Tutorial: Uploading data sets

Step 1: Create a template

The default template (template 1) includes columns for gene, escape state, and start position.

1. CREATE A TEMPLATE			
Template Type			
Template 1 (gene, state)			
Template 2 (gene, state, tiss_samp)			
✓ Include 'start' position			
♣ Download Template			

Template 1:

gene	samp_state	start	

Example data:

gene	samp_state	start
CD99	escape	154021573
CD99	escape	154021573
CD99	inactive	154021573

The term "samp_state" refers to the state of the gene within a sample or tissue. The sample or tissue can be specified by using template 2.

Template 2:

gene	samp_state	sample	start

Example data:

gene	samp_state	sample	start	
CD99	escape	STMACH	154021573	
CD99	escape	SKINS	154021573	
CD99	inactive	LUNG	154021573	

The "start" position is optional an only necessary for plotting the genes on the x chromosome coordinate system. Deselecting "Include 'start' position" will remove the 'start' column for both templates.

Template Type		
Template 1 (gene, state)		
Template 2 (gene, state, tiss_samp)		
☐ Include 'start' position		

Click the boundard Template button to download the template in csv format.

Step 2: Upload data set



Populate the downloaded template with the necessary data. Note, the final file **must include the column names which reflect one of the available templates**.

Example csv files:

Ł	Example A			
	gene	samp_state	sample	start
	ASMTL	escape	S100	1403139
	ASMTL	escape	S200	1403139
	ASMTL	escape	S300	1403139
	ASMTL	escape	S400	1403139
	ASMTL	inactive	S500	1403139
	ASMTL	escape	S600	1403139
	MECP2	inactive	S100	154021573
	MECP2	inactive	S200	154021573
	MECP2	inactive	S300	154021573
	MECP2	escape	S400	154021573
	MECP2	escape	S500	154021573
	MECP2	escape	S600	154021573

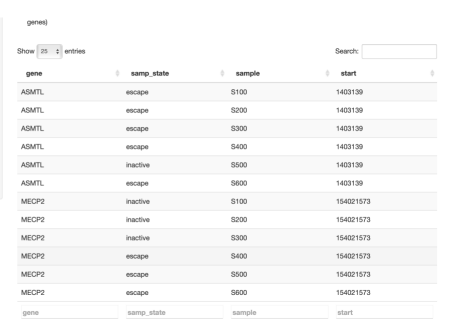
Example B

gene	samp_state	sample
SHOX	escape	AORT
SHOX	escape	AORT
SHOX	escape	AORT
SHOX	inactive	AORT
SHOX	escape	AORT
SHOX	escape	AORT
SHOX	escape	AORT
MECP2	inactive	AORT
MECP2	inactive	AORT
MECP2	inactive	AORT
MECP2	escape	AORT
MECP2	escape	AORT
MECP2	escape	AORT

Click Browse... to select your csv file and upload to the application. Once uploaded, a preview of your data set should be displayed.

Preview of upload:





Step 3: Submit data set

Once you've verified your data set, select

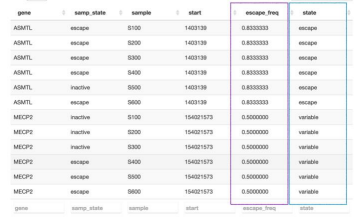
Submit Uploaded Data Set

3. SUBMIT

Submit Uploaded Data Set

The application will produce the following levels of information:

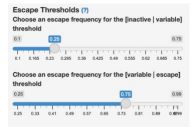
Escape frequencies and escape statuses based upon samp_state



Note: the default thresholds applied are:

- %escape ≤ .25 → Silenced (S)
- .25 < %escape < .75 → Variable escape (VE)
- %escape ≥ .75 → Escape (E)

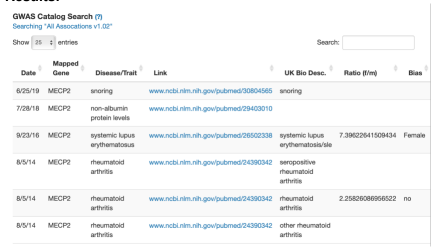
These can be changed with the sliders in the left side panel.



- GWAS associations of selected genes of interest
 - 1. Select the Association Data tab
 - 2. Select genes of interest in left side panel



3. Results:



Genomic positions of genes and escape statuses (if "start" position is included)

