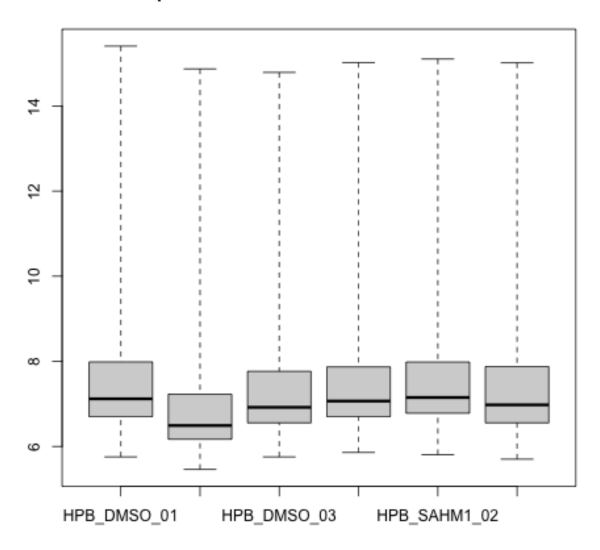
1  %%R
2  save(eset_HPB, file="eset_HPB.RData")
3  save(eset_KOPT, file="eset_KOPT.RData")
```

### 1.8 Quality plots

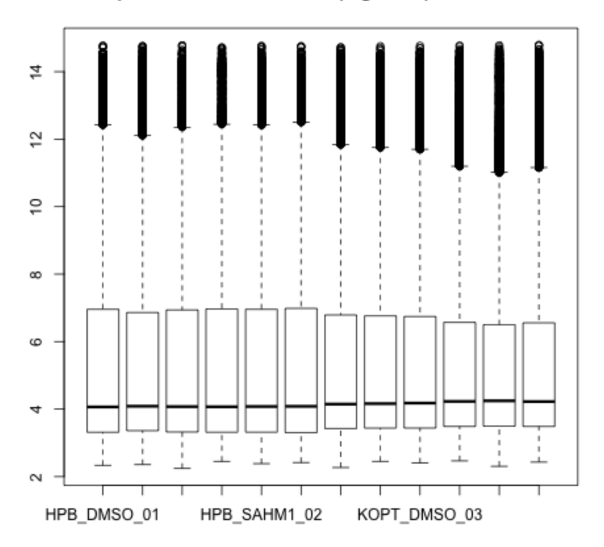
```
Script 1.8.1 (R)

1  %%R
2  boxplots(affyBatch_HPB, eset, "HPB Cell Line")
3  boxplots(affyBatch_KOPT, eset, "KOPT Cell Line")
```

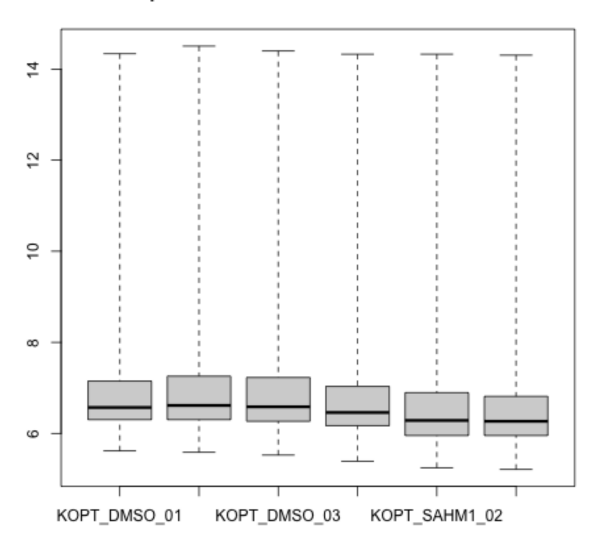
### **Boxplot Before Normalization HPB Cell Line**



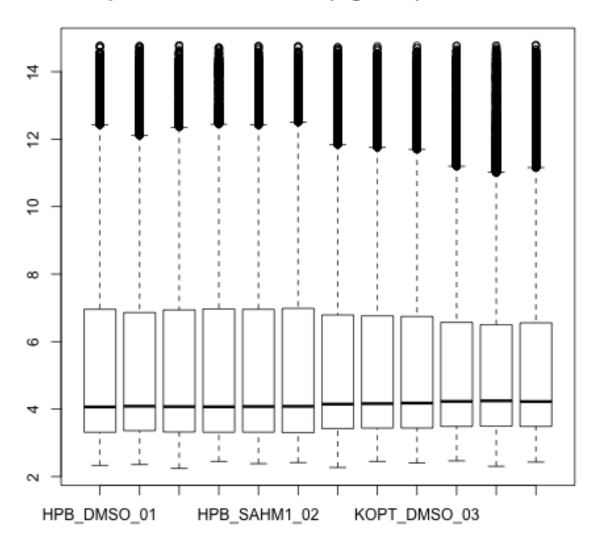
### Boxplot After Normalization (log scale) HPB Cell Line



### **Boxplot Before Normalization KOPT Cell Line**



### Boxplot After Normalization (log scale) KOPT Cell Line



### 1.9 Differential expressed genes

## [1] "Down-regulated genes" Symbol

	Symbol	logFC	adj.P.Val
225342_at	AK4	-2.1078858	1.128982e-05
230710_at	MIR210HG	-1.9761492	7.688286e-06
227336_at	DTX1	-1.3841443	8.636324e-05
201842_s_at	EFEMP1	-1.3678548	6.004820e-05
204348_s_at	AK4	-1.3491612	3.350727e-04
227347_x_at	HES4	-1.2640712	1.228528e-04
200953_s_at	CCND2	-1.2357909	7.997079e-05
202464_s_at	PFKFB3	-1.1937490	1.228528e-04
202022_at	ALDOC	-1.1753644	1.248092e-04
240546_at	LINC01120	-1.1131135	1.552628e-04
227337_at	ANKRD37	-1.1079168	2.110727e-04
200894_s_at	FKBP4	-1.0737202	5.300330e-04
217078_s_at	CD300A	-1.0719092	8.239876e-04
201170_s_at	BHLHE40	-1.0402130	2.549012e-04
202934_at	HK2	-1.0377714	2.104071e-04
201848_s_at	BNIP3	-1.0052154	6.744018e-04
218051_s_at	NT5DC2	-1.0049990	2.851346e-03
203394_s_at	HES1	-0.9993973	7.294799e-04
219371_s_at	KLF2	-0.9800918	3.918155e-04
201251_at	PKM	-0.9787897	4.328581e-03
201849_at	BNIP3	-0.9724936	6.167814e-04
213746_s_at	FLNA	-0.9705676	1.430899e-03
201516_at	SRM	-0.9428084	1.627743e-03
203523_at	LSP1	-0.9360851	2.263449e-03
225944_at	NLN	-0.9317256	1.421369e-03
214183_s_at	TKTL1	-0.9290362	2.452591e-03
236180_at	NA	-0.9287828	3.049302e-03
201194_at	SELENOW	-0.9285463	6.167814e-04
231922_at	ZNF276	-0.9269927	4.098501e-03
209933_s_at	CD300A	-0.9049911	8.239876e-04
214752_x_at	FLNA	-0.9018607	1.636952e-03
226348_at	FUT11	-0.8976507	1.075839e-03
201212_at	LGMN	-0.8972886	1.381530e-03
218305_at	IPO4	-0.8783786	1.923264e-03
205544_s_at	CR2	-0.8502424	1.206402e-03
202145_at	LY6E	-0.8406636	2.534145e-03
200859_x_at	FLNA	-0.8368746	2.851346e-03
202887_s_at	DDIT4	-0.8271607	1.813467e-03
200965_s_at	ABLIM1	-0.8229490	1.522219e-03
203504_s_at	ABCA1	-0.8211483	1.430899e-03
208116_s_at	MAN1A1	-0.7931451	4.037398e-03
202472_at	MPI	-0.7831169	2.929531e-03
207543_s_at	P4HA1	-0.7783781	2.263449e-03
201563_at	SORD	-0.7663457	4.281959e-03
222150_s_at	GSAP	-0.7532244	4.098501e-03
207539_s_at	IL4	-0.7404704	4.098501e-03

```
NOLC1 -0.7357033 4.328581e-03
205895_s_at
219389_at
                SUSD4 -0.7347418 3.852969e-03
201562_s_at
                 SORD -0.7343969 4.098501e-03
218984_at
                 PUS7 -0.7076878 4.281959e-03
[1] "Up-regulated genes"
                  Symbol
                              logFC
                                       adj.P.Val
204962_s_at
                      NA 0.8016614 1.927646e-03
222670_s_at
                    MAFB 0.8314472 1.414249e-03
244075_at
                      NA 0.8555977 1.927646e-03
205047_s_at
                    ASNS 0.8566562 1.373571e-03
                      NA 0.8615325 1.373571e-03
236153_at
                    CHD2 0.8640573 1.731300e-03
228999_at
                    PCK2 0.8839066 1.373571e-03
202847_at
242388_x_at
                   TAGAP 0.8933876 1.116736e-03
243368_at
                      NA 0.9077303 1.731300e-03
1558212_at
                      NA 0.9381646 1.381530e-03
212907_at
                 SLC30A1 0.9572462 5.305252e-04
241505_at
                      NA 0.9630797 1.430899e-03
                      NA 0.9742286 8.892420e-04
230659_at
203279_at
                   EDEM1 0.9785129 3.944123e-04
218923_at
                    CTBS 0.9839137 1.116736e-03
1558920_at
              SLC8A1-AS1 0.9839933 1.272470e-03
215071_s_at
               HIST1H2AC 0.9867407 1.634677e-03
                 SLC7A11 1.0002602 6.787587e-04
217678_at
235795_at
                    PAX6 1.0066337 6.744018e-04
                   FXYD2 1.0219109 1.381530e-03
1556294_at
                  IQGAP3 1.0315469 5.922793e-04
229538_s_at
206864_s_at
                     HRK 1.0324569 1.969355e-03
                  ZNF652 1.0513033 1.522219e-03
243495_s_at
218145_at
                   TRIB3 1.0673820 2.149263e-04
                  TM6SF1 1.0684116 7.475097e-04
219892_at
244377_at
                  SLC1A4 1.0880295 2.042060e-04
201010_s_at
                   TXNIP 1.1062690 1.552628e-04
209921_at
                 SLC7A11 1.1333321 1.248092e-04
209822_s_at
                   VLDLR 1.1350680 2.104071e-04
230795_at
                      NA 1.1699268 2.104071e-04
213931_at
                      NA 1.1702639 3.804126e-04
201009_s_at
                   TXNIP 1.2328092 1.663743e-04
                      NA 1.2501748 1.080676e-03
244042_x_at
218559_s_at
                    MAFB 1.2553755 5.305252e-04
                  CREBRF 1.2981952 4.382754e-04
225957_at
                   FLRT3 1.3254635 1.138577e-04
222853_at
219270_at
                   CHAC1 1.3362724 3.651273e-05
202672_s_at
                    ATF3 1.3481402 3.651273e-05
218280_x_at
                      NA 1.3495011 1.272470e-03
207076_s_at
                    ASS1 1.4918960 3.651273e-05
201008_s_at
                   TXNIP 1.4920023 1.663743e-04
            LOC100130476 1.5055489 4.060560e-04
243871_at
```

```
201464_x_at
                      JUN 1.5541848 7.688286e-06
                      NA 1.5966520 1.522219e-03
236962_at
235412_at
                 ARHGEF7 1.5978442 2.999663e-04
229541_at
                      NA 1.6143542 2.042060e-04
                  RASSF6 1.6458323 7.688286e-06
229147_at
235638_at
                  RASSF6 1.7804932 7.997079e-05
201466_s_at
                      JUN 2.1333931 2.108830e-06
201465_s_at
                      JUN 2.2743608 1.053822e-05
```

### Script 1.9.2 (R)

```
1 %%R
```

- toptable\_KOPT <- create\_TopTable(eset\_KOPT)</pre>
- 3 toptable\_anot\_KOPT <- anotate\_TopTable(toptable\_KOPT)</pre>
- generank\_table(toptable\_anot\_KOPT, "generank\_KOPT")

#### [1] "Down-regulated genes"

```
Symbol
                           logFC
                                    adj.P.Val
              SLC7A11 -2.916468 2.066610e-11
209921_at
205047_s_at
                 ASNS -2.757747 2.764706e-11
209369_at
                ANXA3 -2.661459 2.066610e-11
219270_at
                CHAC1 -2.264373 7.569090e-10
226517_at
                BCAT1 -2.216157 2.899776e-10
                BCAT1 -2.166267 3.402665e-09
214452_at
225285_at
                BCAT1 -2.072317 2.630231e-10
217678_at
              SLC7A11 -2.058710 7.569090e-10
230748_at
              SLC16A6 -2.004791 6.729951e-09
219892_at
               TM6SF1 -1.957664 1.012031e-09
220892_s_at
                PSAT1 -1.938100 4.136414e-10
204351_at
                S100P -1.902944 5.924315e-09
223195_s_at
                SESN2 -1.889417 7.569090e-10
214079_at
                DHRS2 -1.873374 5.924315e-09
209822_s_at
                VLDLR -1.863734 4.214321e-09
212290_at
               SLC7A1 -1.860735 2.786337e-08
202847_at
                 PCK2 -1.828534 1.164949e-09
225520_at
                   NA -1.790127 1.147212e-09
223062_s_at
                PSAT1 -1.786577 1.164949e-09
223196_s_at
                SESN2 -1.730910 3.463629e-08
200924_s_at
               SLC3A2 -1.698282 2.186020e-08
1553972_a_at
                  CBS -1.682036 3.595785e-09
                 ASS1 -1.675254 2.113541e-08
207076_s_at
229787_s_at
                  OGT -1.662192 1.743772e-07
222632_s_at
               LZTFL1 -1.643945 9.106004e-08
212816_s_at
                  CBS -1.599989 1.068633e-07
224839_s_at
                 GPT2 -1.553796 1.734549e-08
```

```
215411_s_at
             TRAF3IP2 -1.540251 2.113541e-08
             EIF4EBP1 -1.533442 1.149078e-08
221539_at
201195_s_at
               SLC7A5 -1.511384 1.336464e-08
214390_s_at
                BCAT1 -1.459915 2.424870e-07
                 ATF5 -1.451634 2.113541e-08
204999_s_at
210512_s_at
                VEGFA -1.448871 5.654616e-08
200878_at
                EPAS1 -1.435966 3.072051e-07
                CEBPB -1.428290 2.844253e-08
212501_at
224580_at
              SLC38A1 -1.428164 1.068633e-07
204744_s_at
                 IARS -1.419418 2.844253e-08
                 GARS -1.406661 4.090508e-08
208693_s_at
              FAM107B -1.392809 5.070038e-08
223059_s_at
218437_s_at
               LZTFL1 -1.332751 1.769993e-07
1558212_at
                   NA -1.308392 1.982077e-07
205653_at
                 CTSG -1.279331 2.502696e-07
217078_s_at
               CD300A -1.278830 2.424870e-07
203627_at
                IGF1R -1.275358 3.751987e-07
               NLGN4X -1.272235 4.113388e-07
221933_at
231894_at
                 SARS -1.263321 2.424870e-07
214095_at
                SHMT2 -1.235111 1.982077e-07
201263_at
                 TARS -1.197960 3.751987e-07
226181_at
                TUBE1 -1.189891 3.344154e-07
[1] "Up-regulated genes"
                           Symbol
                                       logFC
                                                adj.P.Val
224429_x_at
                                NA 0.9259213 6.376426e-06
220725_x_at
                            DNAH3 0.9423323 6.447180e-06
210686_x_at
                         SLC25A16 0.9631964 4.491576e-06
213605_s_at
                                NA 0.9632616 6.833714e-06
1556206_at
                        LINC00408 0.9653901 4.353339e-06
                               NA 0.9679377 6.750428e-06
244114_x_at
241632_x_at
                               NA 0.9686426 4.885508e-06
239017_at
                               NA 0.9834736 7.316357e-06
                        LINCO2053 0.9853247 5.091816e-06
1558496_at
                             NPAT 0.9926606 6.429684e-06
211585_at
236389_x_at
                                NA 1.0027588 4.491576e-06
208120_x_at
                           FKSG49 1.0184438 2.815681e-06
206323_x_at
                            OPHN1 1.0212889 5.601191e-06
                           FKSG49 1.0251308 3.950099e-06
224284_x_at
201464_x_at
                               JUN 1.0348832 3.490668e-06
                        LINC01949 1.0397500 4.491576e-06
220828_s_at
                     LOC100505915 1.0462560 5.662707e-06
81737_at
210800_at
                           TIMM8A 1.0479516 4.730744e-06
215182_x_at
                               NA 1.0559763 4.491576e-06
243489_at
                                NA 1.0626878 4.070717e-06
240988_x_at
                                NA 1.0629756 4.779299e-06
224288_x_at
                           FKSG49 1.0813463 1.113017e-06
AFFX-r2-Ec-bioB-5_at
                               NA 1.1020958 5.322529e-06
```

CARS -1.552793 5.615021e-08

240983\_s\_at

```
242862_x_at
                               NA 1.1113007 1.113017e-06
                            FCRL2 1.1127799 6.945728e-06
1563674_at
AFFX-BioC-5_at
                               NA 1.1199525 6.430788e-06
210718_s_at
                               NA 1.1396478 6.670158e-06
                               NA 1.1425399 2.235176e-06
1562755_at
232964_at
                               NA 1.1460961 1.656041e-06
220232_at
                             SCD5 1.1545960 1.183928e-06
1569940_at
                          SLC6A16 1.1781696 8.464513e-07
                          FKSG49 1.1800267 5.834686e-07
211454_x_at
AFFX-BioB-M_at
                               NA 1.1849198 6.376426e-06
209700_x_at
                          PDE4DIP 1.1871538 8.406263e-07
                               NA 1.1918754 3.003518e-07
1566145_s_at
                           FRG1BP 1.1922760 1.231065e-06
234949_at
                               NA 1.2077566 1.790306e-06
1560144_at
                            RASEF 1.2129105 6.160739e-07
1553185_at
231597_x_at
                               NA 1.2169048 1.477715e-06
242619_x_at
                               NA 1.2257718 4.302148e-07
1553186_x_at
                            RASEF 1.2330332 6.881976e-07
227952_at
                           ZNF595 1.2408024 5.034675e-07
1561754_at
                               NA 1.2457988 9.800827e-07
224159_x_at
                            TRIM4 1.2471076 7.904835e-07
                           FRG1BP 1.3748167 6.160739e-07
243689_s_at
231598_x_at
                               NA 1.4418759 1.469974e-07
228919_at
                               NA 1.4589605 1.682330e-06
1562527_at
                        L0C441666 1.4936054 7.062932e-08
                               NA 1.5005510 2.186020e-08
1558048_x_at
                           SH3GL3 1.5671795 2.844253e-08
211565_at
```

### 1.10 Generate GSEA gct, cls files

```
4 echo "GID
                                                                         HPB_DMSO_03
                                                                                             HPB_SA
                                 HPB_DMSO_01
                                                     HPB_DMSO_02
                     HPB_SAHM1_02
                                    HPB_SAHM1_03" >>
   \hookrightarrow HM1_01
   \rightarrow gct.head.HPB
6 echo "#1.2" > gct.head.KOPT
  echo "$(cat eset_KOPT.tsv | wc -1) 6" >> gct.head.KOPT
8 echo "GID
                   NAME
                                                                                                KOP |
                                 KOPT_DMSO_01
                                                      KOPT_DMSO_02
                                                                           KOPT_DMSO_03
                                              KOPT_SAHM1_03" >>
   \rightarrow T_SAHM1_01
                         KOPT_SAHM1_02

→ gct.head.KOPT

10 cat gct.head.HPB eset_HPB.tsv > eset_HPB.gct
cat gct.head.KOPT eset_KOPT.tsv > eset_KOPT.gct
12
13 echo "6
                  2
                           1" > phenotypes.cls
echo "#DMSO SAHM1" >> phenotypes.cls
                  0
                            0
                                                        1" >> phenotypes.cls
```

### 1.11 Processing all samples

```
background correction: rma
normalization: quantiles
PM/MM correction : pmonly
expression values: medianpolish
background correcting...done.
normalizing...done.
54675 ids to be processed
1
| ########### |
[1] "Down-regulated genes"
               Symbol
                           logFC
                                      P.Value
227347_x_at
                 HES4 -1.2391846 4.541483e-05
227336_at
                 DTX1 -1.0787341 1.772044e-03
230263_s_at
               DOCK5 -1.0589452 2.174812e-04
218051_s_at
              NT5DC2 -0.9507155 1.126824e-03
205544_s_at
                  CR2 -0.9438572 5.961088e-05
202464_s_at
              PFKFB3 -0.9408542 1.077045e-04
```

```
PDK1 -0.8325181 2.019305e-05
226452_at
223364_s_at
                DHX37 -0.8207408 1.605856e-03
                 PDK1 -0.8172945 9.936732e-04
206686_at
203627_at
                IGF1R -0.8050123 1.277490e-03
203867_s_at
                 NLE1 -0.7963720 7.746991e-04
                ELM01 -0.7885410 1.839908e-04
204513_s_at
207543_s_at
                P4HA1 -0.7663495 3.287902e-05
212063_at
                 CD44 -0.7621344 1.769410e-03
227337_at
              ANKRD37 -0.7620441 3.022164e-04
239410_at
                  HK2 -0.7615742 2.557385e-03
1554918_a_at
                ABCC4 -0.7263246 1.912084e-03
231094_s_at
                   NA -0.7232606 9.131629e-04
200965_s_at
               ABLIM1 -0.7085777 6.009393e-04
210625_s_at
                AKAP1 -0.6987526 1.183409e-04
231310_at
               TRIM71 -0.6679915 6.024183e-04
             TMEM185B -0.6494920 1.313180e-03
219253_at
215195_at
                PRKCA -0.6258312 2.255091e-04
228205_at
                  TKT -0.6126859 8.915350e-04
                 VAV3 -0.5936019 2.050231e-03
218806_s_at
206923 at
                PRKCA -0.5924819 2.694018e-04
201367_s_at
              ZFP36L2 -0.5879670 1.331411e-03
236180_at
                   NA -0.5535885 1.051495e-03
221989_at
                   NA -0.5531493 1.744579e-04
223058_at
              FAM107B -0.5499447 2.198129e-03
208858_s_at
                ESYT1 -0.5411513 2.844208e-04
1555434_a_at SLC39A14 -0.5382917 2.440482e-03
1553138_a_at
               ANKLE1 -0.5306391 5.104033e-04
227099_s_at C11orf96 -0.5289498 2.627354e-03
203612_at
                 BYSL -0.5257364 1.818973e-04
226938_at
                DCAF4 -0.5161954 1.874088e-03
              SIGMAR1 -0.5153579 1.465927e-03
214484_s_at
206653_at
               POLR3G -0.5067532 2.177288e-03
226498_at
                 FLT1 -0.5041981 2.319967e-03
                 ATIC -0.4943349 1.053895e-03
208758_at
208997_s_at
                 UCP2 -0.4905291 2.479831e-03
                WDR18 -0.4862286 8.241993e-04
209461_x_at
201161_s_at
                 YBX3 -0.4839871 9.730003e-04
225883_at
              ATG16L2 -0.4745402 2.113101e-03
201692_at
              SIGMAR1 -0.4727785 1.380736e-03
217139_at
                VDAC1 -0.4553310 2.588750e-03
                SFXN4 -0.4251656 1.858039e-03
229236_s_at
                COX20 -0.4173264 2.375617e-03
224824_at
204027_s_at
               METTL1 -0.4116541 2.046282e-03
201250_s_at
               SLC2A1 -0.3947427 2.225655e-03
[1] "Up-regulated genes"
                   Symbol
                               logFC
                                          P. Value
232059_at
                  DSCAML1 0.3346000 0.0068497056
209392_at
                    ENPP2 0.3540105 0.0052561029
```

205381_at	T DDC17	0 26/57/0	0.0064690771
214710_s_at			0.0004090771
1569680_at			0.0073803254
1559023_a_at		0.3845018	
236353 at			0.0082999985
206448_at			0.0032538740
226936_at			0.0055882829
243469_at			0.0048561664
201896_s_at			0.0084642203
1568596_a_at			0.0017179111
238875_at			0.0078367183
242966_x_at	RFX2	0.4321379	0.0027616154
236253_at	ZNF546	0.4657924	0.0075436998
1557290_at			0.0041564711
243992_at	NA	0.4761079	0.0040639088
204641_at	NEK2	0.4770373	0.0034259514
220167_s_at	NA	0.4845848	0.0045882849
241685_x_at	PURA	0.4879466	0.0058069390
239735_at	NA	0.4897705	0.0030891034
202644_s_at	TNFAIP3	0.4934166	0.0004891236
242476_at	NA	0.5030038	0.0056605223
238595_at	NA	0.5058669	0.0059647137
213605_s_at	NA	0.5068902	0.0050211914
244427_at	KIF23	0.5209074	0.0015710943
242637_at	NA	0.5222779	0.0047282688
232953_at	LINC00266-1	0.5259613	0.0041673250
1559156_at	NA	0.5325258	0.0057922093
228390_at	RAB30	0.5332846	0.0071758418
238407_at	NA	0.5390014	0.0066182068
216756_at	NA	0.5406353	0.0019487381
239248_at	SDCBP2-AS1	0.5441412	0.0010999281
213544_at	ING2	0.5513826	0.0046805162
239531_at	NA	0.5536821	0.0026133238
243709_at	SLC38A9	0.5569113	0.0009694726
241745_at	LOC100507557	0.5600510	0.0027569686
1557813_at	NA	0.5691647	0.0029037060
215599_at	NA	0.5931984	0.0012579337
244114_x_at	NA	0.6411913	0.0034400322
213281_at	JUN	0.6419108	0.0061333797
228834_at	TOB1	0.6427144	0.0006729806
244532_x_at	NA	0.6470301	0.0046383310
230795_at	NA	0.6509000	0.0039752814
216094_at	NA	0.6647132	0.0040965858
234759_at	L0C100287497	0.6758224	0.0008985553
244075_at	NA	0.6914067	0.0071383024
215071_s_at	HIST1H2AC	0.7357238	0.0038095135
210718_s_at	NA	0.7879121	0.0013751119
201465_s_at	JUN	1.5137055	0.0061879051

```
Script 1.11.2 (R)

1  %/R
2  # Create files for GSEA
3  # Add description column for gct
4  exprs2 <- cbind(c(" "), exprs(eset))
5  write.table(exprs2,row.names=TRUE,col.names=FALSE,quote=FALSE,file="eset.tsv", sep = "\t")
```

```
Script 1.11.3 (bash)
1 %%bash
2 echo "#1.2" > gct.head
get.head
4 echo "GID NAME HPB_DMSO_01 HPB_DMSO_02 HPB_DMSO_03
                                                                        HPB_SA |
            HPB_SAHM1_02 HPB_SAHM1_03 KOPT_DMSO_01 KOPT_DMSO_02
  \hookrightarrow HM1_01
     KOPT_DMSO_03 KOPT_SAHM1_01 KOPT_SAHM1_02
                                                              KOPT_SAHM1_03" >>
  \rightarrow gct.head
  #head qct.head
  cat gct.head eset.tsv > eset.gct
8 echo "12
             2
                     1" > phenotypes_all.cls
  echo "#DMSO SAHM1" >> phenotypes_all.cls
10 echo "0
              0
                    0
                                   1
                                           1
                                                  0
                                                                         1 |
                   1" >>
           1

→ phenotypes_all.cls
```

#### 1.12 GSEA results

See figures 1 to 8.

#### 1.13 Methods

### **Pipeline**

We have created the working environment under an i-python notebook of the *jupyter* platform configured to be able to execute R in code cells that start with %%R. To do so we have used the python package *rpy2*. This allows us to keep the documentation unified with the execution pipeline. It also becomes a good environment to launch hybrid pipelines with steps in R, python or even bash. It would not be difficult to develop on top a checkpoint and restart system for those developments highly time consuming.

The interface with python has not been necessary but we have developed in *bash* part of the conversions to the *gct* format and the generation of phenotype files (*.cls*)

In R we used the *affy*, *limma*, and *simpleaffy* libraries and developed a pipeline similar to the one followed in the course. We group in functions the most used methods to be able to launch in a more compact way the different experiments.

### Input data

We download from the GEO platform the raw data belonging to expression arrays Human Genome U133 Plus 2.0 with code *GSE18198*. These data have RNA information from cell cultures of the KOPT-K1 and HPB-ALL lines treated for 24h with SAHM1. Also their respective controls with the same amount of DMSO.

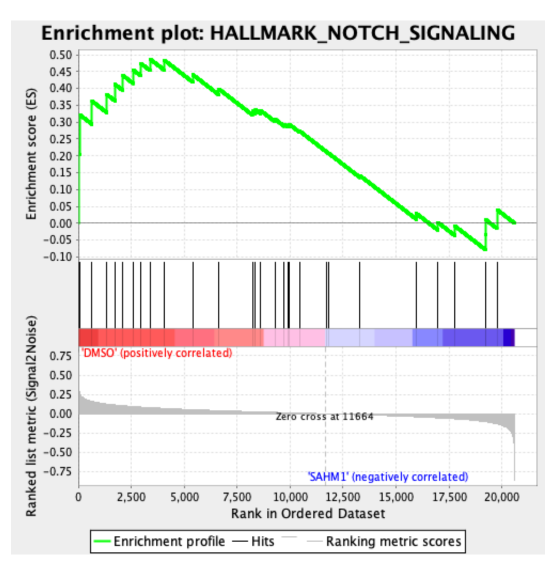


Figure 1: Enrichment plot Notch Signalling Pathway

NAME	SIZE	ES	NES	NOM p-val	FDR q-val	FWER p-val	RANK AT MAX	LEADING EDGE
HALLMARK_WNT_BETA_CATENIN_SIGNALING	40	0.49850842	1.3996072	0.0	0.9853856	277	2947	tags=23%, list=14%, signal=26%
HALLMARK_NOTCH_SIGNALING	29	0.48723552	1.3520839	0.09484536	0.7166934	374	3383	tags=31%, list=16%, signal=37%
HALLMARK_GLYCOLYSIS	186	0.42253992	1.3206861	0.0	0.64004976	473	4630	tags=35%, list=22%, signal=45%
HALLMARK_INTERFERON_ALPHA_RESPONSE	88	0.34315822	1.279187	0.096114516	0.6929842	527	4447	tags=27%, list=22%, signal=35%
HALLMARK_REACTIVE_OXIGEN_SPECIES_PATHWAY	46	0.3981605	1.2701172	0.09168444	0.5729008	527	7669	tags=57%, list=37%, signal=90%
HALLMARK_INTERFERON_GAMMA_RESPONSE	187	0.3087745	1.2654618	0.08958333	0.49679276	527	5351	tags=32%, list=26%, signal=43%
HALLMARK_ADIPOGENESIS	184	0.2826048	1.2613991	0.0911017	0.43812412	527	7541	tags=42%, list=37%, signal=65%
HALLMARK_OXIDATIVE_PHOSPHORYLATION	188	0.35985056	1.2546434	0.10103093	0.4135756	577	9489	tags=56%, list=46%, signal=104%
HALLMARK_MTORC1_SIGNALING	183	0.26467624	1.2359246	0.19502075	0.4407044	0.69	5436	tags=35%, list=26%, signal=47%
HALLMARK_IL2_STAT5_SIGNALING	182	0.31952778	1.2251521	0.0	0.44391555	0.69	2989	tags=23%, list=15%, signal=27%
HALLMARK_FATTY_ACID_METABOLISM	150	0.27257207	1.2249079	0.09663866	0.40792337	0.69	4963	tags=24%, list=24%, signal=31%
HALLMARK_PEROXISOME	98	0.30541733	1.2229942	0.091649696	0.38731882	0.69	4429	tags=26%, list=21%, signal=32%
HALLMARK_HYPOXIA	188	0.38151017	1.2137458	0.0	0.3664674	0.69	4409	tags=35%, list=21%, signal=44%
HALLMARK_UV_RESPONSE_UP	150	0.26630923	1.2116135	0.18930042	0.3437194	0.69	5540	tags=34%, list=27%, signal=46%
HALLMARK_ALLOGRAFT_REJECTION	191	0.3419296	1.1964117	0.09033614	0.3680125	739	4439	tags=28%, list=22%, signal=36%
HALLMARK_MYC_TARGETS_V2	47	0.65528375	1.1840152	0.0	0.3599941	739	4315	tags=66%, list=21%, signal=83%
HALLMARK_INFLAMMATORY_RESPONSE	189	0.34822693	1.1837027	0.097363085	0.34164155	739	4313	tags=33%, list=21%, signal=41%
HALLMARK_MYC_TARGETS_V1	173	0.4086298	1.1757741	0.19381443	0.3362698	788	7640	tags=44%, list=37%, signal=69%
HALLMARK_APICAL_JUNCTION	193	0.31684956	1.1519539	0.28781512	0.3974107	835	4296	tags=26%, list=21%, signal=33%
HALLMARK_ESTROGEN_RESPONSE_EARLY	183	0.29886743	1.119735	0.2801636	0.43835357	888	4844	tags=32%, list=24%, signal=42%
HALLMARK_ANGIOGENESIS	34	0.43509555	1.1002749	0.29411766	0.47263375	888	3335	tags=26%, list=16%, signal=32%
HALLMARK_ESTROGEN_RESPONSE_LATE	194	0.29403758	1.0978482	0.28305784	0.45791104	888	4138	tags=30%, list=20%, signal=38%
HALLMARK_MYOGENESIS	196	0.37049508	1.0864803	0.18958333	0.4909446	888	4781	tags=37%, list=23%, signal=47%
HALLMARK_CHOLESTEROL_HOMEOSTASIS	71	0.2772793	1.0851066	0.28661087	0.47539067	888	5068	tags=35%, list=25%, signal=47%
HALLMARK_COMPLEMENT	192	0.26265764	1.0589281	0.31237322	0.4991516	888	3189	tags=18%, list=15%, signal=21%
HALLMARK_PROTEIN_SECRETION	88	0.20329766	1.0475459	0.38381743	0.49959674	888	6061	tags=24%, list=29%, signal=34%
HALLMARK_ANDROGEN_RESPONSE	94	0.2609074	1.039261	0.2897959	0.52019876	0.95	5980	tags=31%, list=29%, signal=43%
HALLMARK_DNA_REPAIR	145	0.17039137	1.031202	0.2790224	0.526112	0.95	7845	tags=36%, list=38%, signal=58%
HALLMARK_KRAS_SIGNALING_DN	188	0.33348864	1.0201786	0.38655463	0.5383922	0.95	5627	tags=38%, list=27%, signal=52%
HALLMARK_UV_RESPONSE_DN	135	0.19810459	0.95020545	0.77867204	0.70491356	1.0	2015	tags=11%, list=10%, signal=12%
HALLMARK_KRAS_SIGNALING_UP	192	0.26940593	0.9457102	0.70416665	0.6904847	1.0	4576	tags=28%, list=22%, signal=35%
HALLMARK_HEDGEHOG_SIGNALING	35	0.30286688	0.94172853	0.58906883	0.6778487	1.0	3103	tags=26%, list=15%, signal=30%
HALLMARK_EPITHELIAL_MESENCHYMAL_TRANSITION	192	0.2777393	0.9384078	0.4926004	0.6730365	1.0	4711	tags=30%, list=23%, signal=38%
HALLMARK_IL6_JAK_STAT3_SIGNALING	85	0.2657967	0.93745136	0.5821501	0.6546528	1.0	4439	tags=28%, list=22%, signal=36%
HALLMARK_XENOBIOTIC_METABOLISM	194	0.2366412	0.93364125	0.68541664	0.6441654	1.0	5724	tags=29%, list=28%, signal=40%
HALLMARK_PI3K_AKT_MTOR_SIGNALING	98	0.17684619	0.9051278	0.4888438	0.69095284	1.0	2157	tags=12%, list=10%, signal=14%
HALLMARK_SPERMATOGENESIS	124	0.25800297	0.8618422	0.7	0.7451759	1.0	2215	tags=15%, list=11%, signal=17%
HALLMARK_APICAL_SURFACE	42	0.21887365	0.7824094	0.8102767	0.90781873	1.0	7749	tags=45%, list=38%, signal=72%
HALLMARK COAGULATION	133	0.23493658	0.76809555	0.9	0.89633375	1.0	7086	tags=38%, list=34%, signal=58%

Figure 2: Gene sets enriched in phenotype DMSO(Cell Line HPB-ALL)

288 <mark>988</mark>			
DMSO DMSO DMSO SAHMI SAHMI			
	SampleName		
	AK3L1	AK3L1	adenylate kinase 3-like 1
	DTX1	DTX1	GGF-containing fibulin-like extracellular matrix protein 1 deltex homolog 1 (Drosophila)
	TKTL1 CCND2	TKTL1 CCND2	transketolase—like 1 cyclin D2
	L0C54103 BHLHB2	LOC54103 BHLHB2	- basic helix-loop-helix domain containing, class B, 2 hairy and enhancer of split 4 (Drosophila)
	NKG7	HES4 NKG7	natural killer cell group 7 sequence
	ALDOC NT5DC2	ALDOC NT5DC2	aldolase C. fructose—bisphosphate 5'—nucleotidase domain containing 2
	FUT11 HTR2B	FUT11 HTR2B	5'-nucleotidase domain containing 2 fucosyltransferase 11 (alpha (1,3) fucosyltransferase) 5-hydroxytryptamine (serotonin) receptor 2B
	MLC1 C140RF148	MLC1 C140RF148	megalencephalic leukoencephalopathy with subcortical cysts 1 chromosome 14 open reading frame 148
	LSP1 /// L0C649377 F7	F7	coagulation factor VII (serum prothrombin conversion accelerator)
	ANKRD37 L0C283999	ANKRD37 L0C283999	ankyrin repeat domain 37
	TUBB4 PFKFB3	TUBB4 PFKFB3	tubulin, beta 4 6-phosphofructo-2-kinase/fructose-2,6-biphosphatase 3
	GL0XD1 L0C653127 PN0C	GLOXD1 PNOC	glyoxalase domain containing 1
	HK2 BGN	HK2 BGN	prepronociceptin hexokinase 2
	VARS CNTN5	VARS CNTN5	biglycan valyl-tRNA_synthetase contactin 5
	TPO ANKRD47	TP0 ANKRD47	thyroid peroxidase ankyrin repeat domain 47
	ANKRD13B L0C440455	ANKRD13B	ankyrin repeat domain 13B
	L0C389043 MAP3K8	L0C389043 MAP3K8	- mitogen-activated protein kinase kinase 8
	L0C51252 HDC	LOC51252 HDC	histidine decarboxylase
	TRAF4 TNS4	TRAF4 TNS4	TNF receptor-associated factor 4 tensin 4
	KLF2 PLEKHB1	KLF2 PLEKHB1	Kruppel-like factor 2 (lung) pleckstrin homology domain containing, family B (evectins) member 1
	CYP17A1 LDHC	CYP17A1 LDHC	cytochrome P450, family 17, subfamily A, polypeptide 1 lactate dehydrogenase C
	DOCK5 NAV3 /// LOC652725	DOCK5	dedicator of cytokinesis 5
	EFNA3 ADM	EFNA3 ADM	ephrin-A3 adrenomedullin
	CD40LG FCRL4	CD40LG FCRL4	(D40 ligand (TNE superfamily, member 5, hyper-ToM syndrome)
	KLRF1 PLOD1	KLRF1 PL0D1	Fc receptor-like 4 killer cell lectin-like receptor subfamily F, member 1 procollagen-lysine 1, 2-oxoglutarate 5-dioxygenase 1 Ras association (RalGDS/AF-6) domain family 6
	RASSF6 HIST1H1B	RASSF6 HIST1H1B	histone cluster 1, Hib
	HIST1H1E HIST1H1D	HIST1H1E HIST1H1D	histone cluster 1, H1e histone cluster 1, H1d
	L0C644241 FLRT3	FLRT3	fibronectin leucine rich transmembrane protein 3
	CHAC1 ASS1	CHAC1 ASS1	ChaC, cation transport regulator homolog 1 (E. coli) argininosuccinate synthetase 1
	L0C284952 JUN	L0C284952 JUN	jun oncogene
	LRRC2 DDX6	LRRC2 DDX6	leucine Fich repeat containing 2 DEAD (Asp-Glu-Ala-Asp) box polypeptide 6
	PAR5 /// SNORD108 // L0C648832	EL 135034	
	FLJ35024 HIST1H2AJ	FLJ35024 HIST1H2AJ	histone cluster 1, H2aj
	HIST2H2AA3 C30RF20	C30RF20	histone cluster 2, H2aa3 chromosome 3 open reading frame 20
	HRK ZMAT1	HRK ZMAT1	harakiri, BCL2 interacting protein (contains only BH3 domain) zinc finger, matrin type 1
	SLC8A1 IOGAP3 HIST1H3D	SLC8A1 IOGAP3	solute carrier family 8 (sodium/calcium exchanger), member 1 10 motif containing GTPase activating protein 3
	TEX9 ATF3	HIST1H3D TEX9 ATF3	histone cluster 1, H3d testis expressed sequence 9
	HIST1H1C CCDC18	HIST1H1C CCDC18	activating transcription factor 3 histone cluster 1, H1c coiled-coil domain containing 18
	EGR1 STYX /// L0C653890	EGR1	early growth response 1
	ZNF257 VLDLR	ZNF257 VLDLR	zinc finger protein 257 very low density lipoprotein receptor
	LOC57400 MAFB	LOC57400 MAFB	v—maf musculoaponeurotic fibrosarcoma oncogene homolog B (avian)
	L0C400723 /// L0C400 SLC35E4	SLC35E4	solute carrier family 35, member E4
	LOC158160 PLEKHH2	PLEKHH2	pleckstrin homology domain containing, family H (with MyTH4 domain) member 2
	RBAK HIST1H2AM	RBAK HIST1H2AM	RB-associated KRAB zinc finger histone cluster 1, H2am
	SERPINII HIST1H2AC	SERPINI1 HIST1H2AC	serpin peptidase inhibitor, clade I (neuroserpin), member 1 histone cluster 1, H2ac
	TNFSF13B DPY19L2 /// FLJ36166	TNFSF13B	tumor necrosis factor (ligand) superfamily, member 13b
	EMB BLNK	EMB BLNK	embigin homolog (mouse) B-cell linker
	SLC7A11 A0P12B	SLC7A11 A0P12B	solute carrier family 7, (cationic amino acid transporter, y+ system) member 11 aquaporin 12B
	CXORF42 C90RF126	CX0RF42 C90RF126	chromosome X open reading frame 42 chromosome 9 open reading frame 126
	HIST1H3I	HIST1H3I	histone cluster 1. H3i

Figure 3: Heatmap(Cell Line HPB-ALL)

NAME	SIZE	ES	NES	NOM p-val	FDR q-val	FWER p-val	RANK AT MAX	LEADING EDGE
HALLMARK_ANDROGEN_RESPONSE	94	0.293221	1.4200062	0.0	0.4326947	263	3888	tags=30%, list=19%, signal=37%
HALLMARK_MITOTIC_SPINDLE	185	0.386933	1.3821146	0.0	0.23884742	308	4208	tags=38%, list=20%, signal=47%
HALLMARK_MTORC1_SIGNALING	183	0.5215603	1.3441344	0.0	0.26536438	403	2776	tags=27%, list=13%, signal=31%
HALLMARK_G2M_CHECKPOINT	178	0.3373856	1.3382939	0.0	0.21027331	403	6753	tags=45%, list=33%, signal=66%
HALLMARK_PROTEIN_SECRETION	88	0.3678097	1.3335762	0.10330579	0.19757529	403	6221	tags=52%, list=30%, signal=75%
HALLMARK_MYC_TARGETS_V1	173	0.40847418	1.3236173	0.0	0.20407657	454	6363	tags=35%, list=31%, signal=51%
HALLMARK_E2F_TARGETS	173	0.33912787	1.3138137	0.0	0.20767564	454	7519	tags=48%, list=36%, signal=75%
HALLMARK_DNA_REPAIR	145	0.2616415	1.3059548	0.20245399	0.18734114	454	5134	tags=26%, list=25%, signal=34%
HALLMARK_INTERFERON_ALPHA_RESPONSE	88	0.29699197	1.3052112	0.1002004	0.17152534	454	3260	tags=26%, list=16%, signal=31%
HALLMARK_MYC_TARGETS_V2	47	0.66136664	1.2859757	0.0	0.18164015	0.5	5065	tags=62%, list=25%, signal=82%
HALLMARK_CHOLESTEROL_HOMEOSTASIS	71	0.33102265	1.2468052	0.21237114	0.22184616	604	1292	tags=15%, list=6%, signal=16%
HALLMARK_IL2_STAT5_SIGNALING	182	0.3490652	1.2443198	0.0	0.20710902	604	3186	tags=30%, list=15%, signal=35%
HALLMARK_UNFOLDED_PROTEIN_RESPONSE	104	0.6079186	1.227846	0.0	0.23289144	604	4903	tags=48%, list=24%, signal=63%
HALLMARK_ALLOGRAFT_REJECTION	191	0.26240367	1.2218562	0.0	0.23272453	604	3260	tags=23%, list=16%, signal=26%
HALLMARK_P53_PATHWAY	182	0.2778233	1.2039973	0.0	0.24092272	604	3999	tags=23%, list=19%, signal=28%
HALLMARK_HEME_METABOLISM	184	0.25559813	1.1812468	0.0	0.26794052	646	2746	tags=14%, list=13%, signal=16%
HALLMARK_PI3K_AKT_MTOR_SIGNALING	98	0.29646376	1.1782368	0.11434511	0.26249218	646	5469	tags=32%, list=27%, signal=43%
HALLMARK_GLYCOLYSIS	186	0.29053313	1.1596437	0.0	0.2764851	646	5803	tags=38%, list=28%, signal=52%
HALLMARK_HYPOXIA	188	0.26141116	1.1396848	0.2238193	0.32033673	898	3547	tags=27%, list=17%, signal=32%
HALLMARK_TGF_BETA_SIGNALING	50	0.36157095	1.1116209	0.20977597	0.39753282	945	2675	tags=24%, list=13%, signal=28%
HALLMARK_ADIPOGENESIS	184	0.2005336	1.0934803	0.22154471	0.4228268	945	5096	tags=27%, list=25%, signal=35%
HALLMARK_COMPLEMENT	192	0.25042418	1.0874768	0.18383838	0.43187666	1.0	2949	tags=19%, list=14%, signal=22%
HALLMARK_INTERFERON_GAMMA_RESPONSE	187	0.22316597	1.0854584	0.106177606	0.4260444	1.0	3581	tags=24%, list=17%, signal=29%
HALLMARK_NOTCH_SIGNALING	29	0.38857377	1.0616399	0.28716904	0.45897862	1.0	2954	tags=34%, list=14%, signal=40%
HALLMARK_FATTY_ACID_METABOLISM	150	0.21143477	1.0521207	0.28947368	0.4667693	1.0	5347	tags=33%, list=26%, signal=44%
HALLMARK_UV_RESPONSE_UP	150	0.23067562	1.0252469	0.2275574	0.5528972	1.0	2525	tags=13%, list=12%, signal=15%
HALLMARK_HEDGEHOG_SIGNALING	35	0.3210423	1.0227196	0.49588478	0.5402487	1.0	3048	tags=26%, list=15%, signal=30%
HALLMARK_TNFA_SIGNALING_VIA_NFKB	185	0.2405712	1.005899	0.40368852	0.5560466	1.0	2882	tags=20%, list=14%, signal=23%
HALLMARK_UV_RESPONSE_DN	135	0.32195312	0.99710524	0.51934826	0.5491897	1.0	3304	tags=26%, list=16%, signal=31%
HALLMARK_APOPTOSIS	159	0.20274885	0.9836564	0.39130434	0.5896945	1.0	2882	tags=18%, list=14%, signal=20%
HALLMARK_ESTROGEN_RESPONSE_EARLY	183	0.2312291	0.9552753	0.58943087	0.67311716	1.0	2938	tags=18%, list=14%, signal=21%
HALLMARK_OXIDATIVE_PHOSPHORYLATION	188	0.16545603	0.92549586	0.5323887	0.7178825	1.0	16491	tags=99%, list=80%, signal=491%
HALLMARK_PEROXISOME	98	0.16470067	0.85989714	0.7217742	0.8381856	1.0	5944	tags=28%, list=29%, signal=39%
HALLMARK_APICAL_SURFACE	42	0.24479878	0.7153503	0.81287724	0.98692274	1.0	2692	tags=21%, list=13%, signal=25%
HALLMARK_REACTIVE_OXIGEN_SPECIES_PATHWAY	46	0.101305366	0.43637094	0.9117647	1.0	1.0	4676	tags=17%, list=23%, signal=22%

Figure 4: Gene sets enriched in phenotype DMSO(Cell Line KOPT-K1)

SAHM		
SampleName		
ANXA3	ANXA3	annexin A3
SLC7A11 CHAC1	SLC7A11 CHAC1	solute carrier family 7, (cationic amino acid transporter, y+ system) member 11 ChaC, cation transport regulator homolog 1 (E. coli)
GPT2 DHRS2	GPT2 DHRS2	glutamic pyruvate transaminase (alanine aminotransferase) 2 dehydrogenase/reductase (SDR family) member 2
ASNS	ASNS	asparagine synthetase
S100P SLC16A6	S100P SLC16A6	S100 calcium binding protein P solute carrier family 16. member 6 (monocarboxylic acid transporter 7)
SESN2 ULBP1	SESN2 ULBP1	sestrin 2 UL16 binding protein 1
VLDLR	VLDLR	very low density lipoprotein receptor
PCK2 CST7	PCK2 CST7	phosphoenolpyruvate carboxykinase 2 (mitochondrial) cystatin F (leukocystatin)
CBS ASS1	CBS ASS1	cystathionine-beta-synthase argininosuccinate synthetase 1
BCAT1 KIAA1211	BCAT1 KIAA1211	branched chain aminotransferase 1, cytosolic
LZTFL1	LZTFL1	leucine zipper transcription factor-like 1 solute carrier family 4, sodium bicarbonate cotransporter, member 5
SLC4A5 JDP2	SLC4A5 JDP2	-
EPAS1 TRAF3IP2	EPAS1 TRAF3IP2	endothelial PAS domain protein 1 TRAF3 interacting protein 2
RAG1 SIPA1L2	RAG1 SIPA1L2	recombination activating gene 1
BIC	BIC	signal—induced proliferātīon—associated 1 like 2 —
FLJ35024 SYDE2	FLJ35024 SYDE2	synapse defective 1, Rho GTPase, homolog 2 (C. elegans)
MTHFD1L HES4	MTHFD1L HES4	methylenetetrahydrofolate dehydrogenase (NADP+ dependent) 1-like hairy and enhancer of split 4 (Drosophila)
CDK5R1	CDK5R1	cyclin-dependent kinase 5, regulatory subunit 1 (p35) dpy-19-like 3 (C. elegans)
DPY19L3 PLEKHC1	DPY19L3 PLEKHC1	pleckstrin homology domain containing, family C (with FERM domain) member 1
CTSG GPD2	CTSG GPD2	cathepsin G glycerol-3-phosphate dehydrogenase 2 (mitochondrial)
SLC3A2 NLGN4X	SLC3A2 NLGN4X	solute carrier family 3 (activators of dibasic and neutral amino acid transport), m neuroligin 4, X-linked
DOCK5	DOCK5	dedicator of cytokinesis 5
IGF1R L0C203427	IGF1R	insulin-like growth factor 1 receptor
MNS1 ARSG	MNS1 ARSG	meiosis-specific nuclear structural 1 arvlsulfatase G
ATF5	ATF5	activating transcription factor 5 family with sequence similarity 107, member B
FAM107B RNF125	FAM107B RNF125	ring finger protein 125
VEGF LRRC34	VEGF LRRC34	vascular endothelial growth factor leucine rich repeat containing 34
ADAM9 C60RF170	ADAM9 C60RF170	leucine rich repeat containing 34 ADAM metallopeptidase domain 9 (meltrin gamma) chromosome 6 open reading frame 170
C20RF31	C20RF31	chromosome 6 open reading frame 170 chromosome 2 open reading frame 31
ARRDC3 L0C283027	ARRDC3 L0C283027	arrestin domain containing 3
SLC6A16 DSPP	SLC6A16 DSPP	solute carrier family 6, member 16 dentin sialophosphoprotein
L0C653117 C210RF114	L0C653117 C210RF114	-
PLSCR4	PLSCR4	chromosome 21 open reading frame 114 phospholipid scramblase 4
ADH4 LOC401131	ADH4 LOC401131	alcohol dehydrogenase 4 (class II), pi polypeptide
SH3GL3 SLC36A1	SH3GL3 SLC36A1	SH3-domain GRB2-like 3 solute carrier family 36 (proton/amino acid symporter), member 1
L0C284825	L0C284825	-
L0C440117 JUN	L0C440117 JUN	jun oncogene
ALDH1B1 L0C644450	ALDH1B1 L0C644450	aldehyde dehydrogenase 1 family, member B1
ZNF528 L0C282980	ZNF528 L0C282980	zinc finger protein 528
L0C644488 RP11-151A6.2	RP11-151A6.2	-
0IT3	OIT3	oncoprotein induced transcript 3
MEFV ANKRD20A1 /// ANKRD	MEFV 02	Mediterranean fever
DKFZP667M2411 FLJ11292	DKFZP667M2411 FLJ11292	-
SLC15A1 FCRL2	SLC15A1	solute carrier family 15 (oligopeptide transporter), member 1 Fc receptor-like 2
ABCA5	FCRL2 ABCA5	ATP-binding cassette, sub-family A (ABC1), member 5 serine peptidase inhibitor-like, with Kunitz and WAP domains 1 (eppin)
SPINLW1 MGC72104	SPINLW1 MGC72104	
RBMS3 FAM90A1 /// FLJ1066	RBMS3	RNA binding motif, single stranded interacting protein
GABRB1	GABRB1	gamma-aminobutyric acid (GABA) A receptor, beta 1
MBL2 CNTNAP3 /// CNTNAP3	MBL2 B	mannose-binding lectin (protein C) 2, soluble (opsonic defect)
ZNF595 GAGE3	ZNF595 GAGE3	zinc finger protein 595 G antigen 3
C210RF42 ANKRD20A1 /// LOC39	C210RF42	chromosome 21 open reading frame 42
BAGE	BAGE	B melanoma antigen
TMEFF2 BC37295_3	TMEFF2 BC37295_3	transmembrane protein with EGF-like and two follistatin-like domains 2
GAGE2 /// GAGE4 /// L0C203274	L0C203274	-
F0X01A	F0X01A	forkhead box 01A (rhabdomyosarcoma)
WBSCR19 MLCK	WBSCR19 MLCK	Williams Beuren syndrome chromosome region 19
L0C340089 /// L0C44 L0C644963	1	

Figure 5: Heatmap(Cell Line KOPT-K1)

NAME	SIZE	ES	NES	NOM p-val	FDR q-val	FWER p-val	RANK AT MAX	LEADING EDGE
HALLMARK_GLYCOLYSIS	186	0.43311334	1.6781958	0.004192872	0.036362655	25	5630	tags=47%, list=27%, signal=64%
HALLMARK_PI3K_AKT_MTOR_SIGNALING	98	0.34280387	1.6242404	0.010121457	0.030869437	45	5884	tags=36%, list=29%, signal=50%
HALLMARK_MTORC1_SIGNALING	183	0.4759005	1.6139847	0.004158004	0.023340473	51	4779	tags=42%, list=23%, signal=54%
HALLMARK_UNFOLDED_PROTEIN_RESPONSE	104	0.548177	1.5873568	0.03992016	0.024172837	64	4153	tags=38%, list=20%, signal=48%
HALLMARK_NOTCH_SIGNALING	29	0.5714514	1.5091325	0.016597511	0.04010106	116	4882	tags=55%, list=24%, signal=72%
HALLMARK_UV_RESPONSE_UP	150	0.2950838	1.4034421	0.0041237115	0.124392934	301	2906	tags=22%, list=14%, signal=25%
HALLMARK_REACTIVE_OXIGEN_SPECIES_PATHWAY	46	0.40161857	1.3443292	0.07068607	0.21853316	471	7374	tags=57%, list=36%, signal=88%
HALLMARK_MYC_TARGETS_V2	47	0.71475863	1.3281219	0.06759443	0.23772013	531	3914	tags=74%, list=19%, signal=92%
HALLMARK_UV_RESPONSE_DN	135	0.3603946	1.306755	0.05679513	0.2657795	587	2828	tags=26%, list=14%, signal=30%
HALLMARK_HYPOXIA	188	0.35863873	1.2884322	0.043392505	0.295965	651	5635	tags=44%, list=27%, signal=60%
HALLMARK_FATTY_ACID_METABOLISM	150	0.2589689	1.2816921	0.037113402	0.2849363	661	5652	tags=33%, list=27%, signal=46%
HALLMARK_INTERFERON_ALPHA_RESPONSE	88	0.3972688	1.2601771	0.18774703	0.31406602	717	4741	tags=38%, list=23%, signal=48%
HALLMARK_ADIPOGENESIS	184	0.2585026	1.2498535	0.06625259	0.31768975	754	6432	tags=38%, list=31%, signal=54%
HALLMARK_DNA_REPAIR	145	0.27045995	1.220323	0.25440314	0.37851238	844	6445	tags=34%, list=31%, signal=49%
HALLMARK_ESTROGEN_RESPONSE_EARLY	183	0.29162228	1.2086581	0.057494868	0.39013368	877	1687	tags=14%, list=8%, signal=15%
HALLMARK_PEROXISOME	98	0.26575372	1.2073532	0.115384616	0.36860234	881	6877	tags=38%, list=33%, signal=56%
HALLMARK_MYC_TARGETS_V1	173	0.481558	1.1877586	0.38492063	0.40412134	915	7998	tags=57%, list=39%, signal=92%
HALLMARK_ANDROGEN_RESPONSE	94	0.2428143	1.1861122	0.102713175	0.38849032	915	4883	tags=31%, list=24%, signal=40%
HALLMARK_WNT_BETA_CATENIN_SIGNALING	40	0.34349075	1.1845423	0.15767635	0.37267235	916	4889	tags=38%, list=24%, signal=49%
HALLMARK_HEDGEHOG_SIGNALING	35	0.3436916	1.183618	0.18837675	0.35571563	916	1072	tags=17%, list=5%, signal=18%
HALLMARK_CHOLESTEROL_HOMEOSTASIS	71	0.33077985	1.1479423	0.17760618	0.42038524	0.95	2256	tags=18%, list=11%, signal=20%
HALLMARK_PROTEIN_SECRETION	88	0.35464782	1.1016811	0.42352942	0.5150438	972	6980	tags=42%, list=34%, signal=63%
HALLMARK_OXIDATIVE_PHOSPHORYLATION	188	0.32158187	1.0274464	0.5450902	0.69437057	986	7107	tags=34%, list=34%, signal=51%
HALLMARK_IL2_STAT5_SIGNALING	182	0.3837396	0.99643314	0.5445545	0.75440615	989	2494	tags=24%, list=12%, signal=27%
HALLMARK_ALLOGRAFT_REJECTION	191	0.3501197	0.9865181	0.5752577	0.7505662	989	4113	tags=28%, list=20%, signal=34%
HALLMARK_APICAL_SURFACE	42	0.30094463	0.95951355	0.57715434	0.7958004	993	4367	tags=31%, list=21%, signal=39%
HALLMARK_TGF_BETA_SIGNALING	50	0.286461	0.8809257	0.72745097	0.96194637	995	4256	tags=30%, list=21%, signal=38%
HALLMARK_TNFA_SIGNALING_VIA_NFKB	185	0.25158665	0.8724303	0.70178926	0.9456293	995	3627	tags=23%, list=18%, signal=27%
HALLMARK_COMPLEMENT	192	0.25019673	0.86665404	0.68937874	0.9242008	995	3369	tags=20%, list=16%, signal=24%
HALLMARK_HEME_METABOLISM	184	0.19858962	0.85600936	0.8579882	0.91439253	999	6128	tags=30%, list=30%, signal=43%
HALLMARK_INTERFERON_GAMMA_RESPONSE	187	0.29006875	0.8416879	0.63842976	0.90728647	1.0	4763	tags=33%, list=23%, signal=42%
HALLMARK_APICAL_JUNCTION	193	0.21188706	0.8365217	0.87649405	0.8868689	1.0	6030	tags=33%, list=29%, signal=46%
HALLMARK_MITOTIC_SPINDLE	185	0.31453067	0.8285555	0.678501	0.8730112	1.0	4322	tags=28%, list=21%, signal=35%
HALLMARK_MYOGENESIS	196	0.20663904	0.7016324	0.9665971	0.9819069	1.0	2350	tags=15%, list=11%, signal=17%
HALLMARK_INFLAMMATORY_RESPONSE	189	0.25482258	0.6774285	0.79761904	0.969574	1.0	1679	tags=15%, list=8%, signal=17%
HALLMARK_G2M_CHECKPOINT	178	0.19088042	0.601792	0.8170974	0.9818409	1.0	4846	tags=24%, list=24%, signal=31%
HALLMARK_E2F_TARGETS	173	0.17057835	0.59205115	0.85265225	0.9589176	1.0	7214	tags=34%, list=35%, signal=52%

Figure 6: Gene sets enriched in phenotype DMSO(Cell Lines: HPB-ALL KOPT-K1)

122 <mark>122</mark> 22	es es		
KOPT DMSO 6 KOPT DMSO 6 KOPT DMSO 6 HPB SAHM1 6 HPB SAHM1 6 KOPT SAHM1 KOPT SAHM1	AHWII		
DET OF THE PET OF THE	S .		
   점점점 <mark>류뷰뷰점점 </mark>	SampleName		
	AK3L1 ANXA3	AK3L1 ANXA3	adenylate kinase 3-like 1 annexin A3
	GPT2 DHRS2	GPT2 DHRS2	glutamic pyruvate transaminase (alanine aminotransferase) 2 dehydrogenase/reductase (SDR family) member 2
	HES4 NT5DC2	HES4 NT5DC2	hairy and enhancer of split 4 (Drosophila) 5'-nucleotidase domain containing 2
	SLC16A6 S100P	SLC16A6 S100P	solute carrier family 16, member 6 (monocarboxylic acid transporter 7) S100 calcium binding protein P
	DOCK5 BHLHB2	DOCK5 BHLHB2	dedicator of cytokinesis 5 basic helix-loop-helix domain containing, class B, 2
	SYDE2 DTX1	SYDE2 DTX1	synapse defective 1, Rho GTPase, homolog 2 (C. elegans) deltex homolog 1 (Drosophila)
	KIAA1211 CST7	KIAA1211 CST7	cvstatin F (leukocvstatin)
	HES1 HK2	HES1 HK2	hairy and enhancer of split 1, (Drosophila) hexokinase 2
	HIST1H3F RHOBTB1	HIST1H3F RHOBTB1	histone cluster 1, H3f Rho-related BTB domain containing 1
	MPI ADM	MPI ADM	mannose phosphate isomerase adrenomedullin
	CFH /// CFHR1 TKTL1	TKTL1	transketolase-like 1
	L0C653127 L0C54103	L0C54103	<u>-</u>
	PFKFB3 NLE1	PFKFB3 NLE1	6-phosphofructo-2-kinase/fructose-2,6-biphosphatase 3 notchless homolog 1 (Drosophila)
	SLC7A11 ASNS	SLC7A11 ASNS	solute carrier family 7, (cationic amino acid transporter, y+ system) member 11 asparagine synthetase
	IP04 CR2	IP04 CR2	importin 4 complement component (3d/Epstein Barr virus) receptor 2
	GLOXD1 CD300A	GLOXD1 CD300A	glyoxalase domain containing 1 CD300a molecule
		IGF1R TUBA1	insulin-like growth factor 1 receptor tubulin, alpha 1
	PDK1 CD44	PDK1 CD44	pyruvate dehydrogenase kinase, isozyme 1 CD44 molecule (Indian blood group)
	ELM01	ULBP1 ELM01	UL16 binding protein 1 engulfment and cell motility 1
	CHAC1 PPRC1	PPRC1	ChāC, cation transport regulator homolog 1 (E. coli)  peroxisome proliferative activated receptor, gamma, coactivator-related 1
		ARSG L0C441108	arylsulfatase G
	DDIT4 SRG LY6E	DDIT4 SRG LY6E	DNA-damage-inducible transcript 4
	GBP5 CTSG	GBP5 CTSG	lymphocyte antigen 6 complex, locus E guanylate binding protein 5 cathepsin G
	GZMB	GZMB ANKRD37	granzyme B (granzyme 2, cytotoxic T-lymphocyte-associated serine esterase 1) ankyrin repeat domain 37
	TBC1D9 JUN	TBC1D9 JUN	TBC1 domain family, member 9 (with GRAM domain) jun oncogene
	RASSF6 FLRT3	RASSF6 FLRT3	Ras association (RalGDS/AF-6) domain family 6 fibronectin leucine rich transmembrane protein 3
	HERC2P7 ANKRD20A1 /// ANKRD2	HERC2P7	hect domain and RLD 2 pseudogene 7
	HIST1H1B WBSCR19	HIST1H1B WBSCR19	histone cluster 1, H1b Williams Beuren syndrome chromosome region 19
	L0C401131 L0C644450	L0C401131 L0C644450	-
	HRK DSPP	HRK DSPP	harakiri, BCL2 interacting protein (contains only BH3 domain) dentin sialophosphoprotein
	BLNK SH3GL3	BLNK SH3GL3	B-cell linker SH3-domain GRB2-like 3
	L0C400723 /// L0C400 C210RF114	C210RF114	chromosome 21 open reading frame 114
	TEX9 PLSCR4	TEX9 PLSCR4	testis expressed sequence 9 phospholipid scramblase 4
		L0C283027 L0C285535	aladahain baralan, darain ambaining da-23, U L245 M-904 da-22,
	PLEKHH2 SLC36A1	PLEKHH2 SLC36A1	pleckstrin homology domain containing, family H (with MyTH4 domain) member 2 solute carrier family 36 (proton/amino acid symporter), member 1
	L0C203274 L0C282980	L0C203274 L0C282980	-
	LOC158160 RP11-151A6.2	RP11-151A6.2	champking (C.V.C.matif) limand 2
	TREML4 C80RF70	CXCL2 TREML4 C80RF70	chemokine (C—XC motif) ligand 2 triggering receptor expressed on myeloid cells-like 4 chromosome 8 open reading frame 70
	DPY19L2 /// FLJ36166 L0C653071	L0C653071	- Chromosome 6 open reading frame 76
	SLC6A16 ABCC9	SLC6A16 ABCC9	solute carrier family 6, member 16 ATP-binding cassette, sub-family C (CFTR/MRP), member 9
		L0C401321 SLC15A1	solute carrier family 15 (oligopeptide transporter), member 1
	L0C644660 L0C644488	L0C644660	- 101141 101141 AN 101490PENAMO (10112PO) (ELT) HEHIVEL A
	IMPG2 L0C284825	IMPG2 LOC284825	interphotoreceptor matrix proteoglycan 2
	LOC643452 HIST1H2AC	HIST1H2AC	histone cluster 1. H2ac
	NUDT16P LOC157627	NUDT16P L0C157627	nudix (nucleoside diphosphate linked moiety X)—type motif 16 pseudogene
	C10RF103 SPINLW1	C10RF103 SPINLW1	chromosome 1 open reading frame 103 serine peptidase inhibitor-like, with Kunitz and WAP domains 1 (eppin)
	BAGE OMD	BAGE OMD	B melanoma antigen osteomodulin
	LRRC2 ANKRD20A1 /// LOC392	LRRC2	leucine rich repeat containing 2
	HIST1H1D	HIST1H1D	histone cluster 1, H1d

Figure 7: Heatmap(Cell Lines: HPB-ALL KOPT-K1)

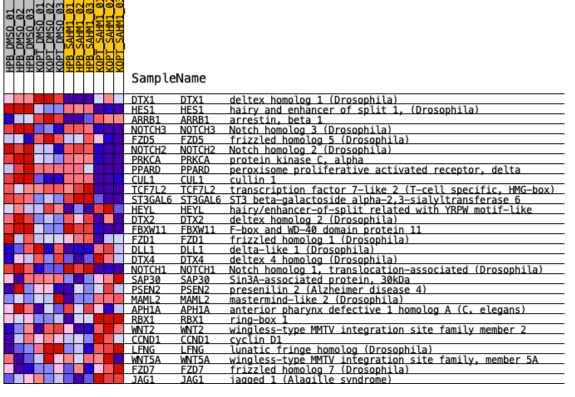


Figure 8: Heatmap(NOTCH signaling pathway)

#### **Analysis strategy**

The quality analysis is carried out with the MAS 5.0 algorithm provided by simpleaffy.

The two cell lines were analyzed separately and jointly in R-limma and in GSEA.

In R-limma we use the Benjamini, Hochberg algorithm to control the false discovery rate derived from multiple testing and obtain adjusted p-values.

The conversion to GSEA is done from the expression set objects of affybacth, without processing in R any data. To do so we had to perform the conversions to the gct format.

The results in GSEA and R are coherent, at least in regard to our target: NOTCH signaling pathway on which we have concentrated exclusively.

### 1.14 Analysis

### Quality

We use the simpleaffy R package that generates a series of metrics recommended by the manufacturer Affymetrix:

- 1. Average background
- 2. Scale factor
- 3. Percentage of genes called present.
- 4. 3 'to 5' ratios (related with RNA degradation)

It is observed that all the indicators are within the acceptance margins (see graph of section 1.5), but that the patterns are clearly different between the samples of both cell lines.

### Differential expression

We performed three different analysis to discover the effectiveness of SAHM1 in the inhibition of NOTCH.

- 1. Comparison between control (DMSO) and inhibitor (SAHM1) in the HBP-ALL cell line.
- 2. Comparison between control (DMSO) and inhibitor (SAHM1) in the KOPT-K1 cell line.
- 3. Comparison between control (DMSO) and inhibitor (SAHM1) joint for both cell lines.

It seems more correct to isolate each cell line separately in the analysis, according to the results of the quality analysis, where the expression patterns within cell lines appear more homogeneous than between. However, in figure 3 of the paper a heatmap is shown where the 12 samples seem to have been treated together, so we reproduced this analysis in case it could really show significant differences with the individual ones.

On the HPB-ALL cell line (1.9) and on the analysis of the two lines together (1.11) we found several direct targets of Notch TF among the 50 most significantly infra-regulated probes on inhibition scenario (lowest values of adjusted p-value and logFC <0): HES1, HES4, and DTX1, which are also investigated in the article. This result is also reproduced in the parallel analysis performed on GSEA. See figures 3 & 7.

GSEA also provides hallmarks that are overexpressed in the absence of inhibitory treatment, and among them we find Notch signaling (figures 2, 4,6)

These results are compatible with the expected effectiveness of SAHM1 as a NOTCH inhibitor.

In the analysis on the KOPT-K1 line, the hallmark NOTCH signaling still appears overexpressed in absence of inhibitor, but there are no traces of HES1, HES4 or DTX1. We may need to consider other NOTCH targets, but really, we have no clues to extract any conclusion in this case.