Feature linkage analysis

While processing 10x Chromium Single Cell Multiome ATAC + Gene Expression sequencing data via 'cellranger-arc count' pipeline, Feature linkages analysis is performed as pairs of genomic features, such as peaks and genes, that have significant correlation in signals across cells. Because it provides a basis for inferring enhancer-gene targeting relationships and constructing transcriptional networks. The features with strong linkages are considered to be "co-expressed" and enriched for a shared regulatory mechanism.

Partek Flow provides the opportunity to our users to explore the linkage relationships among different features including peaks and genes, peaks and peaks, and genes and genes. A tab-delimited file containing information of feature linkages inferred from Flow Cell Ranger - ATAC task will be loaded into Integrative Genome Viewer (IGV)[1] for exploration if Feature linkage analysis task has been completed successfully.

Running Feature linkage analysis

To run Feature linkage analysis task (Figure 1),

- Click one datanode that has both features of ATAC and gene expression;
- Click the Feature linkage analysis task under Peak analysis section in the toolbox;
- Click the Finish button to complete the submission.



Figure 1. Feature linkage analysis task in Flow

There will be no inputs needed if the FASTQ is converted to counts matrix within Flow. However, if users processed the FASTQ files outside of Partek, and imported the counts matrix into Flow later. The feature_linkage.bedpe file in outs/analysis/feature_linkage from Cell Ranger output will be needed for each sample (Figure 2) to complete the analysis.

Feature linkages files	Sample name	File	Browse files		
	mouseBrain5K	feature_linkage.bedpe	Browse		
Back Finish			-		

Figure 2. Specify the .bedpe file for projects started with counts matrix in Flow.

Task report

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A new datanode will be displayed as the task is finished. Double click on the output datanode, Flow will bring you to the IGV browser where you could explore the "co-expressed" feature pairs (Figure 3).

IGV hg38 chr10 ∽ c	hr10:29,919,288-46,643,	965 Q 16	mb						(Cursor Guide	Center Line Track Labels Save SVG) 🖨 🔫	
0_mb 31 mb 32 mb 34 mb 35 mb 36 mb 37 mb 38 mb 39 mb 40 mb 0_mb 31 mb 32 mb 34 mb 35 mb 36 mb 37 mb 38 mb 39 mb 40 mb 1_mg38_ensembl_release98_v28 N2 H 11 H									b 42		43 mb 44 mb 44 11 1 MB 1 MB 1 H 1 H 1 H 1 1 1 H 30 H 10 H 1 H 1 H 8889 PATTO LINCOLEO OXCL12 RAG	i mb II-1111 114 - 1111 II-1111 114 - 1111 II-1111 114 - 1111 11 II-1111 114 - 1111 11 II-1111 114 - 1111 11	46 mb
pbmc_unsorted_3k_83			V			,	2			ŀ			
Sample name ©	Chromosome 1 \$	Start 1 ≎	End 1 \$	Chromosome 2 \$	Start 2 ≎	End 2 ≎	Name ≎	Score ≎	Strand 1 ≎	Strand 2 \$	Significance (-log10 pvalue) \$	Distance \$	Linkage type \$
pbmc_granulocyte_sorted_3k_S12	chr1	822,869	823,760	chr1	999,980	1,000,172	<linc00115_distal><hes4></hes4></linc00115_distal>	0.48			5.66	176,761	peak-gene
pbmc_granulocyte_sorted_3k_S12	chr1	955,134	956,039	chr1	999,980	1,000,172	<noc2l_distal><hes4></hes4></noc2l_distal>	0.48			5.13	44,489	peak-gene
pbmc_granulocyte_sorted_3k_S12	chr1	816,877	817,780	chr1	1,001,137	1,013,497	<fam87b_promoter><isg15></isg15></fam87b_promoter>	0.47			7.71	189,988	peak-gene
pbmc_granulocyte_sorted_3k_S12	chr1	999,859	1,000,448	chr1	1,001,137	1,013,497	<hes4_promoter><isg15></isg15></hes4_promoter>	0.57			10.84	7,163	peak-gene

Figure 3. Feature linkage task report in Flow.

The feature_linkage.bedpe file[2] outputted from Cell Ranger pipeline is available in task report as a table. In the report table, each row is a region of a peak and includes the following information:

- Sample name: name for each sample
- . Chromosome 1: the name of the chromosome on which the first feature exists.
- Start 1: the starting position of the first feature on that chromosome.
- End 1: the ending position of the first feature on that chromosome.
- Chromosome 2: the name of the chromosome on which the second feature exists.
- ٠ Start 2: the starting position of the second feature on that chromosome.
- End 2: the ending position of the second feature on that chromosome.
- Name: the name of the linkage features with the format of <name1><name2>, in which name1 and name2 are based on gene symbol or peak annotation.
- Score: linkage correlation, ranging from -1 to 1.
- Strand 1: all set to "." Strand 2: all set to "."
- Significance: linkage significance: -log10 (p-value) after multiple testing correction (FDR, false discovery rate). Capped at 299.
- Distance: distance in base pairs from feature 2 to feature 1. •
- Linkage type: can be "peak-peak", "gene-peak" or "peak-gene" depending on the type of gene or peak for feature 1 and feature 2.

Filter feature linkage task

To filter out and visualize only the linkages that users are interested in is also made possible through the Filter feature linkages task in Flow (Figure 4). Users are able to download the .bedpe file from Flow and explore them via their stand-alone IGV.



Figure 4. Filter feature linkage task is available in Flow.

References

- https://software.broadinstitute.org/software/igv/
 https://support.10xgenomics.com/single-cell-multiome-atac-gex/software/pipelines/latest/output/analysis

Additional Assistance

If you need additional assistance, please visit our support page to submit a help ticket or find phone numbers for regional support.

