HYENA User Manual

Version 0.5.3

1 Introduction

HYENA (Hijacking of Enhancer Activity) is a computational tool to detect genes activated by nearby somatic structural variations (SVs), a phenomenon known as enhancer hijacking. It takes somatic SVs and gene expression data as inputs and outputs enhancer hijacking candidate genes. This vignette describes the components of HYENA with an example pipeline that uses Pan-Cancer Analysis of Whole Genomes (PCAWG) thyroid cancer (THCA) dataset.

2 Prerequisites and input files

2.1 Required packages

R 4.0.3 and the following list of R packages are required to run HYENA properly:

- optparse 1.6.6 or above
- data.table 1.14.2 or above
- stringr 1.4.0 or above
- stats 4.0.3 or above
- MatrixGenerics 1.2.0 or above
- ggplot2 3.3.5 or above
- RMThreshold 1.1 or above
- tidyr 1.1.2 or above
- GenomicRanges 1.42.0 or above
- IRanges 2.24.1 or above
- dplyr 1.0.7 or above
- S4Vectors 0.28.1 or above
- lattice 0.22.44 or above
- preprocessCore 1.52.1 or above
- EDASeg 2.24.0 or above

All R packages can be installed within a few minutes. For package installation issues, please contact your system admin.

2.2 Testing data set and an example pipeline

The user can run the following code to execute the entire pipeline on the THCA example:

```
cd example
sh ./example.sh
```

This will run HYENA using 0 to 5 principal components in the normal-score regression model while calculating empirical p-values for each run by permuting gene expression data 5 times. The final results for each run will be under "./example/results/" within their respective run folders (e.g. "run PCo". "run PC1", etc.). The user should be able generate "DAT.annot_sv_pur_cn_age_sex_PCo_posEstimate_adj_sig.txt" and "DAT.annot_sv_pur_ cn_age_sex_PCo_posEstimate_adj_sig_rmeqtl.txt" files. The number of significant genes and pvalues may be slightly different from the files we provide. The example pipeline can be completed in a few minutes on a normal desktop.

If the user is unable to run HYENA with all the packages properly installed, or HYENA runs smoothly but the results are dramatically different from the files we provide, it is likely due to the specific

version of the R packages the user is loading. In this case, we recommend using the package versions listed above to ensure the results are reproduced.

2.3 Overview of the pipeline

HYENA is a multi-step pipeline. STEP 1 involves creating transcription start site (TSS) windows for each gene where SV breakpoints will be mapped based on a reference genome. STEP 2 is where the breakpoint locations from each SV are mapped to these TSS windows. STEP 3 prepares expression data by normalizing them across samples, adding a small Gaussian noise, calculating normal scores, and performing principal component analysis. STEP 4 performs normal-score regression on observed regression and identifies candidate genes. STEP 5 calculates empirical p-values by permutation. Finally, STEP 6 looks for associations between known eQTLs of the candidate genes and their SV genotypes to determine whether the observed expression changes can be explained by eQTLs. **Users can choose to end the analysis afterthe normal-score regression step (STEP 4), but we highly recommend running STEP 5 for more reliable results.**

2.4 Input file format

Note that empty lines are not allowed in any input files.

There are three input files that are essential to the HYENA pipeline: SV calls, gene expression data, and sample/aliquot IDs. Individual bedpe files for each sample, a file containing expression data from all samples in a single matrix, and a file with sample IDs are the bare minimum input files. For best performance, the users should include gene level copy number, sample purity, and clinical data with patient age and sex information to be modeled as well.

The SV data should be provided as individual bedpe files and named as "sampleid.bedpe". For samples with no SVs, do not generate empty bedpe files. The contents of each file should follow the standard bedpe format as shown in Figure 1 where the first three columns are the genomic location of the first breakpoint of an SV and the following three columns are the genomic location of the second breakpoint for the same SV. The smaller coordinate should be "start1" and the larger one should be "start2". "strand1" and "strand2" indicate breakpoint orientation. For example, a deletion should have "+" and "-", whereas a tandem duplication should be "-" and "+". Note that the strand annotation may be different for different SV callers and it's important to annotation strands according to PCAWG definition (del: +-, etc.). The column order should be exactly in the indicated order. All other columns are not necessary, but the columns should be present and filled with any string.

chroml	startl	endl	chrom2	start2	end2	sv id	pe support	strandl	strand2	svclass	svmethod
1	74385662	74385663		74439877	74439878	SVMERGE2	16			h2hINV	SNOWMAN DELLY
1	74591786	74591787		74612783	74612784	SVMERGE3				DEL	SNOWMAN DELLY
10	43611774	43611775		57561502	57561503	SVMERGE5	85			DEL	SNOWMAN BRASS DELLY
10	43611833	43611834	10	61630996	61630997	SVMERGE1	48			t2tINV	SNOWMAN BRASS DELLY
10	57561150	57561151		61631185	61631186	SVMERGE4	57			h2hINV	SNOWMAN BRASS DELLY

Figure 1: SV data format (bedpe)

Note that the quality of input SVs is very important. We recommend using multiple SV callers. In our experience, many SV callers produce caller-specific artifacts. The artifacts are often small deletions, small inversions or translocations. They typically have few read support and/or overlap with repetitive sequences. We recommend using approach described in our paper (PMID: 32813322) to identify SV artifacts.

The gene expression matrix should be provided in the format depicted in Figure 2 with gene ID as row names and sample ID as column names. HYENA takes any forms of gene expression quantification as input, such as FPKM, TPM, or other normalized values. The example given in Figure 2 is FPKMs from the PCAWG study. The expression matrix can have os to indicate no expression for a particular gene. However, "NA"s or empty cells are not allowed. Gene IDs should be the same across all input and reference files and the sample IDs should match aliquot_id_rnaseq in Sample_ID file.

Figure 2: Gene expression matrix format

Sample-ID file serves two purposes for HYENA: (1) It provides the list of samples that the user would like to analyze. If the user would like to run only a subset of the samples, they can do so just by limiting their list of sample IDs in the "sample_ids.txt" file without subsetting other input files. (2) It facilitates mapping of different sample aliquot IDs from a single donor/patient. In studies where the donor and the samples collected are identified with the same unique ID, matching the corresponding RNA and DNA samples may not be an issue. However, in most com prehensive multi-omic studies such as PCAWG, International Cancer Genome Consortium (ICGC), and The Cancer Genome Atlas (TCGA), RNA and DNA samples have different aliquot numbers are different from donor IDs. Therefore, the "sample_ids.txt" file is formatted in four columns as depicted in Figure 3: "submitter_donor_id", "aliquot_id_wgs", "aliquot_id_wgs_norm" (ID for the matched normal sample, this column is required for eQTL analysis) and "aliquot_id_rnaseq". This allows HYENA to identify the correct whole genome sequencing data (used for SV calls, copy number calls etc.) and RNA sequencing data. Samples listed in the sample-ID file must be in the gene expression matrix. If all IDs are the same, the user can simply populate all columns with the same values. Note that some special characters in sample IDs may cause the algorithm to crash.

```
        submitter donor id
        aliquot id wgs
        aliquot id wgs
        aliquot id maseq

        085ffcdc-8600-42e2-952a-506465f716a2
        07a7c634-bd9a-4fc2-b9fe-87b060e3d1f
        1c6abcc3-la61-4jaa-92id-6d06f762c1a3
        0809dd6f-0f8c-4j07-bb6a-6leb10feaddd

        2501ee46-8388-48b-8765-e9c9706cbbes
        262c6a0-2341-49c4-a6dd-656e7bb505dd
        62c780b5-335-62b-bff1-6919adda895d
        48f7b6d-d38e-4jb5-864-bb66266da9jb

        2581c1bc-7c64-dch1-ala-3-ba85dafb8f4
        48c2f08-437-6991-acbs-8bs9912f913
        42f5c568-964-6c4-a7r-a-a3df7bb648-26000-2E13-4EC-B097-R2645b77ab

        2589499-8549-4da5-b391-0421f18facf1
        39d30499-8549-4da5-b391-0421f18facf1
        5bdca82-2274419-bc63-3227d3lbf704
        66101692-5616-4968-3893-66c5e87d333
        28C6743-CD3D-4693-8962-651-495-863-8967d33

        368Cc58-336-4dd5-833d-5895b898016
        0cl88b7-bad1-4aa4-a6df-472f389d8761
        9ce50075-10f8-4915-3891-66c5e87d333
        28C6743-CD3D-4693-8950-F5538146dd486

        36787a0e-b542-401e-b1e5-077376090b3a
        4303953-3941-4358-a06b-41e92f7e8d20
        97808f8d-720c-91c2-bad0-5992e8f8a
        486f511-f479-4393-864b-f15386c36e

        364d518-6427b-4911-3b10-055224886ff
        5663951-4368-8580-91c2-bad0-5992e8f8a
        486f516-6-303-3746c8-8580-20c1295e8f3a
        486f511-f479-4393-864b-f15386c36e
```

Figure 3: List of sample IDs

HYENA also incorporates copy number data. The users can feed integer copy number calls for each gene in order to mitigate the effect of differential gene expression due to copy number differences between samples. The copy number input file has similar format as expression data as shown in Figure 4. The sample IDs should be aliquo id wgs from Sample-ID file. NAs are allowed in copy number data.

```
    gene_id
    07a7c634-bd9a-4fc2-b9fe-87b060ec3dlf
    248fd0ed-fil4c-40b8-9f14-e9c7adf16e22
    64c2e6a0-2341-49c4-a6dd-656e7bb505dd
    4bae2f08

    -da75-4991-acb8-5ba9912f9131
    b8cd6882-be27-4742-bc63-3227d31bf704
    5bdca282-c671-48ff-b32b-2380996016c0

    ENSG00000002016.12
    2
    2
    2
    2

    ENSG0000000478.5
    2
    2
    2
    2

    ENSG0000000501.5
    2
    2
    2
    2

    ENSG00000006102.8
    2
    2
    2
    2

    ENSG00000006194.6
    2
    2
    2
    2

    ENSG00000006327.9
    2
    2
    2
    2

    2
    2
    2
    2
    2

    2
    2
    2
    2
    2

    2
    2
    2
    2
    2

    2
    2
    2
    2
    2

    2
    2
    2
    2
    2

    ENSG0000006192.8
    2
    2
    2
    2

    ENSG0000006192.9
    2
    2
    2
    2
```

Figure 4: Absolute copy number calls for each gene in WGS samples

In studies where sample purity is available, users can choose to correct gene expression for sample purity by providing a file with a "samplename" (WGS) column and a "purity" column (Figure 5). Purity should be values between 0 and 1.

```
samplename
                purity
0009b464-b376-4fbc-8a56-da538269a02f
                                         0.885
003819bc-c415-4e76-887c-931d60ed39e7
0040b1b6-b07a-4b6e-90ef-133523eaf412
                                         0.8
00493087-9d9d-40ca-86d5-936f1b951c93
00508f2b-36bf-44fc-b66b-97e1f3e40bfa
                                         0.92
005794f1-5a87-45b5-9811-83ddf6924568
                                         0.596
005e85a3-3571-462d-8dc9-2babfc7ace21
                                         0.46
007aab66-2f07-459d-8952-3041d6ea24a8
                                         0.581
008aef39-0c97-48ce-9dfd-f12d67116c59
                                         0.759
```

Figure 5: Sample purity information to be used in the linear model.

Finally, users can choose to incorporate clinical data. For now, HYENA only supports donor sex, age, and cancer type. If users would like to correct for these confounding factors in the regression model, they can provide a clinical data file with four columns: "submitter_donor_id", "donor_sex", "donor_age_at_diagnosis", and "cancer_type" (Figure 6). The order of these columns does not matter.

```
donor_age_at_diagnosis
submitter donor id
                                       donor sex
c7a2f394-3e3f-4c90-9f1e-f2be3e5b0d6b
                                      male
                                                                           THCA
4fc8e011-4433-4537-b03f-457a3a70240f
                                      male
                                                                           THCA
a09d0be0-fd41-4b30-b488-9a9f2abef8e7
                                      female
                                                  47
                                                                           THCA
fb83f7d7-2182-4fb7-8e3e-8ad7f1feac72 male
                                                                           THCA
6abc861a-376d-446b-acee-fb0f03a82c09 male
                                                                           THCA
c31b3adf-3fc6-4a72-83b0-bbc0f295b5e7
                                      female
                                                                           THCA
e35b2928-8784-45d0-a7la-a2df161542fa female
                                                  44
                                                                           THCA
77bb90f0-e923-4e28-bb88-4d6b420e0b5c
                                                  44
                                                                           THCA
                                      female
8d54a2ed-03f1-4a23-bdd2-f3395f5d3716
                                                  66
                                                                           THCA
                                      female
```

Figure 6: Format of the clinical meta data for each sample

3 Workflow

3.1 Step 1: Calculating TSS windows

A list of TSS-flanking windows needs to be generated using "calc_wind.R". This script will start with a desired reference genome file and create a file with four new columns that define the TSS window for each gene: (1) "tss", transcription start site; (2) "tts", transcription termination site; (3) "tss_left", the left boundary of the TSS window; (4) "tss_right", right boundary of the TSS window.

Arg	Description
-r,ref	Gene annotation
-u,up	Number of basepairs UPSTREAM of transcription start site (default: 500000)
-d,down	Number of basepairs DOWNSTREAM of transcription start site (default: 500000)
-w,write	Output file (default: "./tss_windows.txt")
-v,verbose	Verbose mode for troubleshooting (default: FALSE)

Table 1: Description of arguments for calc_wind.R

Table 1 shows the arguments available for "calc_wind.R". In the THCA example, this can be done with the following command:

```
Rscript ../R/calc_wind.R \
-r ../ref/toy_gene_annot_hg19.txt \
-w ../ref/toy_gene_annot_hg19_tsswindow.txt
```

The example pipeline uses "toy_gene_annot_hg19.txt" for demonstration purposes. Users should not use the toy annotation file other than the test run, but instead use "gene_annot_hg19.txt" and "gene_annot_hg38.txt" provided for actual analysis. Running the above code will generate a file

```
transcript id
    1 gene 11869 14412
                         + ENSG00000223972.4 ENSG00000223972.4 pseudogene
    1 gene 14363 29806
                            - ENSG00000227232.4 ENSG00000227232.4 pseudogene
    1 gene 29554 31109
                            + ENSG00000243485.2 ENSG00000243485.2
    1 gene 34554 36081
                            - ENSG00000237613.2 ENSG00000237613.2
                           + ENSG00000268020.2 ENSG00000268020.2 pseudogene
    1 gene 52473 54936
                            + ENSG00000240361.1 ENSG00000240361.1 pseudogene
    1 gene 62948 63887
                   tts tss left tss right
 gene name
           tss
   DDX11L1 11869 14412
                       -488131
                                   511869
   WASH7P 29806 14363
                        -470194
                                   529806
MIR1302-11 29554 31109
                        -470446
                                   529554
   FAM138A 36081 34554
                        -463919
                                   536081
   OR4G4P 52473 54936
                        -447527
                                   552473
   OR4G11P 62948 63887
                        -437052
                                   562948
```

named "toy_gene_annot_hg19_tsswindow.txt" (Figure 7). Notice that for genes that are close to the beginning of the chromosome, left boundary can be a negative basepair location. Similarly, for genes that are close to the end of the chromosome, the right boundary may extend beyond the length of the chromosome. This will not cause any issues moving forward.

Figure 7: TSS windows spanning 500Kb on either side of the TSS for each gene in "hg19"

The windows with respect to TSS sites can be customized. For example, if we wanted to use the hg38 reference genome and consider 400Kb upstream and 10kb downstream of TSS, we would run the command below and create "gene_annot_hg38_tsswindow.txt" (Figure 8).

```
Rscript ../R/calc_wind.R \
-r ../ref/gene_annot_hg38.txt \
-w ../ref/gene_annot_hg38_tsswindow.txt \
-u 400000 -d 10000
```

```
CHROM TYPE START
                       END STRAND
                                       transcript id
                                                                gene id
1:
       1 gene 11869 14409
                                + ENSG00000223972.5 ENSG00000223972.5
2:
         gene 14404 29570
                                  ENSG00000227232.5 ENSG00000227232.5
3:
         gene 17369 17436
                                - ENSG00000278267.1 ENSG00000278267.1
                                + ENSG00000243485.5 ENSG00000243485.5
4:
       1 gene 29554 31109
                                  ENSG00000284332.1 ENSG00000284332.1
5:
         gene 30366 30503
6:
       1 gene 34554 36081
                                  ENSG00000237613.2 ENSG00000237613.2
                             gene type
                                          gene name
                                                       tss
                                                             tts tss left
   transcribed unprocessed pseudogene
                                            DDX11L1 11869 14409
                                                                   -388131
                                             WASH7P 29570 14404
2:
               unprocessed pseudogene
                                                                     19570
3:
                                          MIR6859-1 17436 17369
                                 miRNA
                                                                      7436
4:
                                lncRNA MIR1302-2HG 29554 31109
                                                                   -370446
5:
                                 miRNA
                                          MIR1302-2 30366 30503
                                                                  -369634
6:
                                            FAM138A 36081 34554
                                                                     26081
                                 1ncRNA
   tss right
       21869
2:
      429570
3:
      417436
4:
       39554
5:
       40366
      436081
```

Figure 8: Customized TSS window

Alternatively, users can define any window they wish in any reference genome without using the "calc_wind.R" script as long as they provide a file in the same format described above.

3.2 Step 2: Mapping SVs to TSS windows

Second step of HYENA involves mapping SV breakpoints to the TSS windows created for each gene using the "mapsv.R" script. For this analysis, breakpoints are considered individually and as long as one of the two breakpoints that define an SV falls within a TSS window, that SV is considered to have the potential to influence the gene expression. However, if both of the breakpoints of an SV fall within gene body (i.e. intragenic SVs), these SVs are left out of the analysis as they are unlikely to change promoter-enhancer interactions for that gene.

Arg	Description
-r,ref	Gene annotation with TSS window
-i,id	Sample id file
-b,bedpe	Folder for bedpe files
ingene	Remove SVs located entirely in a gene (default: 1)
del	Remove small deletions (default: 0)

dels	Small deletion size cutoff (default 10000)
dup	Remove small duplications (default: 1)
dups	Small duplications size cutoff (default: 10000)
inv	Remove small inversion (default: 0)
invs	Small inversion size cutoff (default: 10000)
-w,write	Folder for intermediate files (default: "./intermediate/")
-x,prefix	Prefix for output files (default: "DAT")
-v,verbose	Verbose mode for troubleshooting (default: FALSE)

Table 2: Description of arguments for mapsv.R

Small tandem duplications typically will not lead to new promoter-enhancer interactions. By default, they will be filtered out. However, small deletions may delete TAD boundaries or other repressive elements and new promoter-enhancer interactions can form. So small deletions are not filtered by default. However, certain tumor types have excessive number of small deletions, such as breast cancer. In this case, filter out small deletions may be beneficial.

To execute Step 2 of HYENA for THCA example, run:

```
Rscript ../R/mapsv.R \
-r ../ref/toy_gene_annot_hg19_tsswindow.txt \
-i ./data/sample_ids.txt \
-b ./data/bedpe/ \
-w ./intermediate/
```

By default, SVs located entirely in gene body are removed. Small tandem duplications are also removed, but not small deletions or inversion.

The file "sv_mapped_filtered_numSV.txt" with gene level SV genotype will be used for future steps. The file format is gene by sample ID (Figure 9). If a sample has a breakpoint within the TSS window of a gene, the sample is annotated as "sv" for that particular gene. Otherwise, the sample receives a "no_sv" status. The last two columns titled "num.sv" and "num.nosv" summarize the number of samples that have or do not have SVs, respectively, for a particular gene.

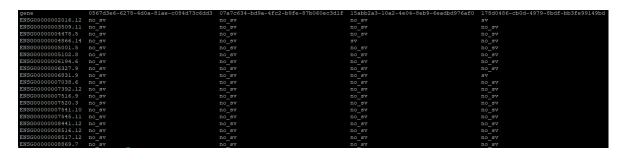


Figure 9: "sv_mapped_filtered.txt" output file

When the sample size is large or the number of SVs is large, this step can be very slow. In this case, users can split the gene annotation file into smaller chunks (e.g. 1000 genes per file), run "mapsv.R" to generate multiple "sv_mapped_filtered.txt" files, and merge them into one file.

3.3 Step 3: Calculating normal scores for gene expression

If gene expression data are not normalized, they will be quantile normalized using the "quantNorm.R" script. If they are already normalized, this step can be skipped.

Arg	Description
-е,exp	Gene expression input
-w,write	Folder for intermediate files (default: "./intermediate/")
-x,prefix	Prefix for output files (default: "DAT")
-i,id	Sample id file

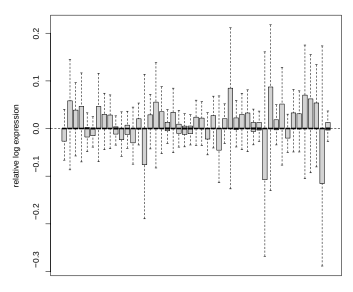
Table 3: Description of arguments for quantNorm.R and add_noise.R

Example use of quantNorm.R:

```
Rscript ../R/quantNorm.R \
-e ./data/exp.txt \
-w ./intermediate/ \
-w ./data/sample ids.txt
```

This step generates a gene by sample ID matrix called "*.exp_quant.txt" and two relative log expression (RLE) plots showing the distribution of FPKM values before and after quantile normalization (Figure 10). These plots can be used to identify problematic or outlier samples that can skew downstream analysis.

RLE before quantile normalization



RLE after quantile normalization

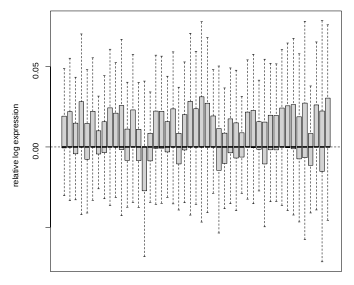


Figure 10: Relative log expression across all samples before and after quantile normalization

To break ties of gene expression values during ranking, a very small Gaussian noise is added to the

quantile-normalized expression matrix using "add_noise.R". The input gene expression file is the output file of "quantNorm.R". Gene expression normal scores are calculated. Then principal component analysis (PCA) is performed on gene expression scores. For THCA example, run:

```
Rscript ../R/add_noise.R \
-e ./intermediate/DAT.exp_quant.txt \
-w ./intermediate/
```

An R object file "*.exp.Rdata" will be generated and is the input gene expression file for "HYENA.R" and "pmt_exp.R".

3.4 Step 4: Normal-score regression

"HYENA.R" tests gene expression associated with SV status by normal-score regression while correcting for the confounding factors such as gene copy number, sample purity, donor age and sex, as well as unknown factors captured by the first n principal components (PCs) of gene expression. Table 4 describes the options for "HYENA.R".

Expression norma score $\sim sv$ status + CN + purity + age + sex + PC_1 + PC_2 + + PC_2	Expression 1	norma	$score \sim sv$	status + CN +	puritu + aae	e + sex + PC	$1 + PC_2 +$	$ + PC_n$
--	--------------	-------	-----------------	---------------	--------------	--------------	--------------	-----------

Arg	Description
-e,exp	Expression Rdata file generated by add_noise.R or pmt_exp.R
-s,sv	*sv_mapped_filtered_numSV.txt file generated by mapsv.R
-i,id	Sample id file
-a,annot	Gene annotation with TSS window
-p,purity	Sample purity file
-c,cna	Gene level copy number file
-m,clinical	Clinical file
-n,npc	Number of first n PCs to be included in the model (default: 0)
-d,dir	Output folder (default: "./results/")
-w,write	Folder for intermediate files (default: "./intermediate/)
-x,prefix	Prefix for output files (default: "DAT")
-v,verbose	Verbose mode for troubleshooting (default: FALSE)
-t,pmt	Run in permutation mode (default: FALSE)
pur	Include purity in regression (default: FALSE)
cn	Include gene copy number in regression (default: FALSE)
age	Include donor age in regression (default: FALSE)
sex	Include donor sex in regression (default: FALSE)
type	Include cancer type in regression for pancancer analysis (default: FALSE)
PC	Include principal components in regression (default: FALSE)
-f,fcutoff	SV Frequency (%) cutoff (default: 5)
-C,cncutoff	Maximum copy number cutoff (default: 10)
-g,genes	A list of genes of interest to be tested (default: FALSE)

Table 4: Description of arguments for HYENA.R

While selecting the first n PCs, HYENA tests the correlation between each PC and sv_status to make sure that those PCs that are highly correlated with sv_status for a particular gene are not included in the regression model. For example, if we wanted to include 5 PCs in the model for Gene A but the 3rd and the 5th PCs are correlated with sv_status, HYENA would include PCs 1,2,4,6, and 7 in the model. We recommend testing number of PCs up to 10% of the sample size for small cohort. For example, in THCA example, we test 0 to 5 PCs. For large cohort, we recommend testing up to 20 PCs, but not more than that.

```
Exp\_normal\_score_A \sim sv\_status_A + CN_A + purity + age + sex + PC_1 + PC_2 + PC_4 + PC_6 + PC_7
```

The users can run the full model with o principal component with the following command:

```
Rscript ../R/HYENA.R \
```

```
-e ./intermediate/DAT.exp.Rdata \
-s ./intermediate/DAT.sv_mapped_filtered_numSV.txt \
-i ./data/sample_ids.txt
-a ../ref/toy_gene_annot_hg19_tsswindow.txt
-p ./data/purity.txt
-c ./data/cna.txt
-m ./data/clindat.txt
-d ./results/run_PCO/
-w ./intermediate/
--pur --cn -C 10 --age --sex -f 5 --PC -n 0
```

This will generate multiple output files in the "./results/run_PCo/" directory. The most important output file is "*annot_* posEstimate.txt" with genes ranked by their false discovery rates (FDRs) (Figure 11).

```
__
Estimate
                                                    pvalue p.onesided
NSG00000136231.9
                                                          23510086
                                                                                                                                         23349828
                                                  1.6911886490216 0.35349983171933
                                                                                                                                 2.23986164867718e-05
93082433859e-05
                           23405237
23905237
90533
                    23905237 7/40
1.84690538842761e-05 0.0
                                                 7/40 14.9 1.63012
0.000304739389090555
                                                                 1.63012363652897
                                                                                                                                4.62671505504283
                                                                                                                                                             3.6938
                       7 23436065 23436135
7/40 14.9 1.47462973468051
-05 0.000821013025819107
10 43572475
  7685521e-05
                                                                                       snoRNA SNORD65 23436065
                                                                                                                                                   22936065
   23936065
7.4637547801737e-05
                                                                                                                                         0.000149275095603474
                             0 43572475 43625799
6/41 12.8 1.74085172085163
0.000844297454752983
                                                                               0.427023207063186
0.000102339085424604
NSG00000252590.1 7
                             23490277
7/40 14 5
                                                                                       snRNA RNU7-143P
                                                 1.16113909951023
    0.00202475613306818
                                                                   + pseudogene
1.22614652891757 0.4
                                                                                                           AC079780.3
NSG00000236654.2
                    0.00259791519548649
```

Figure 11: HYENA output

HYENA has a default cutoff of 10 for the gene's copy number. This is because genes with 10 or more copies are likely amplified in a circular extrachromosomal DNA (also known as double minutes). By default, HYENA removes these cases from downstream analysis. However, if the user would like to include these cases, they can set the "-C" argument to a very high number (e.g. 1000).

If the user has copy ratio data instead of integer copy numbers for genes, they can stilluse the "-c,--can" argument with the full path to their copy ratio file. But the format of their file shouldstill match the CN file. Moreover, if copy ratio is used, we recommend that the user removes "purity" from the regression model as the sample purity information is captured by the copy ratio data already. For copy ratio, we recommend using a cutoff of 3.

```
Rscript ../R/HYENA.R \
-e ./intermediate/DAT.exp.Rdata \
-s ./intermediate/DAT.sv_mapped_filtered_numSV.txt \
-i ./data/sample_ids.txt
-a ../ref/toy_gene_annot_hg19_tsswindow.txt
-c ./data/cn_ratio.txt \
-m ./data/clindat.txt
-d ./results/
-w ./intermediate/
--cn -C 3 --age --sex -f 5 --PC -n 0
```

Users can also automate testing different numbers of PCs to the regression model with the following code, which runs HYENA with 0 to 5 PCs.

```
for i in {0..5}
   do
   Rscript ../R/HYENA.R \
```

```
-e ./intermediate/DAT.exp.Rdata \
-s ./intermediate/DAT.sv_mapped_filtered_numSV.txt \
-i ./data/sample_ids.txt
-a ../ref/toy_gene_annot_hg19_tsswindow.txt
-p ./data/purity.txt
-c ./data/purity.txt
-m ./data/clindat.txt
-m ./data/clindat.txt
-d ./results/run_PC${i}/
-w ./intermediate/run_PC${i}/
-pur --cn -C 10 --age --sex -f 5 --PC -n ${i}
done
```

If users are running pancancer analyses and want to correct for cancer type specific effects, HYENA has the "--type" argument to include cancer type as a random variable in the normal-score regression. In this case, users should provide an extra column of "cancer_type" in the clinical data file. The following code is an example:

```
Rscript ../R/HYENA.R \
-e ./intermediate/DAT.exp.Rdata \
-s ./intermediate/DAT.sv_mapped_filtered_numSV.txt \
-i ./data/sample_ids.txt
-a ../ref/toy_gene_annot_hg19_tsswindow.txt
-p ./data/purity.txt
-c ./data/cna.txt
-m ./data/clindat.txt
-d ./results/
-w ./intermediate/
--pur --cn -C 10 --age --sex --type -f 5 --PC -n 0
```

3.5 Step 5: Calculating empirical p-values

This step is strongly recommended since p-values from the previous step are often inflated. Users can generate a null distribution by randomizing gene expression data using the "pmt_exp.R" script. Empirical p-values can be generated by repeating this permutation process multiple times. To calculate empirical p-values at the 10⁻⁶ level, the user needs to generate at least 1 million permuted p-values. This can be achieved by pooling p-values from multiple permutations. Note that every run of HYENA will generate approximately the number of p-values that is equal to the number of genes that have SVs satisfying the recurrence cutoff in the data set. For example, if a data set has 1,000 genes that have SVs in at least 5% of the tumors, then the user needs to run 1,000 permutations to reach final p-value count of 1 million. We recommend running 50 to 100 permutations for each dataset. Normal-score regression on permuted data should be run in parallel.

"pmt_exp.R" takes "*.exp.Rdata" generated by "add_noise.R" as input and shuffles sample ID at random. The output is a "*exp.Rdata" file to be used as input for "HYENA.R". If HYENA.R is being executed on permuted expression data, "--pmt" argument should be turned on.

Arg	Description
	Expression Rdata file generated by add_noise.R
-w,write	Folder for intermediate files (default: "./intermediate/exp_pmt/")
-x,prefix	Prefix for output files (default: "PMT")

Table 5: Description of arguments for pmt_exp.R

```
Rscript ../R/pmt_exp.R \
-e ./intermediate/DAT.exp.Rdata \
-w ./intermediate/exp_pmt/ \
-x PMT

Rscript ../R/HYENA.R \
```

```
-e ./intermediate/exp_PMT/PMT.exp.Rdata \
-s ./intermediate/DAT.sv_mapped_filtered_numSV.txt \
-i ./data/sample_ids.txt
-a ../ref/toy_gene_annot_hg19_tsswindow.txt
-c ./data/cna.txt \
-m ./data/clindat.txt
-d ./results/
-w ./intermediate/
--cn -C 10 --age --sex -f 5 --PC -n 0 --pmt
```

For demonstration purpose in the example pipeline, we will perform permutation only 5 times on each PC to calculate empirical p-values. The following code can automate this process in the command line:

```
# Permute expression data
for j in \{1...5\}
   do
   Rscript ../R/pmt exp.R \
   -e ./intermediate/DAT.exp.Rdata \
   -w ./intermediate/exp pmt/ \
   -x PMT${j}
   done
# Run HYENA on permuted expression
for i in \{0...5\}
   do
   for j in \{1...5\}
      Rscript ../R/HYENA.R \
      -e ./intermediate/exp pmt/PMT${j}.exp.Rdata \
      -s ./intermediate/DAT.sv mapped filtered numSV.txt \
      -i ./data/sample ids.txt \
      -a ../ref/toy gene annot hg19 tsswindow.txt \
      -p ./data/purity.txt \
      -c ./data/cna.txt \
      -m ./data/clindat.txt \
      -w ./intermediate/run PC${i}/ \
      -d ./intermediate/run PC${i}/ \
      -x PMT${j} \
      -pur -cn -C 10 -age -sex -f 5 -PC -n ${i} --pmt
      done
   done
```

This will generate 5 permuted result files per PC run in the "./intermediate/" directory. Then the permuted p-values from these runs can be extracted and concatenated into a single file using the following command:

```
ls -d intermediate/run_PC* | while read DIR
   do
   [ -e $DIR/pvalpmt.txt ] && rm $DIR/pvalpmt.txt
   touch $DIR/pvalpmt.txt
   tail -q -n +2 $DIR/*_posEstimate.txt >> $DIR/pvalpmt.txt
   cp $DIR/pvalpmt.txt $DIR/temp.txt
   cut -f17 $DIR/temp.txt > $DIR/pvalpmt.txt
   rm $DIR/temp.txt
   done
```

Then the "pvalpmt.txt" file can be used as the null distribution to calculate empirical p-values by the "empiricalp.R" script which outputs a file with the "_adj.txt" suffix. Here, we use two-sided p

values to build null distribution and calculate two-sided empirical p values, then derive one-sided empirical p values and perform FDR correction.

Arg	Description
-o,obs	HYENA.R output file using observed gene expression
-p,pmt	P value null distribution file (pvalpmt.txt)
-d,dir	Output folder (default: "./results/")
qq	Draw a QQ plot (default: FALSE)

Table 6: Description of arguments for empiricalp.R

Example use of empiricalp.R:

```
Rscript ../R/empiricalp.R \
-o ./results/*_posEstimate.txt \
-p ./intermediate/pvalpmt.txt \
-d ./results/
```

To calculate empirical p-values for each PC run:

```
for i in {0..5}
   do
   Rscript ../R/empiricalp.R \
   -o ./results/run_PC${i}/*_posEstimate.txt \
   -r ./intermediate/run_PC${i}/pvalpmt.txt \
   -d ./results/run_PC${i}/
   done
```

Users can also choose to generate a Quantile-Quantile plot (Q-Q plot) of p-values by adding the "-qq" argument. Note that the Q-Q plot argument requires all p-values to be non-zero. If any of the empirical values are 0, the Q-Q plot will not be drawn, and an error message will appear. However, the empirical p-value results will still be output appropriately.

After running multiple models with different numbers of PCs, users can compare the results by "setpc.R" to identify the number of significant hits achieved by each model. A final model that identifies at least 80% of the maximum number of hits across all models with the smallest number of PCs will be picked.

Arg	Description
-l,filelist	Comma separated list of result files
-p,power	Desired power (default: 0.8)
-f,fdrcutoff	FDR cut-off for significant genes (default: 0.1)
-w,write	Folder for intermediate files (default: "./intermediate/")
emp	P values to process are empirical p-values (p.emp.onesided)
-d,dir	Output folder (default: "./results/")
-x,prefix	Prefix for output files (default: "DAT")

Table 7: Description of arguments for setpc.R

It is crucial that the users list their results files from different models in an INCREASING order of parameters used in the model. For example, a model that uses "sv + purity + cn + age" has fewer parameters than "sv + purity + cn + age + 3 PCs". Similarly, "sv + purity + cn + age + 3 PCs" has fewer parameters than "sv + purity + cn + age + 10 PCs". This allows the script to pick the model that reaches 80% power with the fewest number of parameters to avoid over-fitting. "setpc.R" generates a result file with the extension "_sig.txt" that only has the significant hits. It will also generate a file called "Prefix.pc_tally.txt" with the number of significant genes identified in each model listed in the order provided by the user (Figure 12). An example command to run "setpc.R" would be:

```
Rscript ../R/setpc.R \
-l ./results/run_PC0/DAT.annot_sv+pur+cn+age+sex+PC0_posEstimate_adj.txt,
./results/run PC1/DAT.annot sv pur cn age sex PC1 posEstimate adj.txt,
```

```
./results/run_PC2/DAT.annot_sv_pur_cn_age_sex_PC2_posEstimate_adj.txt,
./results/run_PC3/DAT.annot_sv_pur_cn_age_sex_PC3_posEstimate_adj.txt,
./results/run_PC4/DAT.annot_sv_pur_cn_age_sex_PC4_posEstimate_adj.txt,
./results/run_PC5/DAT.annot_sv_pur_cn_age_sex_PC5_posEstimate_adj.txt \
-w ./intermediate/ \
-d ./results/\
--emp

1 ./results/run_PC0/DAT.annot_sv_pur_cn_age_sex_PC0_posEstimate_adj.txt 10
2 ./results/run_PC1/DAT.annot_sv_pur_cn_age_sex_PC1_posEstimate_adj.txt 9
3 ./results/run_PC2/DAT.annot_sv_pur_cn_age_sex_PC2_posEstimate_adj.txt 7
4 ./results/run_PC3/DAT.annot_sv_pur_cn_age_sex_PC3_posEstimate_adj.txt 7
```

Figure 12: Example "pc_tally.txt" output

./results/run_PC4/DAT.annot_sv_pur_cn_age_sex_PC4_posEstimate_adj.txt ./results/run_PC5/DAT.annot_sv_pur_cn_age_sex_PC5_posEstimate_adj.txt

3.6 Step 6: Testing associations between SV status and known eQTLs

This step requires germline single nucleotide variations (SNVs) and is optional. Based on our experience, about 10% of the candidate genes whose expression can be partially explained by known eQTLs. "eqtl.R" can test associations between SV status and known eQTLs and remove genes with significant associations.

Arg	Description
-Q,eqtl	Tissue-specific significant gene-eQTL pairs
-S,snv	Germline SNV genotypes for all samples
-R,results	HYENA results file '_posEstimate.txt' or '_adj.txt' or'_adj_sig.txt'
emp	P values are empirical p-values ('p.emp.fdr'),
-f,fdrcutoff	FDR cutoff for genes (default: 0.1)
-c,sigcutoff	P-value cutoff for eQTLs (default: 0.05)
-g,genome	Reference genome version ('hg19' or 'hg38')
-x,prefix	Prefix for output files (default: "DAT")
-V,svstat	*sv_mapped_filtered_numSV.txt generated by mapsv.R in intermediate folder
-i,id	Sample id file
-a,annot	Gene annotation with TSS window
-w,write	Folder for intermediate files (default "./intermediate/ ")
-d,dir	Output folder (default "./results/")
-v,verbose	Verbose mode for troubleshooting (default: FALSE)

Table 8: Description of arguments for eqtl.R

"eqtl.R" script requires two SNV and eQTL input files: (1) germline SNV genotypes called from normal samples and (2) known eQTL-gene pairs for the relevant tissue. eQTL-gene pairs can be downloaded from the Genotype-Tissue Expression (GTEx) data portal (http://www.gtexportal.org). Each row should indicate one unique eQTL with an ID in the format of "CHROM_POS_REF_ALT_version" (Figure 13. The "variant_id" should end with "_b38" if hg38 is used as the reference genome, and "_b37" if hg19 is used. GTEx portal provides liftover references to convert variant ids between hg38 and hg19. If the user provides two columns for variant id (one for hg19 id and one for hg38 id in the GTEx reference file as in Figure 13), "eqtl.R" will automatically select the appropriate id column based on the reference version indicated in the "--genome" argument.

variant_id_b38 gene_id	l tss_distance ma_samp	les ma_cour	nt	maf pva	l_nominal slope	slope_se	pval_nominal_th	reshold min_pva	l_nominal
pval_beta	variant_id_b37								
10_42142211_G_T_b38	ENSG00000169826.7	-996275 28	28	0.0243902	6.95649e-05	0.446382	0.111284	0.000374254	2.91374e
-115 1.27025e-100	10_42637659_G_T_b37								
10_42173399_A_G_b38	ENSG00000169826.7	-965087 28	28	0.0243902	6.95649e-05	0.446382	0.111284	0.000374254	2.91374e
-115 1.27025e-100	10_42668847_A_G_b37								
10_42205499_G_C_b38	ENSG00000169826.7	-932987 27		0.0235192	9.50302e-05	0.446609	0.113508	0.000374254	2.91374e
-115 1.27025e-100	10_42700947_G_C_b37								
10_42225864_ATAT_A_b38	ENSG00000169826.7	-912622 28		0.0243902	6.95649e-05	0.446382	0.111284	0.000374254	2.91374e
-115 1.27025e-100	10_42721312_ATAT_A_b37								
10_42242034_C_T_b38	ENSG00000169826.7	-896452 12		0.0104712	3.78534e-05	0.698318	0.167977	0.000374254	2.91374e
-115 1 27025e-100	10 42737482 C T h37								

Figure 13: "Thyroid.signifpairs.goi.txt" GTEx eQTL-gene pair data format

SNV data for the normal samples within the data set should be compiled in the format depicted in Figure 14. The contents of the "variant_id" column in the SNV file should match the formatting and reference genome version used in the variant id column of the GTEx file.

```
CHROM POS ID REF ALT QUAL FILTER INFO FORMAT 0569a3fc-932d-4acc-8c2d-3bf9999c3138 06dfc16a-ca37-46ca-8abb-20c1295e6f3a 180bbc11-bde0-4489-ad9d-ec66b632239b 1c4c0193-231e-4559-b362-9410bbdf16f7 1c6abcc3-1a61-441a-921d-6d06f762c133 1f1e18738-felc-4684-b3a5-2985d577536e 2125f0b1-cc6b9-4b56-ad6b-9fd4422e7c7e 22998cc2-01f2-4e03-a2a5-a4c6f9f836ce 23aa8b80-6356-479f5-baee-d06aa7d895dc 3537e160-d1b4-d0be-9152-2db2d8ac8a73 385b324c 48b6e8f8-3784-4f66-892f-5c294e06c306 60ecca33-9f6a-4c77-a862-6c3d723029c9 62c780b8-c3a9-4c61-albc-18a80c5ca48d 6f8f241c-6488-42c0-6660-660b8580563a 7acb93cd-79b5-4b8b-18b80-9ea170d402c5 8f9a995-f558f-162-1472-b561-61948d8895 8b4e9b4-c233-43b8-b7a1-34642e565553 9735-990c-7221-4442-af55-1004bdf61842 97808f8d-720c-41c2-ba0d-59922e48a182 9d555bf-01c2-4422-b5c0-76b0146bbff9 9dcf95bf-fb65-4589-90cb-302da798f00b 9ee50075 ad765-9bb4-a7c25e4f6245 b993ae95-590a-427b-a9c4-5b10-4af5-9a0-40d9-b007-24c54f34ea21 ae5bd94-c5b3-4d12-9b5f-00ad13bdd3a2 af0cf95a-72510-4af5-9bb4-a7c25e4f6245 b993ae95-590a-427b-a9c4-5b105437cf8 bb45ac91-59b6-6a-4a-604-a367-a71765db4 d2657b4b-0306-433d-a2f5-0bd9c7d4e6ba d2f5c569-9664-4cb4-a77e-a3df7bfb85a d60fc127-2b47-4cel-a120-f43adc2ae776 d6ea05f6-20df49ea05f6-20df49ea05f6-20df49ea05f69a-60df4-d0c0-a141-7667a1765db4 d2657b4b-0300-4e16-867e-ca5f8ea0355 e60fd-a0ce-94f3-b918a84c1c2 ea378b05-b95e-460d-b026-5a3b783795dc ea509942 e6
```

Figure 14: "snv.txt" input file format

Running the following command will filter genes based on eQTL and SV status association. If a gene has at least one eQTL that correlates with the SV genotype of the gene, that gene is removed. A candidate gene list after filtering will be generated with the extension "_sig_rmeqtl.txt". Note that if there are multiple "_posEstimate_adj_sig.txt" files in the "./results/" folder, an error will occur when running the command below. There should be only one such file.

```
Rscript ../R/eqtl.R \
-Q ./ref/Thyroid.signifpairs.goi.txt \
-S ./data/snv.txt \
-R ./results/*_posEstimate_adj_sig.txt \
-V ./intermediate/run_PCO/DAT.sv_mapped_filtered_numSV.txt \
-i ./data/sample_ids.txt \
-a ../ref/toy_gene_annot_hg19_tsswindow.txt \
-w ./intermediate/ \
-d ./results/ \
--emp
```

4 Results

The file with candidate gene list depends on how many steps users execute. If HYENA is executed without empirical p-value will labeled calculations, the output file be <Prefix>.<Parameters>_posEstimate.txt in the output folder. If empirical p-values are calculated, the output file will have the "_adj.txt" suffix. If multiple PCs are tested and "setpc.R" script is used to select regression model, then the output will have the "_sig.txt" suffix and contain only the significant genes from the selected model. Note that "setpc.R" can be used on both regression p-values and empirical pvalues. If "eqtl.R" is used, the final output will have the "_rmeqtl.txt" suffix. The output file for THCA example is "DAT.annot_sv_pur_cn_age_sex_PCo_posEstimate_adj_sig_rmeqtl.txt" shown in Figure 15. Users may generate a file with different number of PC being selected.

gene id CHROM	START	END	STRAND Q	gene typ	e g	ene nar	ne	tss	tts	tss left	tss	right	Ratio E	freq
Estimate	StdF	rror		pval	ue p.one	sided	fo	dr.onesid	ed p.e	mp p.en	mp.onesided	p.emp.fdr		
ENSG00000136231.	.9		23349828		23510086			protei	n coding	IGF2BP3	23510086	2334982	8 2	230100
86 240100	086	7/40	14.9	1.6911	886490216	0.353	4998317	71933	4.784	128583021	112 2	.23986164867	718e-05	1.11
993082433859e-05	5 0.00	0304739	389090555											
ENSG00000232627.	.1		23405237		23405595			pseudo	gene	AC021876	5.4 2340	05237	23405595	
22905237		2390523	7	7/40	14.9 1	.63012	3636528	397	0.35232	851324013	31 4.63	267150550428		3.6938
1077685521e-05	1.8469	0538842	761e-05	0.0003	047393890	90555								
ENSG00000212264.	.1		23436065		23436135			snoRNA	SNORD65	23436065	234	36135	22936065	
23936065	5	7/40	14.9	1.474629	73468051		0.3528	308589715	412	4.179687	773342507	0.00014	9275095603	3474
7.4637547801	1737e-05	0.0	0082101302	25819107										
ENSG00000165731.	.13	10	43572475		43625799			protei	n_coding	RET	43572475	4362579	9 4	130724
75 440724	475	6/41	12.8	1.7408	517208516		0.42	270232070	63186	4.0767	71454866396	0.000	2046781708	349208
0.0001023390	085424604	0.0	0084429745	54752983										
ENSG00000252590.	.1		23490277		23490341			snRNA	RNU7-14	3P	23490277	2349034	1 2	229902
77 239902	277	7/40	14.9	1.1611	390995102		0.38	312725563	95076	3.0454	13057200019	0.004	0495122661	13636
0.002024756	613306818		0133633904	17825	0.00	862068	9655172	241 0	.00431034	482758621	0.02370	068965517241		

Figure 15: Final output file for THCA example

If the sample size is large (e.g. more than 1000), HYENA can be slow. It is recommended to split the reference gene annotation file into smaller chunks (e.g. 1000 gene per file). Each subset of genes can be run in parallel. After running HYENA.R, the result files can be combined.

5 Plotting

Users can plot gene expression for the candidate genes. The expression plots are grouped based on the SV genotype ("sv" vs. "no_sv"). If cancer type was a part of the regression model, the plot can be grouped by cancer type as well. For each gene three different expression plots will be generated: (1) Quantile normalized FPKM (fpkm-qn), (2) expression normal scores (fpkm-qn-ns), and (3) Relativelog expression (fpkm-qn-rle).

Arg	Description			
-R,results	HYENA results file with the significant genes '_sig.txt'			
type	Group samples by cancer type (default: FALSE)			
-d,dir	Folder for gene expression matrices generated by HYENA.R			
	(default: " ./intermediate/run_PCo/matrix/")			
-w,write	Output folder (default: " ./results/plots/")			

Table 9: Description of arguments for expr_plotter.R

To generate expression plots for the example data set, run the following code:

```
Rscript ../R/expr_plotter.R \
-R ./DAT.annot_sv_pur_cn_age_sex_PC0_posEstimate_adj_sig_rmeqtl.txt \
-d ./intermediate/run_PC0/matrix/ \
-w ./results/plots/
```

"expr_plotter.R" generates a file in the output folder with "_fc.txt" suffix containing expression fold change (fpkm-qn) for each gene. The fold changes provided here should be used by caution. The significant genes are nominated by normal score regression. The estimate from normal-score regression reflects changes in ranks. So, we provide fold changes in this step calculated by linear regression.

6 Reference and contact

If you have used HYENA package, please cite our publication: TBD.

If you are able to run the THCA example, but unable to run your own data, most likely it is an input file formatting problem. You may double check **gene ID matching**, **chromosome name matching** ("chr1" and "1"), **sample ID matching**, **special characters in sample IDs**, **field separators** ("tab"), **NA** or **missing values** in gene expression matrix, **non-numeric values** for gene expression and copy number, **truncated files**, **mis-spelled column names**, **DOS/Mac new line characters**, **empty lines** in input files, etc.

If you have questions using HYENA, please contact Ali Yesilkanal (aeyesilkanal@gmail.com) and Lixing Yang (lixingyang@uchicago.edu).