# Gene Analysis of Schizophrenia Patient Samples

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## Summary

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### Introduction

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#### Methods

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#### Preview of the Data

The data used is from the Emani et al. (2024) set of cohorts, with its samples. The raw data of the annotated expression matrices can be found here. All the samples come from the prefrontal cortex (PFC). The metadata contains the information of the following 12 cohorts:

- 1. CMC
- 2. DevBrain
- 3. Girgenti-multiome
- 4. IsoHuB
- 5. LIBD
- 6. Ma\_et\_al (Ma et al. 2022)
- 7. MultiomeBrain
- 8. PTSDBrainomics
- 9. SZBDMulti-Seq
- 10. ROSMAP
- 11. UCLA-ASD
- 12. Velmeshev\_et\_al (Velmeshev et al. 2019)

However, because no annotated matrices were provided for the ROSMAP cohort, these were filtered out from the metadata.

This is an overview of each cohort:

Table 1: Each cohort with the total patients studied, the mean of the age of death, and disease studied.

Cohort	Patients	$Mean\_Age$	Disorder_studied
CMC	100	71	Schizophrenia
DevBrain	16	29	Williams Syndrome, ASD
Girgenti-snMultiome	19	49	None
IsoHuB	4	22	None
LIBD	10	50	None
Ma_et_al	2	57	None
MultiomeBrain	21	42	Schizophrenia, Bipolar Disorder
PTSDBrainomics	19	47	MDD, PTSD
SZBDMulti-Seq	72	65	Bipolar Disorder, Schizophrenia
UCLA-ASD	52	23	ASD
$Velmeshev\_et\_al$	27	18	ASD

From Table 1, we can see that the CMC cohort has the biggest number of patient samples with 100 samples, followed by SZBDMulti-seq with 72. Ma\_et\_al has the smallest size, with only 2 samples. CMC and SZBDMulti-seq also have the highest mean of age of death at 71

years and 65 years respectively. Velmeshev\_et\_al has the lowest one, at 18 years. A variety of diseases are studied in these cohorts, with some cohorts studying none. These diseases include schizophrenia, autism spectrum disorder (ASD), bipolar disorder, major depressive disorder (MDD), post-traumatic stress disorder (PTSD), and Williams Syndrome.

Table 2: Number of patients per condition per cohort. Includes the mean age and the biological sex as well.

Cohort	Disorder	Number_of_PatientsMean	_Age N_	_Male N_	_Female
$\overline{\mathrm{CMC}}$	Control	53	71	32	21
CMC	Schizophrenia	47	71	33	14
DevBrain	ASD	9	24	8	1
DevBrain	Control	4	36	4	0
DevBrain	Williams	3	31	2	1
	Syndrome				
Girgenti-	Control	19	49	15	4
snMultiome					
IsoHuB	Control	4	22	2	2
LIBD	Control	10	50	6	4
$Ma\_et\_al$	Control	2	57	1	1
MultiomeBrain	Bipolar Disorder	10	40	6	4
MultiomeBrain	Control	5	39	4	1
MultiomeBrain	Schizophrenia	6	46	3	3
PTSDBrainomics	Control	9	49	7	2
PTSDBrainomics	MDD	4	52	3	1
PTSDBrainomics	PTSD	6	41	5	1
SZBDMulti-Seq	Bipolar Disorder	24	70	12	12
SZBDMulti-Seq	Control	24	62	12	12
SZBDMulti-Seq	Schizophrenia	24	64	12	12
UCLA-ASD	ASD	27	23	22	5
UCLA-ASD	Control	25	24	21	4
$Velmeshev\_et\_al$	ASD	13	15	10	3
$Velmeshev\_et\_al$	Control	14	20	9	5

Table 2 indicates the number of patients per condition in each cohort. The CMC cohort has the highest numbers, with 53 control samples and 47 schizophrenia samples. The UCLA-ASD is second, with 27 ASD samples and 25 control samples. The mean of age at death and the distribution of the biological sexes is also shown in this table.

It must be noted that many patients' age at death was marked as "89+". In order to determine the age distribution and the mean age of death, these values were converted to "89". The number of samples that were marked as "89+" from the Schizophrenia cohorts is in Table 3.

Table 3: Number of patients annotated as "89+" in Schizophrenia cohorts.

Cohort	Disorder	Plus89_patients
$\overline{\mathrm{CMC}}$	Control	9
CMC	Schizophrenia	5
SZBDMulti-Seq	Bipolar Disorder	1
SZBDMulti-Seq	Control	1
${\bf SZBDMulti\text{-}Seq}$	Schizophrenia	2

The age distribution of each condition per cohort was also examined.

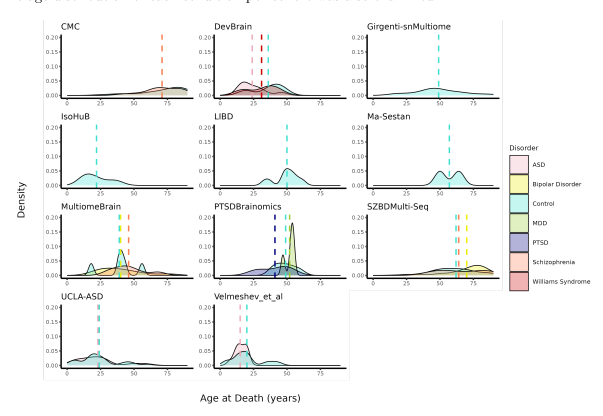


Figure 1: Age distribution in each cohort.

In Figure 1, the dotted vertical lines are the mean of the age at death for each condition.

The number of genes and single cells per each Schizophrenia cohort was also looked at in Table 4. The number of genes is per sample, whereas the number of cells is the total number for all samples in a cohort.

Table 4: Number of genes and total number of cells per each Schizophrenia cohort.

Cohort	Genes	Total_Cells
CMC	33792	456560
SZBDMulti-Seq	34361	603281
MultiomeBrain	33822	134666

The number of genes is quite similar across 3 cohorts. SZBDMulti-Seq has the highest number of total cells, at 603281. MultiomeBrain has the lowest number of total cells, at 134666.

#### References

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