

Application of TPAC method to TCGA liver cancer RNA-seq data using MSigDB Hallmark collection

H. Robert Frost

1 Load and process TCGA liver cancer RNA-seq data

The following logic loads FPKM normalized counts for The Cancer Genome Atlas (TCGA) [1] liver cancer (LIHC) cohort. The TPAC function *tpacForCancer()* leverages Human Protein Atlas (HPA) normal tissue gene expression data ("HPA.normal.FPKM.GDCpipeline.csv") that was specially processed by the HPA group as FPKM values using a pipeline similar to that employed by GDC for the TCGA data (this data was generated for the "Human Pathology Atlas" paper [2]). For consistency with this HPA normal tissue data, the TCGA data is retrieved from the HPA provided TCGA gene expression data file *rna_cancer_sample.tsv*, which contains FPKM normalized counts and can be downloaded from https://www.proteinatlas.org/download/rna_cancer_sample.tsv.zip.

Generation of the LIHC-specific matrix from the *rna_cancer_sample.tsv* data was performed using the following R code (which is not executed here given the size of the data and processing time):

```
library(data.table)
library(reshape2)
tcga.file = "rna_cancer_sample.tsv"
data = read.table(file=tcga.file, header=T, sep="\t", stringsAsFactors=F,
                  # Columns: "Gene", "Sample", "Cancer", "FPKM"
                  colClasses=c("factor", "factor", "factor", "numeric"))
data = as.data.table(data)
setkey(data, Cancer, Gene)
lihc.data = data[.(LIHC)]
lihc.matrix = acast(lihc.data, Sample~Gene, value.var="FPKM")
saveRDS(lihc.matrix, file="lihc.rna.matrix.rds")

> # read in saved matrix
> liver.counts.fpkm = readRDS(file="lihc.rna.matrix.rds")
> # change to FPKM + 1
> liver.counts.fpkm = liver.counts.fpkm + 1
```

2 Load the MSigDB Hallmark collection

The following logic loads the MSigDB Hallmark collection using the *msigdbr* R package. The data frame returned by *msigdbr* is then converted into a list of gene ID vectors (each list element corresponds to a gene set and is a vector of Ensembl IDs). The *tpacForCancer()* function automatically transforms this into a list of vectors of gene indices using the *createGeneSetCollection()* helper function.

```
> # Load the MSigDB Hallmark collection using the msigdbr package
> hallmark.collection = msigdbr::msigdbr(category="H")
> # Create a gene.set.collection list of Ensembl IDs
```

```

> gene.set.names = unique(hallmark.collection$gs_name)
> num.sets = length(gene.set.names)
> message("Number of sets in MSigDB Hallmark collection: ", num.sets)
> gene.set.names[1:5]

[1] "HALLMARK_ADIPOGENESIS"      "HALLMARK_ALLOGRAFT_REJECTION"
[3] "HALLMARK_ANDROGEN_RESPONSE" "HALLMARK_ANGIOGENESIS"
[5] "HALLMARK_APICAL_JUNCTION"

> gene.set.collection = list()
> for (i in 1:num.sets) {
+   gene.set.name = gene.set.names[i]
+   gene.set.rows = which(hallmark.collection$gs_name == gene.set.name)
+   gene.set.ensembl.ids = hallmark.collection$human_ensembl_gene[gene.set.rows]
+   gene.set.collection[[i]] = unique(gene.set.ensembl.ids)
+ }
> names(gene.set.collection) = gene.set.names

```

3 Execute TPAC method

Since we are processing TCGA RNA-seq liver cancer data, we can execute TPAC using the `tpacForCancer()` wrapper function. Note that the cancer types supported by `tpacForCancer()` can be accessed via the `getSupportedCancerTypes()` function.

```

> # Display the full list of cancer types supported by tpacForCancer()
> TPAC::getSupportedCancerTypes()

[1] "urothelial cancer"      "breast cancer"           "cervical cancer"
[4] "colorectal cancer"     "glioma"                  "head and neck cancer"
[7] "renal cancer"          "liver cancer"            "lung cancer"
[10] "ovarian cancer"        "pancreatic cancer"       "prostate cancer"
[13] "colorectal cancer"     "melanoma"                "stomach cancer"
[16] "testis cancer"          "thyroid cancer"          "endometrial cancer"

> # Get the normal tissue corresponding to liver cancer
> cancer.type = "liver cancer"
> # Execute TPAC
> tpac.out = TPAC::tpacForCancer(cancer.gene.expr=liver.counts.fpkm,
+                                 cancer.type=cancer.type,
+                                 gene.set.collection=gene.set.collection)

```

Look at a subset of the TPAC scores in the generated S, S- and S+ matrices:

```

> tpac.out$S[1:5, 1:5]

          HALLMARK_ADIPOGENESIS HALLMARK_ALLOGRAFT_REJECTION
TCGA-2Y-A9GS-01A      3.064720e-03      3.018793e-01
TCGA-2Y-A9GT-01A      7.926992e-14      7.994741e-05
TCGA-2Y-A9GU-01A      9.918417e-01      8.723984e-01
TCGA-2Y-A9GV-01A      0.000000e+00      1.707913e-04
TCGA-2Y-A9GW-01A      5.827508e-01      3.248046e-03

          HALLMARK_ANDROGEN_RESPONSE HALLMARK_ANGIOGENESIS

```

| | | |
|------------------|--------------|--------------|
| TCGA-2Y-A9GS-01A | 2.386972e-01 | 0.7538041194 |
| TCGA-2Y-A9GT-01A | 2.586985e-06 | 0.9999999828 |
| TCGA-2Y-A9GU-01A | 4.637972e-01 | 0.0003027669 |
| TCGA-2Y-A9GV-01A | 5.906668e-04 | 0.9997277821 |
| TCGA-2Y-A9GW-01A | 9.060539e-01 | 0.9999999999 |

HALLMARK_APICAL_JUNCTION

| | |
|------------------|------------|
| TCGA-2Y-A9GS-01A | 0.24475713 |
| TCGA-2Y-A9GT-01A | 0.03069919 |
| TCGA-2Y-A9GU-01A | 0.14960836 |
| TCGA-2Y-A9GV-01A | 0.05386065 |
| TCGA-2Y-A9GW-01A | 0.41299687 |

> *tpac.out\$S.neg[1:5,1:5]*

HALLMARKADIPOGENESIS HALLMARK_ALLOGRAFT_REJECTION

| | | |
|------------------|--------------|--------------|
| TCGA-2Y-A9GS-01A | 1.642178e-03 | 0.4488871715 |
| TCGA-2Y-A9GT-01A | 2.716716e-13 | 0.0002430214 |
| TCGA-2Y-A9GU-01A | 9.848749e-01 | 0.9616570059 |
| TCGA-2Y-A9GV-01A | 0.000000e+00 | 0.0010408253 |
| TCGA-2Y-A9GW-01A | 6.410074e-01 | 0.0089029673 |

HALLMARK_ANDROGEN_RESPONSE HALLMARK_ANGIOGENESIS

| | | |
|------------------|--------------|-------------|
| TCGA-2Y-A9GS-01A | 2.649604e-01 | 0.852475176 |
| TCGA-2Y-A9GT-01A | 6.135243e-06 | 0.999999997 |
| TCGA-2Y-A9GU-01A | 5.329255e-01 | 0.001090343 |
| TCGA-2Y-A9GV-01A | 9.895708e-04 | 0.999862611 |
| TCGA-2Y-A9GW-01A | 9.256814e-01 | 1.000000000 |

HALLMARK_APICAL_JUNCTION

| | |
|------------------|------------|
| TCGA-2Y-A9GS-01A | 0.05700461 |
| TCGA-2Y-A9GT-01A | 0.01287834 |
| TCGA-2Y-A9GU-01A | 0.56693484 |
| TCGA-2Y-A9GV-01A | 0.01254875 |
| TCGA-2Y-A9GW-01A | 0.16432979 |

> *tpac.out\$S.pos[1:5,1:5]*

HALLMARKADIPOGENESIS HALLMARK_ALLOGRAFT_REJECTION

| | | |
|------------------|------------|-------------|
| TCGA-2Y-A9GS-01A | 0.85091309 | 0.232426214 |
| TCGA-2Y-A9GT-01A | 0.08466006 | 0.190018804 |
| TCGA-2Y-A9GU-01A | 0.96036440 | 0.001615544 |
| TCGA-2Y-A9GV-01A | 0.36058713 | 0.072145624 |
| TCGA-2Y-A9GW-01A | 0.16157345 | 0.193789454 |

HALLMARK_ANDROGEN_RESPONSE HALLMARK_ANGIOGENESIS

| | | |
|------------------|------------|-----------|
| TCGA-2Y-A9GS-01A | 0.37298345 | 0.1748558 |
| TCGA-2Y-A9GT-01A | 0.03279483 | 0.6134479 |
| TCGA-2Y-A9GU-01A | 0.02420909 | 0.3539980 |
| TCGA-2Y-A9GV-01A | 0.11124720 | 0.8844703 |
| TCGA-2Y-A9GW-01A | 0.20945836 | 0.3179549 |

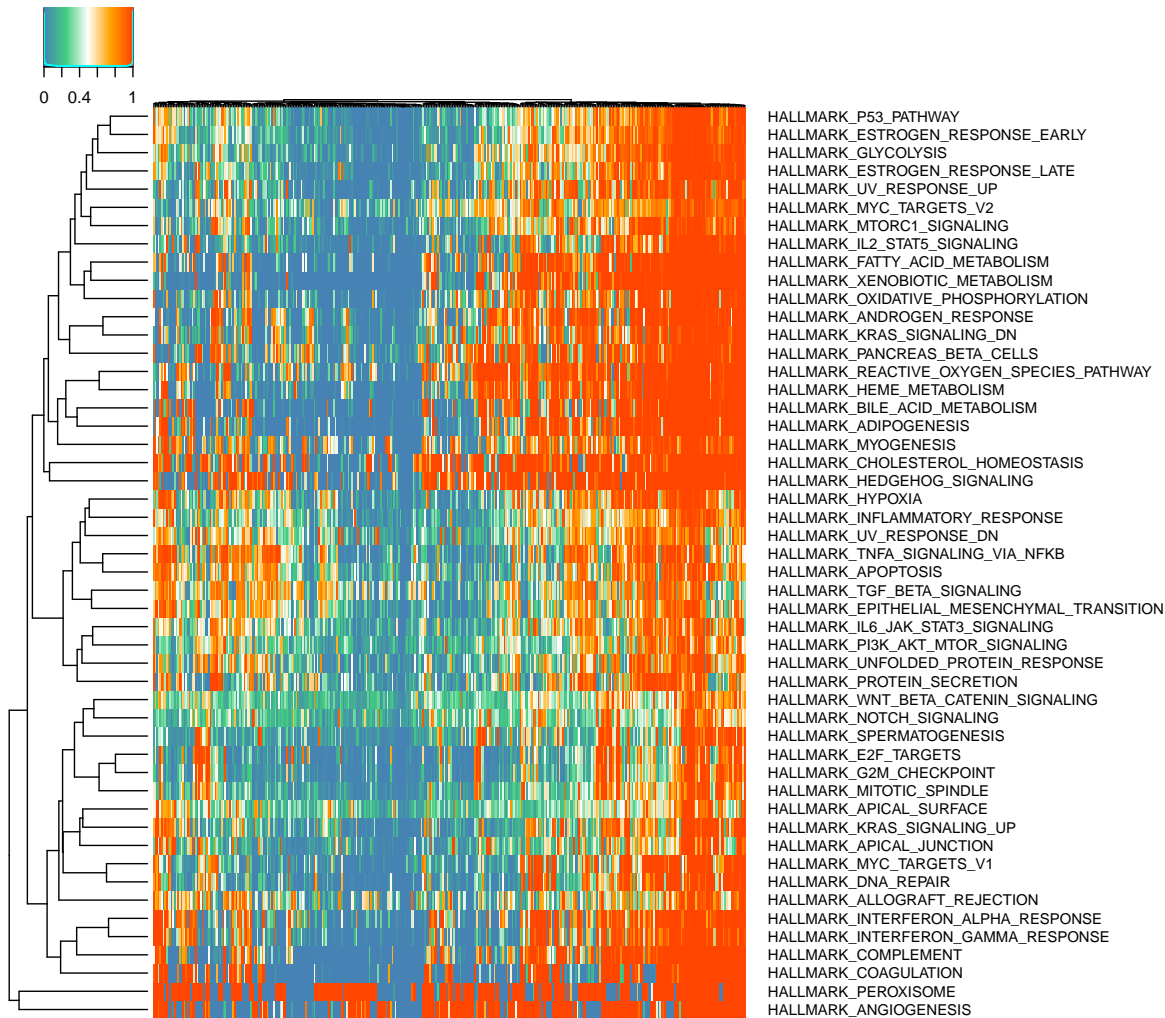
HALLMARK_APICAL_JUNCTION

| | |
|------------------|------------|
| TCGA-2Y-A9GS-01A | 0.47087802 |
| TCGA-2Y-A9GT-01A | 0.10049191 |
| TCGA-2Y-A9GU-01A | 0.09223503 |

| | |
|------------------|------------|
| TCGA-2Y-A9GV-01A | 0.17151841 |
| TCGA-2Y-A9GW-01A | 0.61519815 |

Visualize the TPAC scores in the **S** matrix as a heatmap (this is generated using similar logic as the heatmaps included in the TPAC paper [3]).

```
> library(gplots)
> my_palette = colorRampPalette(c("steelblue", "seagreen3",
+                                     "white", "orange", "orangered"))(n = 299)
> breaks = 300
> heatmap.2(t(tpac.out$S),
+             col = my_palette, dendrogram="both", na.rm=T,
+             symm=F, scale = "none", trace = "none",
+             xlab=NA, ylab=NA, labCol=NA, sepcolor="white",
+             sepwidth=c(0, .2), symkey=F,
+             Rowv=T, Colv=T,
+             breaks=breaks, margins=c(2,27),
+             key.title=NA, key.ylab=NA, key.xlab=NA,
+             key.ytickfun=function() {
+               return(list(labels=FALSE, tick=FALSE))
+             },
+             lwid=c(.5,4), lhei=c(.5,4), main = NA)
```



References

- [1] Cancer Genome Atlas Research Network, Weinstein, J.N., Collisson, E.A., Mills, G.B., Shaw, K.R.M., Ozenberger, B.A., Ellrott, K., Shmulevich, I., Sander, C., Stuart, J.M.: The cancer genome atlas pan-cancer analysis project. *Nat Genet* **45**(10), 1113–20 (2013). doi:10.1038/ng.2764
- [2] Uhlen, M., Zhang, C., Lee, S., Sjöstedt, E., Fagerberg, L., Bidkhor, G., Benfeitas, R., Arif, M., Liu, Z., Edfors, F., Sanli, K., von Feilitzen, K., Oksvold, P., Lundberg, E., Hober, S., Nilsson, P., Mattsson, J., Schwenk, J.M., Brunnström, H., Glimelius, B., Sjöblom, T., Edqvist, P.-H., Djureinovic, D., Micke, P., Lindskog, C., Mardinoglu, A., Ponten, F.: A pathology atlas of the human cancer transcriptome. *Science* **357**(6352) (2017). doi:10.1126/science.aan2507
- [3] Frost, H.R.: Tissue-adjusted pathway analysis of cancer (tpac). *bioRxiv* (2022). doi:10.1101/2022.03.17.484779. <https://www.biorxiv.org/content/early/2022/03/19/2022.03.17.484779.full.pdf>