rnaseq_figure_plotter (by Atsumi Ando, emal; a.ando@utexas.edu)

```
Usage; Rscript rnaseq_figure_plotter.r -i input_file -t bar -o output_file -g gene_list_file ... -c 5 -s 6
```

```
parameter of rnaseq_figure_plotter
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HELP -h, --help show this help message and exit
```

-o, --output

required function

OUTPUT

sample1 sample2

```
INPUT -i, --input input file name

TYPE -t, --type choose plot types (bar, box, density, density fill, dot color, dot shape, heatmap, histogram, line, scatter, or violin)
```

default output; output file name

general optional function

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GENE
                                          file name of specific gene ID list; generate "output" gene selection.txt file
                 -g, --gene
                                  default 0; calculate log value (log2; 2, log10; 10, loge; e)
LOG2
                 -l, --log
LOG2 NUMBER -lgn, --log number
                                                   default 0.00000001; add number to avoid -inf for log value
ZSCORE
                                          default off; apply Z-score transformation in gene (on or off). --log function should be 0 to apply --zscore function.
                 -zs, --zscore
                                          default samples; choose x-axis (gene or sample)
XAXIS
                 -x, --xaxis
                                          default gene; choose fill, color, or shape (gene or sample)
ZAXIS
                 -z, --zaxis
COLOR
                                          default 1; choose color type (1-10)
                 -c, --color
CUSTOM_COLOR
                                                   default None; customize color scales. Split colors by space. Example; red white blue green yellow
                         -cst, -custom color
LETTER SIZE
                 -ls, --letter size default 8 10; type text and title size of legend and axis, respectively. Split two number by space. Example; 20 24
FIGURE SAVE FORMAT -f, --figure save format
                                                           default pdf; choose format of figures (eps, ps, tex (pictex), pdf, jpeg, tiff, png, bmp, svg)
PLOT SIZE
                 -p, -plot size
                                          default 7 7; type width and height of figure. Split two number by space. Example; 10 12
```

optional parameter for individual plot types

```
STYLE
                 -s, --style
                                          default 4; choose backgroud of figures (1-7). This function is for every plots excepts heatmap.
                -lim, --limit
                                          default None; apply individual scale of "data". This function is for every plots excepts heatmap. Split two numbers(e.g.
LIMIT
limit 0 to 200 -> type 0 200) by space. Negative number required double quotation marks such as "negative number". Example; 0 100/"-1" 3
AXIS CHANGE -a, --axis changedefault off; flip axis in figures (on or off). This function is for every plots excepts heatmap.
                                                  default right; choose legend position of figures (none, left, right, bottom, top, or two-element numeric vector).
LEGEND POSITION
                         -lp, --legend position
This function is for every plots excepts heatmap and scatter.
GEOM POSITION
                                                  default 1; choose plot visualize types (geom position) from 1-4 in bar, density, and histogram
                         -gp, --geom position
                                                  default on on; apply column and row cluster function for heatmap (on or off). Column is first and row is
CLUSTER SELECT
                         -cs, --cluster_select
second, split two factor(on or off) by space. Example; on off
SCATTER SELECT
                         -ss, --scatter select
                                                  default None; type column of two samples for comparison in dot plot. Split samples by space. Example;
```

Input Data

-i, --input input_file

Input data should be samples for column and gene ID for row.

sample1 sample2 sample3 sample4 sample5

geneA 1 3 5.5 7 2 geneB 100 267 55 79 62

geneC 0.3 0.65 9.5 0.87 2.1

geneD 205 356 78 67 2900

geneE 1001 3001 5500 7001 2001

geneF 2 2 2 2 2

geneG 0.01 0.03 0.5 0.07 0.02

Optional; if you need to extract specific genes, provide -g, --gene gene_id_list_file

-g, --gene

gene id list file

Gene ID should be in first row and split by \n.

geneA

geneF

geneG

-g, --gene function provide "output"_gene_selection.txt file as followings.

"sample1" "sample2" "sample3" "sample4" "sample5"

"geneA" 1 3 5.5 7 2

"geneF" 2 2 2 2 2

"geneG" 0.010.030.5 0.070.02

Dataframe generation

Optional; add log transformation (log2, log10, loge) or z-score calculation. To use (-zs, -zscore) function, (-l, --log) function requires to be off (0).

<u>Parameter</u>	Setting	<u>Description</u>
-l,log	2, 10, or e	return log2, log10, or loge
-zs,zscore	on	return z-score transformation

Once software finishes it provide dataframe of three columns, gene, data, and sample. Sample, data, and gene refer to sample name, gene expression value, and gene ID, respectively. Example of dataframe is following;

sample	data	gene
sample1	1.0000	geneA
sample1	100.0000	geneB
sample1	0.3000	geneC
sample1	205.0000	geneD
sample1	1001.0000	geneE
sample1	2.0000	geneF
sample1	0.0100	geneG

Once is successfully generate data frame it will show

[&]quot;Complete dataframe generation! Dataframe generation time (seconds); time"

Axis selection

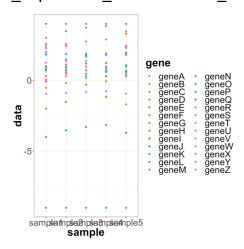
Default of x-axis and z-axis is sample and gene. You can change both axis by followings.

- -x gene
- -z sample

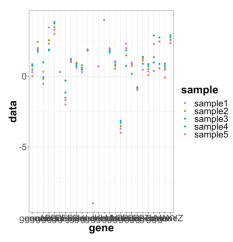
You can modify x and z in the following table.

plots	x-axis	y-axis	color/shape
bar	X	data	Z
box	Χ	data	X
density	data	density	X
dot_color	X	data	Z
dot_shape	X	data	Z
heatmap	sample	gene	
histogram	data	count	X
line	X	data	Z
scatter			
violin	x data	9	х

Rscript rnaseq_figure_plotter.r
-i gene_expression_data.txt -t dot_color -l 10

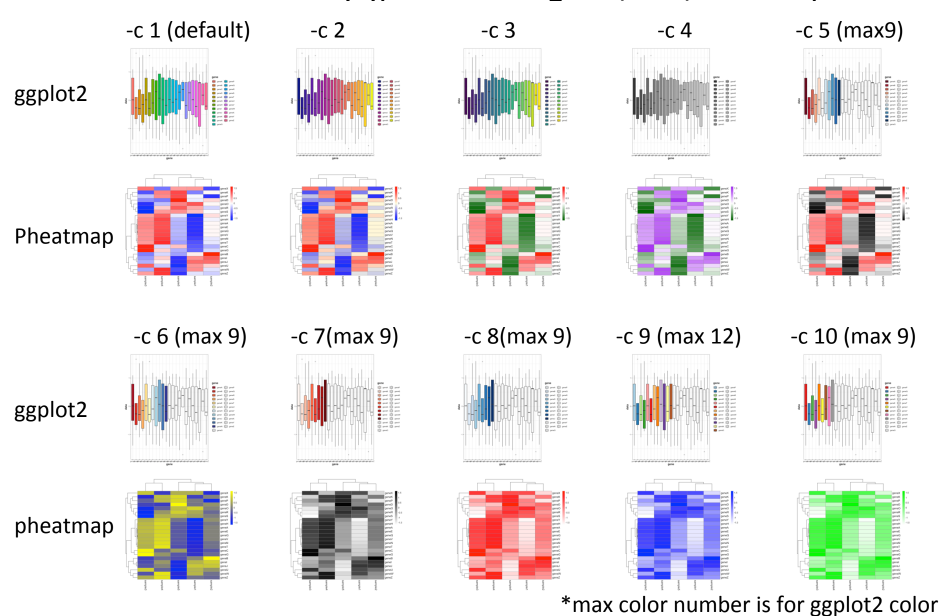


Rscript rnaseq_figure_plotter.r
-i gene_expression_data.txt -t dot_color -l 10
-x gene -z sample



Color (Rscript rnaseq_figure_plotter.r -i gene_expression_data.txt -t box/heatmap -x gene -zs on)

You can customize color by type -cst, --custom_color (colors). Look example 4

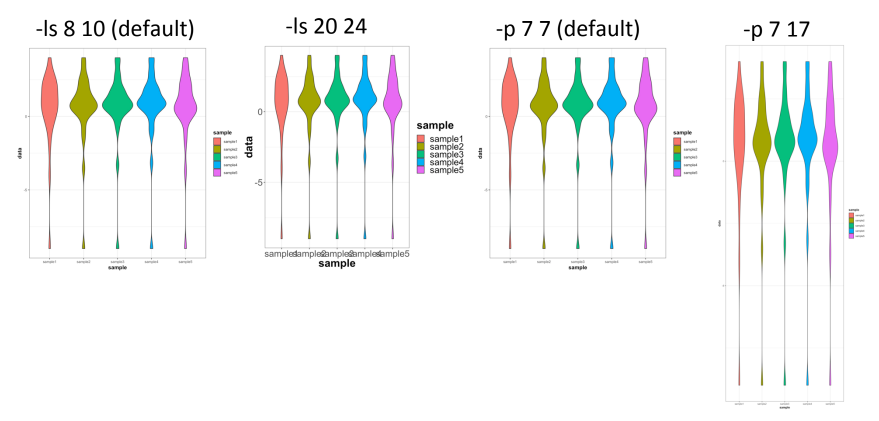


Letter and figure size/file format

Letter Default 8 10; type text and title size of legend and axis, respectively. Split two number by space. Example; 20 24.

Figure Default 7 7; type width and height of figure. Split two number by space. Example; 10 12

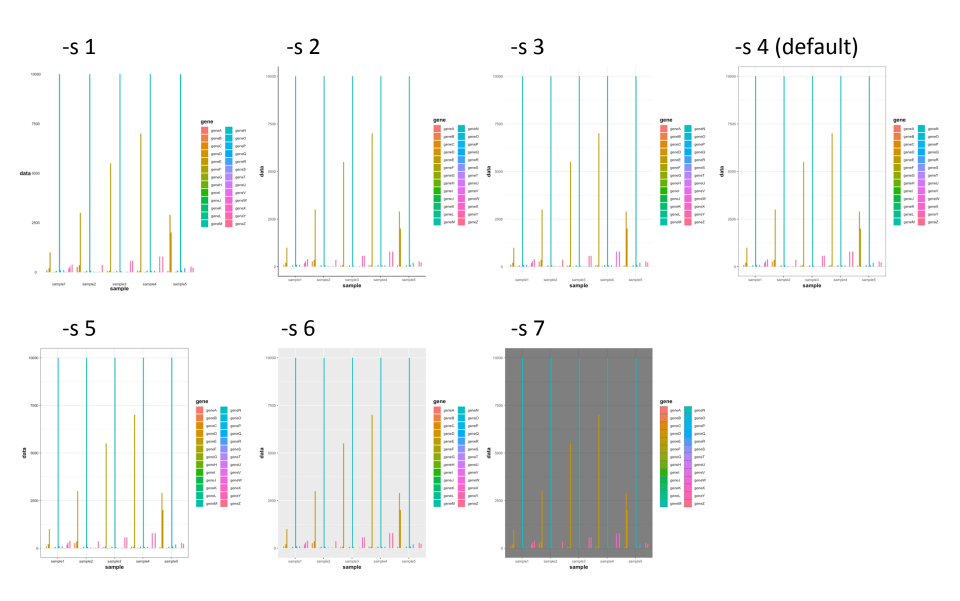
Rscript rnaseq_figure_plotter.r -i gene_expression_data.txt -t violin -l 10



Default is pdf, you can also choose eps, ps, tex (pictex), pdf, jpeg, tiff, png, bmp, svg. Example of usage; save figure in jpeg by (-f jpeg).

 $\textbf{Style} \; (\textbf{Rscript rnaseq_figure_plotter.r -i gene_expression_data.txt -t bar -gp 2 -f jpeg})$

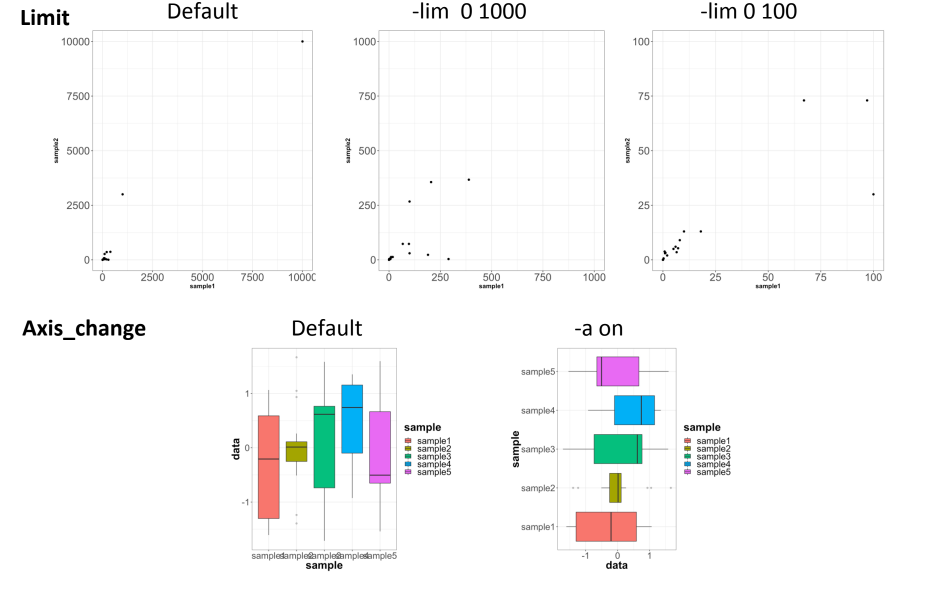
*This function is for every plots excepts heatmap.



Data limit and axis change

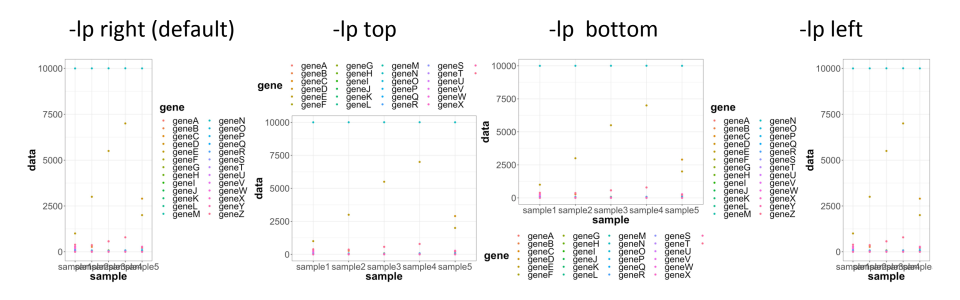
Data limit (Rscript rnaseq_figure_plotter.r -i gene_expression_data.txt -t scatter -ss sample1 sample2)
Axis change(Rscript rnaseq_figure_plotter.r -i gene_expression_data.txt -t box -zs on)

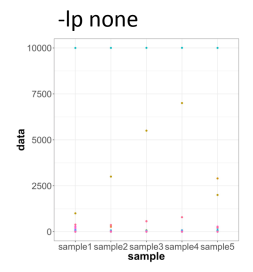
*This function is for every plots excepts heatmap.



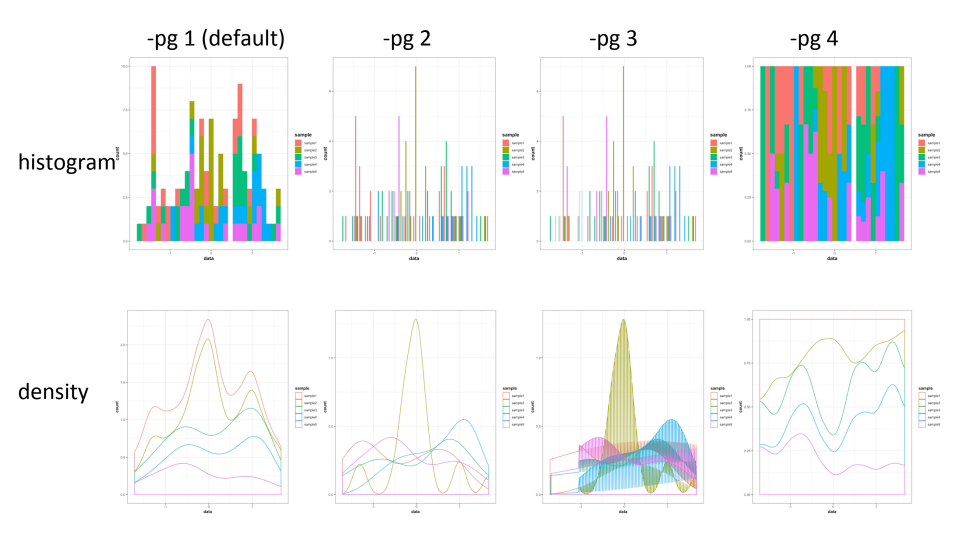
Legend position (Rscript rnaseq_figure_plotter.r -i gene_expression_data.txt -t dot_color -ls 20 24)

*This function is for every plots excepts heatmap and scatter.

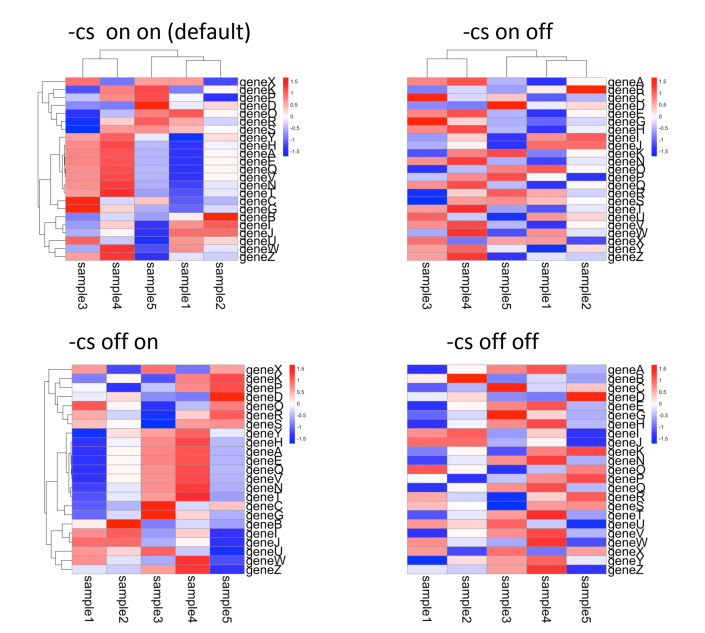




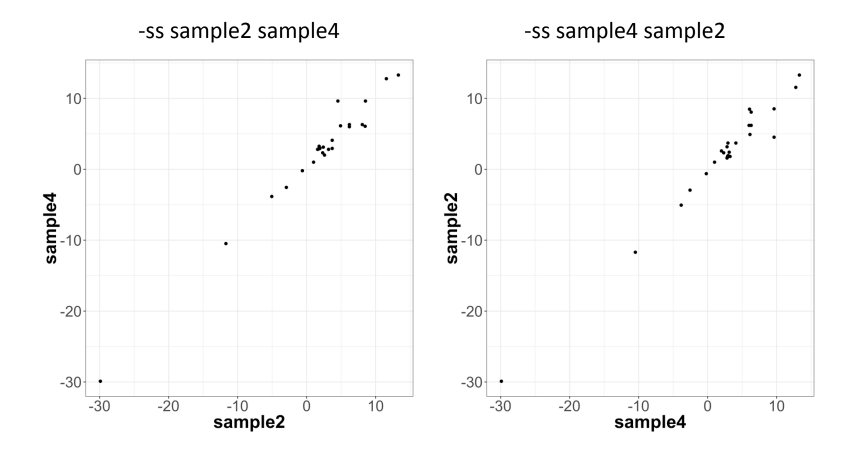
Geom_position (Rscript rnaseq_figure_plotter.r -i gene_expression_data.txt -t histogram/density -zs on) *This function is for bar, density, and histogram.



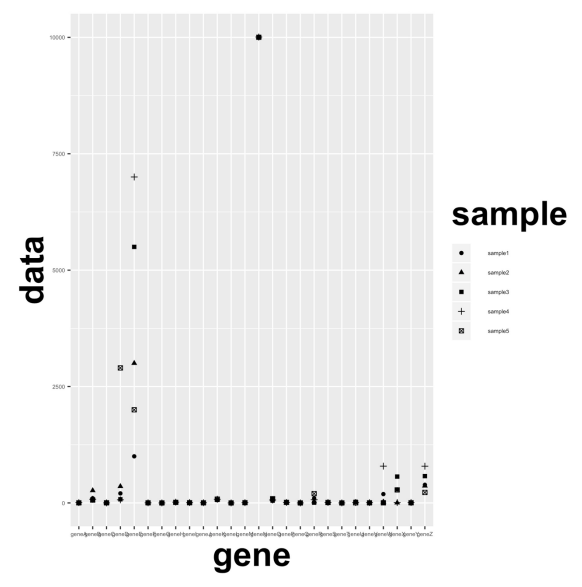
Cluster (Rscript rnaseq_figure_plotter.r -i gene_expression_data.txt -t heatmap -zs on -ls 20 24) *This function is only for heatmap.



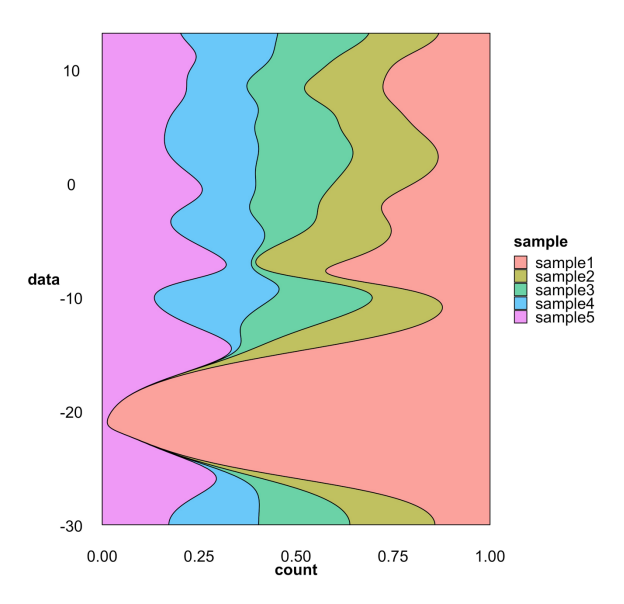
Scatter (Rscript rnaseq_figure_plotter.r -i gene_expression_data.txt -t scatter -ls 20 24 -l e) *This function is only forscatter plot.



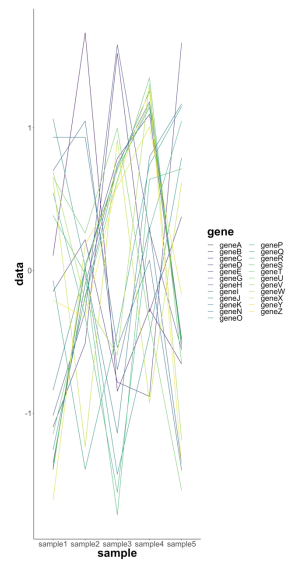
Rscript rnaseq_figure_plotter.r -i gene_expression_data.txt -t dot_shape -o sample_data -x gene -z sample -s 6 -ls 5 30 -f tiff



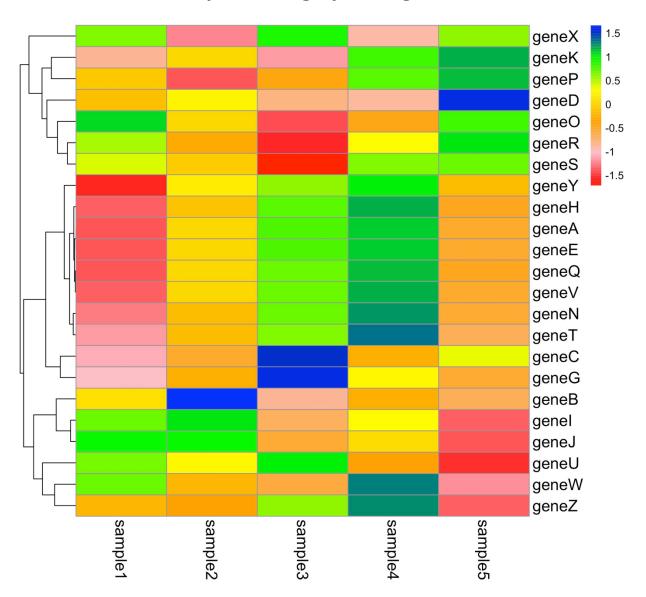
Rscript rnaseq_figure_plotter.r -i gene_expression_data.txt -t density_fill -o sample_test -l e -s 1 -a on -gp 4 -ls 20 20



Rscript rnaseq_figure_plotter.r -i gene_expression_data.txt -t line -zs on -s 2 -c 3 -ls 20 30 -f png -p 10 20



Rscript rnaseq_figure_plotter.r -i gene_expression_data.txt -t heatmap -o sample_heat -zs on -ls 15 15 -cs off on -cst red pink orange yellow green blue #custom color



Rscript rnaseq_figure_plotter.r -i gene_expression_data.txt -t bar -g gene_list.txt -l 10 -c 6 -ls 15 15 -a on -gp 2 -lim "-1" 1.75

