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):

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

[1] "urothelial cancer"

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.

"cervical cancer"

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

"breast 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"
                                                    "stomach cancer"
                            "melanoma"
[16] "testis cancer"
                                                    "endometrial cancer"
                             "thyroid 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,
```

gene.set.collection=gene.set.collection)

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

cancer.type=cancer.type,

> 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	0.999999828
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	0.999999999
	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]

	HALLMARK_ADIPOGENESIS HALLM	MARK_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	0.00000e+00	0.0010408253	
TCGA-2Y-A9GW-01A	6.410074e-01	0.0089029673	
	${\tt HALLMARK_ANDROGEN_RESPONSE}$	HALLMARK_ANGIOGENESIS	
TCGA-2Y-A9GS-01A	2.649604e-01	0.852475176	
TCGA-2Y-A9GT-01A	6.135243e-06	0.99999997	
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	1.00000000	
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		

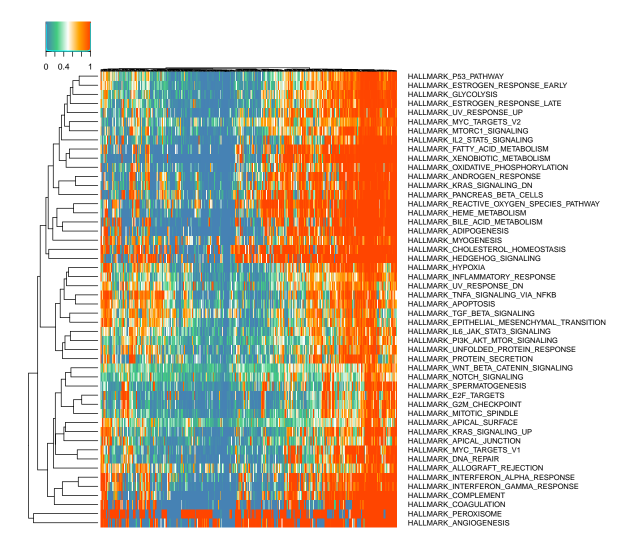
> tpac.out\$S.pos[1:5,1:5]

		HALLMARK_ADIPOGENESIS HALL	MARK_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., Bidkhori, 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