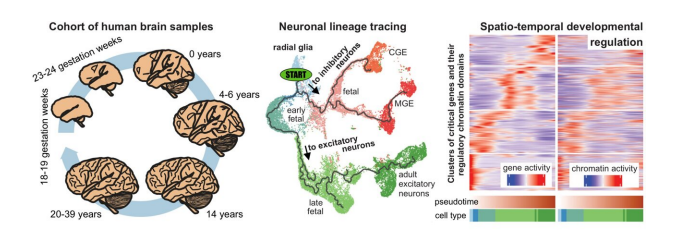


## First 'multiome' atlas of cell development in the human cerebral cortex from before birth to adulthood constructed

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Study summary graphic. Credit: Mount Sinai Health System

A team of researchers from the Icahn School of Medicine at Mount Sinai and Yale University School of Medicine has created the first "multiome" atlas of brain cell development in the human cerebral cortex across six broad developmental time points from fetal development into adulthood, shedding new light on their roles during brain development and disease.

"Multiome" refers to the simultaneous analysis of multiple types of genetic information within the same biological sample. They can include



the genome, the DNA encoded in our cells; the transcriptome, the RNA copies that the cell makes from the genome; and the epigenome, chemical modifications and regulatory factors that determine the accessibility of chromatin,

As described in <u>Science Advances</u>, the researchers used new scientific tools to analyze and describe two kinds of information from each cell—gene expression (transcriptome) as well as DNA structure (epigenome)—enabling them to categorize cell types at different developmental stages.

The data revealed specific changes in the structure of chromatin that precede <u>gene expression</u>. These changes are crucial for numerous processes, including the formation of neurons.

Additionally, their analysis pinpointed regions of chromatin associated with the regulation of genes known to play a pivotal role in human brain development. Notably, they revealed that these regulatory regions are often enriched for genetic signals associated with increased risk for neuropsychiatric disorders such as schizophrenia or bipolar disorder.

"Human brain development starts during embryogenesis and extends postnatally through infancy, childhood, adolescence, and young adulthood," says Panos Roussos, M.D., Ph.D., Professor of Psychiatry, and Genetics and Genomic Sciences, Director of the Center for Disease Neurogenomics at Icahn Mount Sinai, and senior author of the paper.

"Given the variable age of onset of different neurodevelopmental disorders, it is critical to examine the effect of risk factors across the full spectrum of brain development. Through the development of this atlas, we have gained a deeper understanding of the intricate regulatory mechanisms underlying brain development and disease."



The comprehensive atlas is now accessible to other researchers through an online repository so others can engage with the data, visualize it effectively, and use it for their own research. The paper is included in a package of 21 research studies across *Science, Science Advances* and *Science Translational Medicine* that detail research conducted as part of the National Institutes of Health's BRAIN Initiative Cell Census Network (BICCN), a program launched in 2017 to create an atlas of the human and non-human primate brain at the cell-type level in unprecedented detail.

The basis for neuropsychiatric disease in adulthood can often be influenced by alterations in the cellular composition of the brain that arise during development. In addition to creating the first atlas of human brain cell development in the human cerebral cortex, the research team prioritized 152 risk genes that play a causal role in a range of neuropsychiatric disorders.

Their findings go beyond existing knowledge by mapping cell type and temporally specific genetic loci implicated in neuropsychiatric disorders. For example, they discovered that Tourette syndrome is associated with oligodendrocytes, while obsessive-compulsive disorder is associated with astrocytes. Both of these associations between diseases and cell types were previously unknown, and the discoveries contribute to a deeper understanding of the complex relationships between different <u>cell types</u> and <u>neuropsychiatric disorders</u>.

"It is important to recognize that the most effective therapeutic interventions should be customized to target deficiencies in gene function as specific developmental stages," says Jaroslav Bendl, Ph.D., Assistant Professor of Psychiatry, and Genetics and Genomic Sciences, at Icahn Mount Sinai and a co-author of the work. "Only by doing so will we be able to minimize further damage and improve outcomes for individuals affected by these disorders."



Having demonstrated that an atlas of cellular development in the human cerebral cortex was possible, the team is currently expanding their study by analyzing a larger sample cohort and including different brain regions. By so doing, they aim to achieve increased resolution and gain deeper understanding of the intricate regulatory mechanisms and further unravel the complex regulatory logic underling <u>brain development</u> and disease.

**More information:** Kenneth Chiou et al, A single-cell multi-omic atlas spanning the adult rhesus macaque brain, *Science Advances* (2023). DOI: 10.1126/sciadv.adh1914. www.science.org/doi/10.1126/sciadv.adh1914

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