

Data exploratory analysis Antonia Chroni for SJCRH DNB_BINF_Core

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## ##	The following object is masked _byGlobalEnv:	
##	root dir	

PI: NA
Project: test-dataset
Task: NA
Project Lead(s): NA
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DNB Bioinformatics Core Pipeline: Standard sc-/sn-ATAC-Seq Analysis in

10X Genomics data

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1 Information about this notebook

This is an exploratory analysis of the data in the project. We are investigating number of samples overall per condition and variables as defined in the params.

We are looking for cohorts that fit the following criteria.

- Control vs condition (min. 3+3 samples)
- Number of cells/sample size of datasets might determined packages/pipelines to be used
- Single cell/Single-nucleus ATAC
- Integration of scRNA-seq and scATAC-seq data from the same biological system (multiple modalities)
- Pipeline for same samples but not same cells with scRNA-seq and scATAC-seq
- Annotate scATAC-seq cells via label transfer by using scRNA data: cell type annotation
- Available bulk ATAC-seq data for the same samples (matched) this could be used for cell type annotation
- 2 Set up
- 3 Directories and paths to file Inputs/Outputs
- 4 Read raw data file
- 5 Read processed data file
- 6 Read Jackie data file (2025-01-03) and update table
- 7 10x matched scRNA-seq and scATAC-seq cohort

7.1 Type of sequencing assay and unit

We should investigate if there are samples from two different sequencing technologies and unit.

Single cell sequencing was done by using cell, nucleus and there are RNA, ATAC and 10Xv3, 10Xv2 sequencing technologies in the database. Cohort is formed and processed accordingly.

7.2 Number of samples per experiment

Table 1: Summary of samples per experiment

experiment	n
Cerebellum	9
More Retina	7
Stressed Retina	3
Victoria Knockout	8

7.3 Number of samples per experiment, seq_unit, assay, and condition

Table 2: Summary of samples per experiment, seq_unit, assay, and condition $\,$

experiment	seq_unit	assay	condition	n
Cerebellum	cell	RNA	age	2
Cerebellum	cell	RNA	sorted	2
Cerebellum	nucleus	ATAC	age	3
Cerebellum	nucleus	ATAC	sorted	2
More Retina	cell	ATAC	age	2
More Retina	cell	ATAC	none	2
More Retina	cell	ATAC	sorted	1
More Retina	cell	RNA	age	1
More Retina	cell	RNA	sorted	1
Stressed Retina	cell	RNA	LPS Injection	1
Stressed Retina	cell	RNA	PBS Injection	1
Stressed Retina	nucleus	ATAC	LPS Injection	1
Victoria Knockout	cell	RNA	knock-out	2
Victoria Knockout	cell	RNA	wt	2
Victoria Knockout	nucleus	ATAC	knock-out	2
Victoria Knockout	nucleus	ATAC	\mathbf{wt}	2

7.4 Number of samples per experiment, tissue, seq_unit, assay, condition, and sample

 $\label{thm:condition} \begin{tabular}{ll} Table 3: Summary of samples per experiment, tissue, seq_unit, assay, seq_technology, condition, and Sample \\ \end{tabular}$

				seq_technol-	-		
experiment	tissue	${\rm seq_unit}$	assay	ogy	condition	Sample	n
Cerebellum	Cerebel- lum	cell	RNA	10Xv3	age	6 week cerebellum	1
Cerebellum	Cerebel- lum	cell	RNA	10Xv3	age	P0 cerebellum	1
Cerebellum	Cerebel- lum	cell	RNA	10Xv3	sorted	АТОН НІ	1
Cerebellum	Cerebel- lum	cell	RNA	10Xv3	sorted	ATOH Low	1
Cerebellum	Cerebel- lum	nucleus	ATAC	10Xv2	age	6 week cerebellum	2
Cerebellum	Cerebel- lum	nucleus	ATAC	10Xv2	age	P0 cerebellum	1
Cerebellum	Cerebel- lum	nucleus	ATAC	10Xv2	sorted	АТОН НІ	1
Cerebellum	Cerebel- lum	nucleus	ATAC	10Xv2	sorted	ATOH Low	1
More Retina	Retina	cell	ATAC	10Xv2	age	E14.5 Retina	2
More Retina	Retina	cell	ATAC	10Xv2	none	Retina	2
More Retina	Retina	cell	ATAC	10Xv2	sorted	NRL Dep	1
More Retina	Retina	cell	RNA	10Xv3	age	E14.5	1
More Retina	Retina	cell	RNA	10Xv3	sorted	NRL Dep	1
Stressed	Retina	cell	RNA	10Xv3	LPS	LPS	1
Retina	reema	cen	101111	102110	Injection	LI U	
Stressed	Retina	cell	RNA	10Xv3	PBS	PBS	1
	пенна	cen	ΠNA	10AV3		грэ	1
Retina	D 4:	1	А ТА С	10V 0	Injection	IDC ATAC	1
Stressed	Retina	nucleus	ATAC	10Xv2	LPS	LPS_ATAC	1
Retina	D	11	DATA	1077.0	Injection	CELLO 01	
Victoria	Retina	cell	RNA	10Xv3	knock-out	SEKO_21	1
Knockout	-					G7770	
Victoria	Retina	cell	RNA	10Xv3	knock-out	$SEKO_25$	1
Knockout	5	11	D37.4	1077		TTT. 4	_
Victoria	Retina	cell	RNA	10Xv3	wt	Wt1	1
Knockout							
Victoria	Retina	cell	RNA	10Xv3	wt	Wt2	1
Knockout							
Victoria	Retina	nucleus	ATAC	10Xv2	knock-out	$SEKO_21$	1
Knockout						-	
Victoria	Retina	nucleus	ATAC	10Xv2	knock-out	$SEKO_25$	1
Knockout							
Victoria	Retina	nucleus	ATAC	10Xv2	wt	$\mathrm{Wt}1$	1
Knockout							
Victoria Knockout	Retina	nucleus	ATAC	10Xv2	wt	Wt2	1

8 10x Genomics Multiome cohort

8.1 Type of sequencing assay and seq_unit

We should investigate if there are samples from two different sequencing technologies and unit.

Single cell sequencing was done by using nucleus and there are RNA, ATAC and 10X sequencing technologies in the database. Cohort is formed and processed accordingly.

8.2 Number of samples per experiment

Table 4: Summary of samples per experiment

experiment	
Multiome Retina	6

8.3 Number of samples per experiment, seq_unit, assay, and condition

Table 5: Summary of samples per experiment, seq_unit, assay, and condition $\,$

experiment	seq_unit	assay	condition	n
Multiome Retina	nucleus	ATAC	age	3
Multiome Retina	nucleus	RNA	age	3

8.4~ Number of samples per experiment, tissue, seq_unit, assay, seq_technology, condition, and sample

 $\label{thm:condition} \begin{tabular}{ll} Table 6: Summary of samples per experiment, tissue, seq_unit, assay, seq_technology, condition, and Sample \\ \end{tabular}$

				seq_technol-			
experiment	tissue	${\rm seq_unit}$	assay	ogy	condition	Sample	n
Multiome	Retina	nucleus	ATAC	10X	age	Adult Retina	1
Retina							
Multiome	Retina	nucleus	ATAC	10X	age	E14.5 Retina	1
Retina							
Multiome	Retina	nucleus	ATAC	10X	age	P0 Retina	1
Retina							
Multiome	Retina	nucleus	RNA	10X	age	Adult Retina	1
Retina							
Multiome	Retina	nucleus	RNA	10X	age	E14.5 Retina	1
Retina							
Multiome	Retina	nucleus	RNA	10X	age	P0 Retina	1
Retina							

9 Notes

I have identified the following datasets that almost fit the criteria for the testing phase:

- 10x matched scRNA-seq and scATAC-seq: Victoria Knockout experiment with 1 replicate/knock-out group (4 samples for 10x RNA + 4 samples 10x ATAC). Replicates could be potentially grouped together and have 2 replicates for WT and 2 replicates for knock-out.
- 10x Genomics Multiome: Multiome Retina experiment with 1 replicate/age group (3 samples for 10x RNA (Multiome) + 3 samples for 10x ATAC (multiome).

10 Session Info

```
## R version 4.4.0 (2024-04-24)
## Platform: x86_64-pc-linux-gnu
## Running under: Red Hat Enterprise Linux 8.8 (Ootpa)
## Matrix products: default
## BLAS:
           /usr/lib64/libblas.so.3.8.0
## LAPACK: /usr/lib64/liblapack.so.3.8.0
##
## locale:
                                    LC_NUMERIC=C
    [1] LC_CTYPE=en_US.UTF-8
##
    [3] LC_TIME=en_US.UTF-8
##
                                    LC_COLLATE=en_US.UTF-8
    [5] LC_MONETARY=en_US.UTF-8
                                    LC_MESSAGES=en_US.UTF-8
##
    [7] LC_PAPER=en_US.UTF-8
                                    LC_NAME=C
##
    [9] LC_ADDRESS=C
                                    LC_TELEPHONE=C
## [11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
##
## time zone: America/Chicago
## tzcode source: system (glibc)
## attached base packages:
                 graphics grDevices utils
## [1] stats
                                                datasets methods
                                                                     base
##
## other attached packages:
   [1] readxl_1.4.3
                        knitr_1.48
                                         lubridate_1.9.3 forcats_1.0.0
##
    [5] stringr_1.5.1
                        dplyr_1.1.4
                                         purrr_1.0.2
                                                         readr_2.1.5
   [9] tidyr_1.3.1
                        tibble_3.2.1
                                         ggplot2_3.5.1
                                                         tidyverse_2.0.0
##
## [13] yaml_2.3.10
##
## loaded via a namespace (and not attached):
    [1] bit_4.0.5
                          gtable_0.3.5
##
                                             jsonlite_1.8.8
                                                                crayon_1.5.3
##
    [5] compiler_4.4.0
                          tidyselect_1.2.1
                                             parallel_4.4.0
                                                                jquerylib_0.1.4
    [9] scales_1.3.0
                          fastmap_1.2.0
                                             mime_0.12
                                                                R6_2.5.1
## [13] generics_0.1.3
                          munsell_0.5.1
                                             bslib_0.8.0
                                                                pillar_1.9.0
## [17] tzdb_0.4.0
                          rlang_1.1.4
                                             utf8 1.2.4
                                                                stringi_1.8.4
## [21] cachem_1.1.0
                          xfun_0.47
                                             sass_0.4.9
                                                                bit64_4.0.5
## [25] timechange_0.3.0
                          cli_3.6.3
                                             withr_3.0.1
                                                                magrittr_2.0.3
## [29] digest_0.6.37
                          grid_4.4.0
                                             vroom_1.6.5
                                                                hms_1.1.3
## [33] lifecycle_1.0.4
                          vctrs_0.6.5
                                             evaluate_0.24.0
                                                                glue_1.7.0
## [37] cellranger_1.1.0
                          fansi_1.0.6
                                             colorspace_2.1-1
                                                                rmarkdown_2.28
## [41] tools_4.4.0
                          pkgconfig_2.0.3
                                             htmltools_0.5.8.1
```