REVIEW

Application of Computational Biology to Decode Brain Transcriptomes

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Abstract The rapid development of high-throughput sequencing technologies has generated massive valuable \textit{brain transcriptome atlases}, providing great opportunities for systematically investigating gene expression characteristics across various brain regions throughout a series of developmental stages. Recent studies have revealed that the transcriptional architecture is the key to interpreting the molecular mechanisms of brain complexity. However, our knowledge of brain transcriptional characteristics remains very limited. With the immense efforts to generate high-quality brain transcriptome atlases, new computational approaches to analyze these high-dimensional multivariate data are greatly needed. In this review, we summarize some public resources for brain transcriptome atlases and discuss the general computational pipelines that are commonly used in this field, which would aid in making new discoveries in brain development and disorders.

Introduction

The mammalian brain is an evolutionary miracle that contains well-organized molecules, cell types, and neuronal circuits in each subregion; some of these features are closely connected at both the structural and functional levels. Moreover, brain development is an intricate, highly regulated process that continues throughout embryonic growth, and these lifespan program codes are conserved among species \cite{1}. The complicated properties of the brain are mainly reflected in the complexity of its transcriptomic architecture, including highly ordered gene expression and elaborate transcriptional regulation. For example, the majority of genes (>80\%) are expressed in the mammalian brain \cite{2}, and the expression profiles of these genes show great variability during development, with the most remarkable changes occurring during development in prenatal and postnatal stages \cite{3,7}. On the other hand, brain tissues exhibit the smallest transcriptomic changes compared with other organs \cite{8,9}. Therefore, understanding the spatiotemporal characteristics of gene expression can offer valuable insights into brain functional specialization and the
roles of key genes during brain development. Furthermore, analyzing the transcriptomic architecture of normal brain development and function is of vital importance to determine the causes of a variety of complicated neurological disorders.

In the last decade, many quantitative methods have been applied to explore the expression of individual genes, particularly the spatial and temporal patterns across the brain. The development of microarray analysis and various high-throughput sequencing technologies has made it possible to investigate the expression of genes in a high-throughput manner, yielding large datasets. Specifically, single-cell sequencing can be used to quantify the transcriptome of a single cell, providing major opportunities to parse the complex cellular composition of the brain. However, analysis of such high-dimensional data remains substantially complex and requires more effective and sophisticated computational methods and models. Recent progress in computational and systems biology fields has facilitated transcriptomic studies with high precision to obtain new insights into the transcriptional characteristics of the brain.

In this review, we introduce a variety of brain transcriptome atlases and discuss how to apply computational methods to elucidate the relationships between gene expression and brain function as well as the relationships between brain development and disease. Many of these relationships have been discovered by following the general pipeline of brain transcriptome analysis (Figure 1). Finally, we state some limitations in recent transcriptome studies and offer some directions for future studies.

**Brain transcriptomic atlas resources**

In the past decade, an increasing number of researchers have realized the importance of large-scale brain transcriptome data, and various countries have launched big brain research projects, which have greatly promoted the study of the molecular mechanisms of brain organization and function. The rapid development of high-throughput technologies has made it possible to quantify the expression of thousands of genes simultaneously. Currently, various brain transcriptome datasets from humans and other species can be obtained from different molecular platforms, such as microarray, RNA sequencing, and *in situ* hybridization (ISH). For rodents, the Gene Expression Nervous System Atlas (GENSAT) [10,11] and GenePaint [12] have provided expression signals for thousands of genes in developing and adult mouse brains. However, compared with mouse brain atlases, the available human brain expression atlas is less abundant because there are more difficulties in obtaining, storing, and analyzing human postmortem brain tissues [13]. Fortunately, several studies have investigated gene expression variations among different brain regions [14,15] and at different development time points [3–7,16]. Furthermore, a series of transcriptome atlases of the developing and adult mouse brains [2,17], the developing and adult human brains [18,19], and the nonhuman primate (NHP) brains [20–22] have been released. Specifically, the Allen Institute for Brain Science (http://brain-map.org/) possesses comprehensive transcriptomic sources from mouse and human brains and is a great resource for many neuroscience fields [23]. To facilitate the application of these data, we have summarized some available brain transcriptome resources in Table 1. Notably, Jason et al. has provided a detailed user guide for some brain transcriptome databases in another review [24]. In this review, we include a series of data released recently. We believe that these available transcriptome data are essential components for investigating the complex molecular architecture of the brain at a large scale.

**Analyzing brain-wide gene expression patterns**

**Spatial and temporal gene expression analyses**

One important aspect of brain complexity is that the brain is organized into multiple functional regions with distinct
| Species  | Sample  | Age      | Region | Method                  | Web link                               | Data access | Annotation     | Refs. |
|----------|---------|----------|--------|-------------------------|----------------------------------------|-------------|----------------|-------|
| Mouse    | Bulk tissues | Multiple | CNS    | BAC transgenic; ISH     | http://www.gensat.org/                | URL         | Spatiotemporal | [10,11] |
| Mouse    | Bulk tissues | Multiple | Multiple | ISH                     | http://www.genepaint.org               | URL         | Spatiotemporal | [12]   |
| Mouse    | Bulk tissues | Multiple | Multiple | ISH                     | http://www.emouseatlas.org/emage/     | URL         | Spatiotemporal | [25]   |
| Mouse    | Bulk tissues | Lifespan  | Multiple | ISH                     | http://developingmouse.brain-map.org/ | URL         | Spatiotemporal | [17]   |
| Mouse    | Bulk tissues | Postnatal | NCX    | MiD; RNA-seq            | http://hbatlas.org/mouseNCXtranscriptome | SRP031888   | Spatiotemporal | [26]   |
| Mouse    | Bulk tissues | Adult    | NCX layers | RNA-seq               | http://genserv.anat.ox.ac.uk/layers    | GSE27243    | Spatial        | [27]   |
| Mouse    | Bulk tissues | Adult    | Multiple | ISH                     | http://mouse.brain-map.org/           | URL         | Spatial        | [2]    |
| Mouse    | Bulk tissues | Postnatal | Forebrain | FACS; PAN; microarray  | www.ncbi.nlm.nih.gov/geo               | GSE9566     | Cell-type specific | [28] |
| Mouse    | Bulk tissues | Adult    | NCX    | PAN; FACS; RNA-seq      | http://web.stanford.edu/group/barres_lab/brain_rnaseq.html | GSE52564    | Cell-type specific | [29] |
| Mouse    | Bulk tissues | Adult    | CNS    | TRAP; microarray        | http://genetics.wustl.edu/jdlab/csea-tool-2 | GSE13379    | Cell-type specific | [30] |
| Mouse    | Bulk tissues | Embryonic | NCX    | FACS; RNA-seq           | http://decon.rc.fas.harvard.edu/       | GSE63482    | Cell-type specific | [31] |
| Mouse    | Bulk tissues | Adult    | HIP    | Genetic labeling; RNA-seq | http://hipposeq.janelia.org            | GSE74985    | Cell-type specific | [32] |
| Mouse    | Bulk tissues | Adult    | Multiple | Genetic labeling; RNA-seq | http://neuroseq.janelia.org            | GSE79238    | Cell-type specific | [33] |
| Mouse    | Single-cell | Postnatal | Brain; SC | SPLiT-seq             | www.ncbi.nlm.nih.gov/geo               | GSE110823   | Spatiotemporal | [34] |
| Mouse    | Single-cell | Adolescence | NS | FACS; 10X Genomics   | http://mousebrain.org                  | SRP135960   | Spatial        | [35] |
| Mouse    | Single-cell | Juvenile; adult | Multiple | scRNA-seq             | http://linnarssonlab.org/oligodendrocytes/ | GSE75330    | Spatiotemporal | [36] |
| Mouse    | Single-cell | Adult    | NCX; HIP CA1 | Fluidigm C1             | http://linnarssonlab.org/cortex        | GSE60361    | Spatial        | [37] |
| Mouse    | Single-cell | Adult    | HPA    | Drop-seq               | www.ncbi.nlm.nih.gov/geo               | GSE87544    | Spatiotemporal | [38] |
| Mouse    | Single-cell | Adult    | RB neurons | FACS; Drop-seq         | https://portals.broadinstitute.org/single_cell | GSE81905    | Spatial        | [39] |

(continued on next page)
| Species       | Sample         | Age          | Region | Method                          | Web link                                                                 | Data access | Annotation       | Refs. |
|---------------|----------------|--------------|--------|---------------------------------|--------------------------------------------------------------------------|-------------|------------------|-------|
| Mouse         | Single-cell    | Adult        | HIP    | Div-seq (nuclei)                | https://portals.broadinstitute.org/single_cell                           | GSE84371    | Spatial          | [40]  |
| Mouse         | Single-cell    | Adult        | V1 (NCX) | FACS; SMARTer                  | http://casestudies.brain-map.org/celltax                                | GSE71585    | Spatial          | [41]  |
| Mouse         | Single-cell    | Adult        | HIP    | SMART-seq                       | www.ncbi.nlm.nih.gov/geo                                                  | GSE71485    | Spatiotemporal   | [42]  |
| Mouse         | Single-cell    | Adult        | STR    | Mic-scRNA-seq; FACS-scRNA-seq   | www.ncbi.nlm.nih.gov/geo                                                  | GSE82187    | Spatial          | [43]  |
| Mouse         | Single-cell    | Adult        | Multiple | Drop-seq                       | http://dropviz.org/                                                      | GSE116470   | Spatial          | [44]  |
| Mouse         | Single-cell    | Adult        | NCX    | FACS; SMART-seq                 | www.ncbi.nlm.nih.gov/geo                                                  | GSE113576   | Spatial          | [45]  |
| Mouse         | Single-cell    | Adult        | HPA    | MERFISH; Drop-seq               | www.ncbi.nlm.nih.gov/geo                                                  | GSE115685   | Spatial          | [46]  |
| Mouse         | Single-cell    | 1–3 M; 21–22 M | Brain  | 10X Genomics                   | http://shiny.baderlab.org/AgingMouseBrain/                               | GSE129788   | Temporal         | [47]  |
| Mouse         | Mixed          | Multiple     | Multiple | Microwell-seq                  | https://figshare.com/s/863e694ad06d5857db4b                               | GSE108097   | Spatiotemporal   | [48]  |
| Rhesus macaque | Bulk tissues   | Lifespan     | Multiple | LMD; microarray                | http://www.blueprintnhpatlas.org/                                        | URL         | Spatiotemporal   | [21]  |
| Rhesus macaque | Mixed          | Lifespan     | Multiple | RNA-seq; 10X Genomics          | http://www.evolution.psychencode.org/                                    | PRJNA448973 | Spatiotemporal   | [50]  |
| Human         | Bulk tissues   | Lifespan     | Multiple | MaD; exon-array                | http://hbatlas.org/                                                      | GSE25219; GSE13344 | Spatiotemporal | [3,5] |
| Human         | Bulk tissues   | Lifespan     | Multiple | LMD; microarray; ISH; RNA-seq   | http://www.brainspan.org/                                                | URL         | Spatiotemporal   | [19]  |
| Human         | Bulk tissue    | Lifespan     | Multiple | Multi-omics                    | http://development.psychencode.org/                                      | phs000755   | Spatiotemporal   | [51]  |
| Human         | Bulk tissues   | Adult        | Multiple | MaD; LMD; microarray; ISH       | http://human.brain-map.org/                                              | URL         | Spatial          | [18]  |
| Human         | Bulk tissues   | Lifespan     | PFC    | Microarray                     | http://braincloud.jhmi.edu                                             | GSE30272    | Temporal         | [6]   |
| Human         | Bulk tissues   | Fetal; juvenile; adult | NCX; HIP | PAN; RNA-seq                   | http://www.brainrnaseq.org/                                             | GSE73721    | Cell-type specific | [52] |
| Human         | Single-cell    | Fetal        | PFC    | SMART-seq2                     | www.ncbi.nlm.nih.gov/geo                                                  | GSE104276   | Temporal         | [53]  |
| Species   | Sample          | Age     | Region | Method              | Web link                                    | Data access | Annotation     | Refs. |
|-----------|-----------------|---------|--------|---------------------|---------------------------------------------|-------------|----------------|-------|
| Human     | Single-cell     | Fetal   | NCX    | STRT-seq            | www.ncbi.nlm.nih.gov/geo                     | GSE103723   | Spatial        | [54]  |
| Human     | Single-cell     | Fetal   | NCX    | Fluidigm C1         | https://cells.ucsc.edu/?ds=cortex-dev       | phs000989.v3 | Spatiotemporal | [55]  |
| Human     | Single-cell     | Fetal; adult | NCX | Fluidigm C1         | www.ncbi.nlm.nih.gov/geo                     | GSE67835    | Spatiotemporal | [56]  |
| Human     | Single-cell     | Fetal; adult | Multiple | Fluidigm C1         | http://www.psychencode.org/                 | URL         | Spatiotemporal | [51]  |
| Human     | Single-cell     | Adult   | NCX    | Fluidigm C1(nuclei) | http://www.scap-t.org/                       | phs000833.v3.p1 | Spatial      | [57]  |
| Human     | Single-cell     | Adult   | Multiple | snDrop-seq          | www.ncbi.nlm.nih.gov/geo                     | GSE97942    | Spatial        | [58]  |
| Human     | Single-cell     | Adult   | MTG    | SMART-seq v4        | http://celltypes.brain-map.org/             | URL         | Spatial        | [59]  |
| Human     | Mixed           | Adult   | Multiple | RNA-seq             | https://www.gtexportal.org                   | URL         | Spatiotemporal | [14]  |
| Drosophila| Single-cell     | Adult   | Brain  | 10x Genomics; Drop-seq | http://scope.aertslab.org                 | GSE107451    | Spatiotemporal | [60]  |
| Drosophila| Single-cell     | Adult   | Midbrain | Drop-seq            | https://www.ncbi.nlm.nih.gov/sra/SRP128516 | SRP128516   | Spatial       | [61]  |
| Zebrafish | Single-cell     | Juvenile | Brain  | GESTALT; Drop-seq   | http://krishna.gs.washington.edu/content/members/aaron/fate_map/harvard_temp_trees/ | GSE105010   | Spatiotemporal | [62]  |
| Zebrafish | Single-cell     | Adult   | Hab    | FACS; 10X Genomics; SMART-seq2 | http://stackjoint.com/zb/daim/ | GSE105115   | Spatial      | [63]  |
| Multiple  | Bulk tissue     | Adult   | Multiple | RNA-seq             | http://www.psychencode.org/                | PRJNA236446 | Species       | [64]  |
| Multiple  | Mixed           | Multiple | Multiple | Multi-omics         | https://www.encodeproject.org               | URL         | Integrative   | [65]  |

Note: Web links for supporting resources are provided when available. Multiple means that samples were obtained from multiple species, tissues, brain regions, or at multiple time points. NCX, neocortex; HPA, hypothalamus; PFC, prefrontal cortex; HIP, hippocampus; Hab, habenular; RB, retinal bipolar; STR, striatum; CNS, central nervous system; SC, spinal cord; BAC, bacterial artificial chromosome; V1, primary visual cortex; MTG, middle temporal gyrus; MiD, microdissection; MaD, macrodissection; FACS, fluorescence-activated cell sorting; LMD, laser microdissection; PAN, immunopanning; TRAP, translating ribosome affinity purification; GESTALT, genome editing of synthetic target arrays for lineage tracing; ISH, in situ hybridization; MERFISH, multiplexed error robust fluorescence in situ hybridization.
transcriptomic architectures. Therefore, a good strategy for studying the functions of a specific gene is to analyze its expression across different developmental stages and/or brain regions. Many transcriptomic analyses of prenatal and postnatal tissues have shown that the intricate principles of human brain development can be revealed by carefully surveying spatial and temporal gene expression [3–7]. For example, Kang et al. used a high-throughput exon array to characterize the spatial and temporal transcriptomes of the human brain [5]. The authors collected more than 1000 postmortem brain samples, covering 16 different regions of the human brain (the hippocampus, striatum, cerebellar cortex, amygdala, mediodorsal nucleus of the thalamus, and 11 neocortical areas). These tissue samples spanned 15 periods from the prenatal stage (5.7 weeks after conception) to the aging stage (82 years old), making this collection one of the most comprehensive collections of brain transcriptome data. This work provides new insights into the spatiotemporally regulated patterns of brain-related genes and their co-expression relationships [5]. The data also show that the predominately regulated stage is the prenatal stage (70.9% genes are spatially differentially expressed, 89.9% genes are temporally differentially expressed, and 70.0% of all expressed genes are regulated in both patterns) [5]. Furthermore, based on the spatial and temporal transcriptome data, researchers can obtain the developmental trajectories of key genes, such as marker genes of different cell types (Figure 2). For brain development and neurodevelopmental disorders, an important problem that needs to be solved is when and where the key genes are expressed and how such expression is disrupted in neurodevelopmental disorders. These gene expression trajectories are valuable resources to dissect the molecular mechanisms underlying the functional specialization of brain regions. More importantly, these trajectories can also contribute to understanding the causes of various neurodevelopmental diseases.

In addition to analyzing spatiotemporal expressional patterns, some groups have considered temporal gene expression dynamics among different brain regions, reflecting the functional specialization of brain regions. Using the mouse brain, Liscovitch and Chechik [66] identified differently expressed genes in multiple brain regions and determined how regional dissimilarities changed over time. In this study, they calculated the dissimilarity for each pair of regions, defined as $1 – \text{Pearson’s correlation coefficient}$. Their results suggest an hourglass pattern in which dissimilarities increase greatly in early prenatal development, decrease to a minimum at birth, and increase again after birth [66]. Notably, they observed a significant postnatal specialization in the mouse cerebellum, and a similar phenomenon was also observed in human brains [66]. In another study related to the human cortex, a temporal hourglass pattern consisting of three major phases was discovered by Pletikos and the colleagues [7]. Prenatal development is the first phase and is characterized by the highest number of differentially expressed genes. The pre-adolescent phase is the second phase, showing less divergent regional gene expression and a more synchronized gene expression pattern. The last phase is adolescence, showing increased regional differences again [7]. This cup-shaped transcriptional divergence pattern is repeatedly observed in the transcriptome of developmental brains from both humans and macaques. Interestingly, the transcriptional divergence between human and macaque brains also exhibits a cup-shaped pattern, as reported in two recent studies [50,51]. These temporal differences in gene expression among different brain regions provide valuable insights into the specialization of brain function.

![Timeline of major human brain cell types based on gene expression trajectories](image)

*Figure 2* Timeline of major human brain cell types based on gene expression trajectories

The X axis shows the developing time and the Y axis represents the relative gene expression level (percentage of maximum). The occurrence and progression of each cell type are reflected by the expression trajectories of the associated genes (data are from the Human Brain Transcriptome project [5]). Based on these trajectories, the prenatal and early infant stage is the most dynamic phase of different cell types. PCW, postconceptional weeks; M, months; Y, years.
Brain-wide coexpression modularity analysis

In the aforementioned study, Kang et al. found that the brain transcriptome tends to organize into co-expression networks that are implicated in distinct biological processes [5]. Generally, genes that share similar expression patterns among samples or time points are defined as co-expressed genes, and there is a high possibility that these genes are involved in similar biological processes [74]. Thus, identifying the co-expressed network based on expression similarity is a powerful method to obtain context-specific functional annotations.

In practice, the key fundamental part of co-expression analysis is how to measure gene expression similarities. Generally, people choose similarity measures according to the purpose of their studies, such as Pearson’s correlations, Spearman’s correlations, partial correlations, mutual information, Euclidian distances, Cosine similarities, and probabilistic measures. The most widely used are correlation-based measurements. For example, NeuroBlast can identify genes with similar three-dimensional spatial expression based on Pearson’s correlations [75], and the Spearman’s correlation coefficient can be used to analyze co-expression gene pairs in the mouse brain [76]. Another example is a recent study that analyzed the co-expression pattern of chromdomain helicase DNA-binding protein 8 (CHD8), a key autism-associated gene [77]. This study showed that CHD8 is widely expressed in both cortical and subcortical structures, although its expression density decreases during development in both human and macaque brains. Moreover, significant enrichment of autism genes was observed in CHD8-correlated genes [77].

Generally, unsupervised clustering and network analyses are appropriate for exploring molecular interactions between a set of genes that may have similar biological functions or be involved in similar pathways. As an unsupervised method, hierarchical clustering is widely used to group genes and samples. Gofflot et al. applied unsupervised hierarchical clustering to explore the expression of nuclear receptors (NRs) in 104 brain regions [78]. They found that anatomical brain structures are organized in three main clusters in favor of the existing taxonomy models of brain, and NRs are clustered in two major groups, with distinct expression patterns [78]. Besides clustering, another approach is constructing a co-expression network in which the nodes are co-expressed genes and the edges represent co-expression relationships of gene pairs with or without weights. The most widely used co-expression network in practice is weighted gene co-expression network analysis (WGCNA), a computational approach to identify network modules based on the topological profiles of a co-expression network [79]. In WGCNA, there is an eigengene for each module, which represents the overall expression of that module, and hub genes can be identified further based on the connectivity of the module members. In this way, the module’s function can be inferred based on the function or enrichment analysis of those hub genes [79]. In neuroscience, this method has been widely applied to construct transcription networks of the mammalian brain. For example, Oldham et al. used WGCNA to compare the network conservation between human and chimpanzee brains [80]. They observed that functional modules of the cerebral cortex are less likely to be conserved during
evolution than those of other brain regions [80]. Moreover, other studies applied WGCNA to identify modules associated with distinct cell types and functions or corresponding to distinct brain regions in the developing and adult brains of mice, rhesus monkeys, and humans [5,17,18,81]. For example, Hawrylycz et al. identified 13 co-expression modules with specific anatomical distributions to characterize the transcriptional variations across the adult human brain [18].

Complex neurological disorders are not caused by a single gene but multiple dysregulated genes, which may converge in the same dysregulated biological processes. With the increasing number of samples taken into consideration, genomewide association studies have linked an increasing number of variants with complex neurological and neuropsychiatric disorders, including autism spectrum disorders (ASDs) [82–87], schizophrenia [88,89], and Alzheimer’s disease [90,91]. In this context, analyzing co-expressed genes with known disease-related genes can provide an avenue to dissect the molecular underpinnings of complex neurological disorders. Ben-David and Shifman used WGCNA to analyze the co-expression relationships of rare and common autism variants and found two modules affected by rare and common variations corresponding to the plasticity of synapses and neurons and the areas of learning and memory, respectively [92]. In another study, Menashe et al. used cosine similarity as a measurement of expression similarity and constructed a co-expression network of autism genes in the mouse brain [93]. These studies demonstrated that autism-related genes are preferentially co-expressed. Moreover, Menashe et al. identified two modules in which autism-related genes are highly connected and overexpressed in a specific brain region, the cerebellar cortex [93]. These abovementioned studies have shown a link between the network of autism-related genes and specific brain regions. Furthermore, researchers can use co-expression analysis to examine when and where specific genes are expressed and how they change during specific biological processes, such as neuron differentiation and maturation, which may provide another view for research into neurodevelopmental disorders. Some studies have been conducted in this field. For example, Parikhshak et al. constructed brain developmental-related WGCNA networks based on the BrainSpan dataset (www.brainspan.org) and mapped ASD-related and intellectual disability-related genes onto different modules [94]. Their results demonstrated that modules significantly enriched in ASD genes are involved in distinct biological functions, such as the regulation of synaptic development [94]. They further found that ASD genes are preferentially located in superficial cortical layers and expressed in glutamatergic projection neurons [94]. In another study, Mahfouz et al. analyzed 455 autism genes to identify their shared pathways [95]. They showed that modules containing large numbers of ASD genes are related to biological processes involving synaptogenesis, apoptosis, and GABAergic neurons [95]. All of these studies demonstrated that the co-expression network is a powerful strategy to reveal the biological functions of disease-risk genes.

**Cell type-specific gene expression analysis**

The brain is the most heterogeneous organ, in which diverse cell types are assembled into distinct but highly connected circuits and regions. Thus, it is possible to identify functional regions and neural cell types based on their transcriptional architecture, not on their morphological and electrophysiological properties. However, in general transcriptome studies, RNAs are extracted from tissue samples and examined *en masse*, which means the characteristics of a specific cell type are missing, further limiting the utility of bulk transcriptome data, since the expression changes that occur in rare cell types may not be detected. Therefore, it is necessary to directly quantify the transcriptome of a specific cell type. In practice, various methods, such as laser-capture microdissection, immunopanning, fluorescence-activated cell sorting, manual cell sorting, and transgenic engineering, are used to identify and isolate specific cell types. A detailed review has compared these methods [96], and another review has provided an overview of existing studies combining these methods and high-throughput transcriptomes to explore cell-specific expression patterns [24].

In addition, great efforts have been made to extract cell type-specific or region-specific patterns from bulk brain transcriptome data. For example, Kirsch et al. proposed a method to detect layer-specific gene expression in the mouse cerebellum [97]. In this work, the authors used a histogram of local binary patterns to represent each gene’s ISH image and predicted the localization based on a two-level classification. First, a classifier based on a support vector machine was trained to identify images of specific layers. Then, genes were classified based on multiple-instance learning [97]. Similarly, Li et al. developed another method (scale-invariant feature transform) to detect cell type-specific genes from ISH images [98]. Zeng et al. applied a deep convolutional neural network to the developing mouse brain [99]. In this work, they used two approaches to extract features from ISH data, *i.e.*, the invariant image feature descriptors method and regularized learning method [99]. All of these studies have demonstrated that computational approaches, particularly feature exacting methods, are helpful for detecting cell type-specific and/or region-specific genes. However, these methods are based on some known marker genes of specific regions, layers, or cell types, and the accuracy of the results needs to be improved. A better choice is characterizing the total transcriptome at the single-cell level and grouping cells into distinct populations based on their transcriptional pattern, as discussed below.

**Single-cell gene expression analysis**

Combined with physical isolation of specific cell types and computational analysis of brain cell pools, the transcriptional atlas of specific cell types can be depicted. However, the accuracy needs to be improved, and heterogeneity still exists. Recently, advances in the isolation of single cells have made it possible to generate the transcriptome of a single cell, and a series of single-cell transcriptome data have been released (Table 1). Researchers can use single-cell RNA-seq (scRNA-seq) to discriminate distinct cell populations, identify new and rare cell types, and trace cell developmental trajectories.

The mammalian brain is viewed as the most complicated organ largely due to the heterogeneity of diverse specialized cell types. Since scRNA-seq can describe the transcriptome from a single cell and the same types of cells are likely to share similar expression patterns, researchers can assign individual cells to distinct cell populations based on the similarity of
### Table 2: scRNA-seq studies revealing multiple cell types in the brain

| Species | Region | No. of total cells | No. of neuronal cells | No. of non-neuronal cells |
|---------|--------|--------------------|-----------------------|--------------------------|
| Zebrafish | Brain; HIP CA1 | 66,000 | 47,822 | 18,178 |
| Mouse | S1; HIP CA1 | 3005 | 1351 | 1654 |
| Mouse | V1 | 1679 | 609 | 648 |
| Mouse | V1; ALM | 3047 | 1265 | 1782 |
| Mouse | Brain; SC | 31,290 | 15,923 | 15,367 |
| Mouse | Ncx | 3127 | 2092 | 1035 |
| Human | Human | 2309 | 701 | 1608 |
| Human | Human | 1560 | 630 | 930 |
| Human | Human | 16,049 | 690 | 914 |
| Human | NCX | 4123 | 1608 | 2515 |
| Human | MTG | 35,289 | 18,045 | 17,244 |

Note: The numbers in parentheses indicate the number of subpopulations. S1: primary somatosensory cortex; V1: primary visual cortex; V1; ALM: anterior lateral motor cortex; SC: spinal cord; NCX: cerebellum; PFC: prefrontal cortex; FC: frontal cortex; CBL: cerebellum; MTG: middle temporal gyrus; SN: spiny neuron; NPC: neural progenitor cell; OPC: oligodendrocyte progenitor cell; HCAI: corpus amastoma area of hippocampus.

the transcriptome, not just based on the expression of marker genes. scRNA-seq has shown great power to explore the heterogeneity of cells in the brain (Table 2). In practice, unsupervised clustering methods, including hierarchical clustering, k-means clustering, principal component analysis, and t-distributed stochastic neighbor embedding, are widely used to identify cell subpopulations. Notably, it is better to apply these clustering methods to differentially expressed genes or highly variable genes. For example, Zeisel et al. measured the transcriptomes of 3005 cells from two regions of the adult mouse brain, that is the primary somatosensory cortex (S1) and hippocampal CA1 region [37]. First, they selected 5000 genes based on a series of strict criteria. Then, they used an algorithm called BackSPIN to cluster genes and cells simultaneously, and identified 47 subclasses of nine major clusters (S1 and CA1 pyramidial neurons, interneurons, oligodendrocytes, astrocytes, microglia, vascular endothelial cells, mural cells, and ependymal cells). Next, Zeisel and colleagues extracted specific markers of each cell population. Some of these markers are well known, while some are novel, such as Gmi11549 specific for S1 pyramidial cells, Spink8 specific for hippocampal pyramidial cells, and Pnoc specific for interneurons [37]. Notably, the general analysis assumes that the cell types are abundant. If the cells are small in number or rare, it is a challenge to discriminate them from the cell populations. To solve this problem, Grun et al. proposed RaceID, which uses transcript counts to identify the rare and abundant cell types in complex cell pools [100]. Overall, RaceID has two major steps. First, k-means clustering is applied to the similarity matrix, and the cluster number is determined from the gap statistic [101]. Then, outlier cells are identified followed by rare cell type identification [100]. Using RaceID, Grun et al. identified a novel marker for enterovascular cells, Reg4 [102].

Another important implication of scRNA-seq is tracking cell trajectory dynamics during a dynamic process, such as neuronal differentiation. However, it is difficult to determine which cell type at time point $n$ progresses to a cell at time point $n + 1$ in scRNA-seq data since the cell is completely consumed. In addition, the cells collected from a sample may not be completely synchronized. Some algorithms have been developed to address these problems, and these algorithms can be generally divided into two classes. These include pseudotime ordering methods, such as diffusion pseudotime (DPT) [103], single-cell topological data analysis (scTDA) [104], Wanderlust [105], Waterfall [42], and Monocle 2 [106], and probabilistic branch models, such as single-cell clustering using bifurcation analysis (SCUBA) [107] and temporal assignment of single cells (TASIC) [108]. In practice, pseudotime ordering methods usually require dimension reduction first, followed by reconstruction of cell trajectories in the lower dimension space, in which graph analysis is usually required, including the minimum spanning trees and principal curves. Recently, Lin et al. proposed a method called continuous-state hidden Markov model (CSHMMs) to infer branching topology and assign cells to the correct branches [109]. In neuroscience, these aforementioned methods are widely used to track cell trajectories during brain development. For example, Zhong et al. performed monocyte pseudotime analysis [110] of human prefrontal cortex development and revealed the following development branches for neural progenitor cells, including two paths to intermediate progenitor cells and one late path to outer radial glia (RG) [53]. In another study, Polioudakis et al. explored the diversity...
and lineage of cell types during human neocortex development. First, they identified 16 distinct cell populations from ~40,000 cells and then performed pseudotime ordering analysis [111]. Moreover, they found ordered transitions during neural progenitor differentiation, such as RG transitioning to intermediate progenitors (IPs) and IP transitioning to newborn migrating neurons [111].

Although scRNA-seq has shown extraordinary superiority in characterizing neuronal cell types and their distributions, some issues should be considered; for example, high variability in levels of the detected transcripts. In the future, advanced methods are required to improve the coverage of the transcriptome and preserve the physiological microenvironment of cells.

Integrative analysis of brain transcriptome and neuroimaging data

In recent years, neuroimaging technology has been greatly developed, providing an unprecedented opportunity to associate molecular variance with macroscopic changes in the brain. Although a large number of brain transcriptome atlases are available, most lack the capability to cover the entire brain, except the Allen Brain Atlas (ABA). ABA is an anatomically comprehensive atlas, comprising 3702 transcriptomes from six adult brains. Importantly, ABA contains MRI data and Montreal Neurological Institute coordinate data [18], allowing researchers to integrate the relationship between spatial variation at the molecular level and observed neuroimaging phenotypes. Recently, many studies have suggested that gene expression is related to the functional connectivity of the brain. In an early study in this field, Goel et al. explored whether there is a relationship between gene expression and anatomical brain regions [112]. They extracted structurally connected regions based on magnetic resonance (MR) diffusion tractography and found no direct relationship between structural connectivity and similar expression patterns at the individual level [112]. In another study, Wang et al. used fractional amplitude of low-frequency fluctuations, a region-specific index, to associate genes with a network called the brain functional activity default mode network, which contains brain regions that exhibit coherent functional magnetic resonance imaging (fMRI) signal fluctuations under the resting state [113]. They found that these related genes are preferentially expressed in neurons and the expression of these genes is downregulated in the brain of autistic patients [114]. In another similar study, Richiardi et al. found that functionally connected regions have similar gene expression patterns via mapping ABA expression data to 14 functional networks [115]. Furthermore, they identified 136 genes driving the relationship that are significantly enriched in ion channels [115]. In addition to investigating the relationship between variations in gene expression and variations in structural/functional connections of the brain, other researchers have shifted their focus to the relationship between structural changes in the brain and gene expression patterns. One example is a study by Whitaker et al., in which the authors explored the underlying mechanism of brain structure maturation during adolescence [116]. Specifically, they collected 297 samples and measured the thickness and myelination of the cortex via MRI. Their results demonstrate a significant association between the shrinkage and myelination of the cortex and the gene expression patterns of dorsoventral areas [116].

Notably, integrative analysis of transcriptome and imaging data often involves many variables, which requires sophisticated data processing. Over the years, various software and tools have been developed to perform such analyses [117–120]. However, the accuracy and consistency of the results obtained are largely affected by the choice of these tools. Recently, a practical guide for key procedures in analyzing HABA data has been proposed to facilitate studies in this field [121]. In the future, developing methodological guidelines to for more accurate results is still necessary.

Limitations and future directions

The resolution of brain ISH data

Although great progress has been made in quantifying gene expression in the brain, several aspects in the field regarding the analysis of the spatial and temporal patterns of the brain must be improved. One key problem is the low resolution of human brain expression imaging data. Although cellular-level resolution is possible in the original ISH data (~1 μm), much higher resolution data are desired for genome-wide data used in three-dimensional (3D) space (~200 μm) [13]. The low resolution poses challenges to investigations into the detailed characteristics of the organization of the brain. Many researchers have attempted to develop new approaches to solve this problem. For example, Ramsden et al. realigned mouse ISH data using nonlinear regression model, which increased the resolution to approximately 10 μm [122]. Using this method, the expression levels of genes that can define the border and layers of medial entorhinal cortex were identified [122]. In the future, more general methods are needed to integrate spatial gene expression data into the standard 3D space.

Expression of non-coding RNAs

Current transcriptome data of the brain mainly focus on the expression of protein-coding genes (mRNAs), whereas the expression features of non-coding RNAs (ncRNAs) are often ignored. In recent years, a series of studies have shown that ncRNAs are of great importance in brain development and neurological disorders [123,124]. In an early study, Mercer et al. analyzed the ISH data from the adult mouse brain and identified a large number of ncRNAs (849 transcripts) [125]; most of these ncRNAs have specific expression profiles in different brain regions and cell types [125]. In another study, Furtuizhos et al. focused on the transcriptional differences among neocortex layers and how these differences change during brain development. As a result, they profiled the temporal transcriptomes of the mouse S1 region, including protein-coding genes and ncRNAs [26]. Similarly, Zias and Rennert explored the roles of microRNAs (miRNAs) during human brain development, and identified miRNAs with spatial- and/or sex-dependent expression and their putative targets [126]. Further functional analysis revealed that these differentially expressed miRNAs are involved in many basic developmental events and neurological disorders [126]. All the
The aforementioned studies demonstrate that necessity of exploring the expression of ncRNAs and their regulatory basis throughout brain development.

Integrative analysis with other neuro-omics data

The rapid development of high-throughput sequencing technologies provides not only transcriptome atlases but also other omics atlases of the brain. Transcriptomes reflect the abundances of RNA, whereas epigenomics data, such as DNA methylation and histone modifications, describe the underlying regulatory mechanisms of gene expression. Additionally, proteomics data provide a more reliable readout of gene expression. With the available isolation of more homogeneous brain samples and great advances in single-cell analysis [127,128], multiple omics data of the brain can be obtained. For example, Illingworth et al. explored the interindividual variability in the human brain methylome and found that compared to other brain regions, the cerebellum has a distinct methylation pattern, which is consistent with the results of transcriptome analysis [129]. In another study, Vermunt et al. identified cis-regulated elements across brain regions, and further analysis of coregulation of the enhancer network revealed hidden cell type and functional information [130]. Furthermore, the psychENCODE project aims to construct a neurobiological epigenetic landscape of adult and developing human brains that are normal or diseased [131]. Based on these high-dimensional multi-omics, it is necessary to develop systematic approaches to conduct integrative analyses. Integrating different multi omics datasets can help us better explore the molecular mechanisms underlying complex phenotypes and neurological disorders.

Conclusion

In recent years, the hypergrowth of next-generation technologies has enabled high-throughput transcriptome measurement of the brain throughout its main developmental stages. The accompanying brain transcriptome atlases are also valuable sources to reveal the molecular architecture of the brain. Computational methods are important to decode these high-dimensional transcriptome data. Combined with transcriptome data and appropriate approaches, the relationships among spatial and temporal gene expression, the complex brain traits, and neurological disorders can be studied. However, with the emergence of new data and the limitations of current data (such as low resolution and the lack of non-coding genes), developing new computational methods remains necessary to overcome limitations and identify new molecular underpinnings of the brain. Furthermore, new systematic approaches are needed to conduct integrative analyses of transcriptomic data and other neuro-omics data.

Competing interests

The authors have declared no competing interests.

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References

[1] Borrell V, Calegari F. Mechanisms of brain evolution: regulation of neural progenitor cell diversity and cell cycle length. Neurosci Res 2014;86:14–24.
[2] Lein ES, Hawrylycz MJ, Ao N, Ayres M, Bensinger A, Bernard A, et al. Genome-wide atlas of gene expression in the adult mouse brain. Nature 2007;445:168–76.
[3] Johnson MB, Kawasaki YI, Mason CE, Krsnik Z, Coppola G, Bogdanovic D, et al. Functional and evolutionary insights into human brain development through global transcriptome analysis. Neuron 2009;62:494–509.
[4] Ip BK, Wappler I, Peters H, Lindsay S, Clowry GJ, Bayatti N. Investigating gradients of gene expression involved in early human cortical development. J Anat 2010;217:300–11.
[5] Kang HJ, Kawasaki YI, Cheng F, Zhu Y, Xu X, Li M, et al. Spatio-temporal transcriptome of the human brain. Nature 2011;478:483–9.
[6] Colantuoni C, Lipska BK, Ye T, Hyde TM, Tao R, Leek JT, et al. Temporal dynamics and genetic control of transcription in the human prefrontal cortex. Nature 2011;478:519–23.
[7] Pletikos M, Sousa AM, Sedmak G, Meyer KA, Zha Y, Cheng F, et al. Temporal specification and bilateral character of human neocortical topographic gene expression. Neuron 2014;81:321–32.
[8] Sarsopoulos I, Marin R, Cardoso-Moreira M, Kaessmann H. Developmental dynamics of IncRNAs across mammalian organs and species. Nature 2019;571:510–4.
[9] Cardoso-Moreira M, Halbert J, Valloton D, Velten B, Chen C, Shao Y, et al. Gene expression across mammalian organ development. Nature 2019;571:505–9.
[10] Gong S, Zheng C, Doughty ML, Losos K, Didkovsky N, Schambra UB, et al. A gene expression atlas of the central nervous system based on bacterial artificial chromosomes. Nature 2003;425:917–25.
[11] Heintze N. Gene expression nervous system atlas (GENSAT). Nat Neurosci 2004;7:483.
[12] Visel A, Thaller C, Eichele G. GenePaint.org: an atlas of gene expression patterns in the mouse embryo. Nucleic Acids Res 2004;32:D552–6.
[13] Mahfouz A, Huisman SMH, Lelieveldt BPF, Reinders MJT. Brain transcriptome atlases: a computational perspective. Brain Struct Funct 2017;222:1557–80.
[14] GTEx Consortium. The Genotype-Tissue Expression (GTEx) project. Nat Genet 2013;45:580–5.
[15] GTEx Consortium; Laboratory Data Analysis & Coordinating Center (LDACC)—Analysis Working Group; Statistical Methods groups—Analysis Working Group; Enhancing GTEx (eGTEx) groups; NIH Common Fund; NIH/NCI; NIH/NHGRI; NIH/NIMH et al. Genetic effects on gene expression across human tissues Nature 2017;550:204–13.
[16] Oldham MC, Konopka G, Iwamoto K, Langfelder P, Kato T, Horvath S, et al. Functional organization of the transcriptome in human brain. Nat Neurosci 2008;11:1271–82.
[17] Thompson CL, Ng L, Menon V, Martinez S, Lee CK, Glattfelder K, et al. A high-resolution spatiotemporal atlas of gene expression of the developing mouse brain. Neuron 2014;83:309–23.

[18] Hawrylycz MJ, Lein ES, Guillozet-Bongaarts AL, Shen EH, Ng L, Miller JA, et al. An anatomically comprehensive atlas of the adult human brain transcriptome. Nature 2012;489:391–9.

[19] Miller JA, Ding SL, Sunkin SM, Smith KA, Ng L, Szafer A, et al. Transcriptional landscape of the prenatal human brain. Nature 2014;508:199–206.

[20] Bernard A, Lubbers LS, Tanis KQ, Luo R, Podtelezhnikov AA, Finney EM, et al. Transcriptional architecture of the primate neocortex. Neuron 2012;73:1083–99.

[21] Bakken TE, Miller JA, Ding SL, Sunkin SM, Smith KA, Ng L, et al. A comprehensive transcriptional map of primate brain development. Nature 2016;535:367–75.

[22] Bakken TE, Miller JA, Luo R, Bernard A, Bennett JL, Lee CK, et al. Spatiotemporal dynamics of the postnatal developing primate brain transcriptome. Hum Mol Genet 2015;24:4327–39.

[23] Pollock JD, Wu DY, Satterlee JS. Molecular neuroanatomy: a generation of progress. Trends Neurosci 2014;37:106–23.

[24] Keil JM, Quilez AL, Kwan KY. Brain transcriptome databases: a user’s guide. J Neurosci 2018;38:2399–412.

[25] Richardson L, Venkataraman S, Stevenson P, Yang Y, Moss J, Graham L, et al. EMAGE mouse embryo spatial gene expression database: 2014 update. Nucleic Acids Res 2014;42:D2835–44.

[26] Furtuzhins S, Li M, Kawasawa YI, Ivic V, Franje D, Singh D, et al. Laminar and temporal expression dynamics of coding and noncoding RNAs in the mouse neocortex. Cell Rep 2014;6:938–50.

[27] Belgard TG, Marques AC, Oliver PL, Aha HO, Sirey TM, Hoerder-Suabedissen A, et al. A transcriptomic atlas of mouse neocortical layers. Neuron 2011;71:605–16.

[28] Cahoy JD, Emery B, Kaushal A, Foo LC, Zamanian JL, Christopherson KS, et al. A transcriptome database for astrocytes, neurons, and oligodendrocytes: a new resource for understanding brain development and function. J Neurosci 2008;28:264–78.

[29] Zhang Y, Chen K, Sloan SA, Bennett ML, Scholze AR, O’Keefe S, et al. An RNA-sequencing and splicing database of glia, neurons, and vascular cells of the cerebral cortex. J Neurosci 2014;34:11929–47.

[30] Doyle JP, Dougherty JD, Heiman M, Schmidt EF, Stevens TR, Ma G, et al. Application of a translational profiling approach for the comparative analysis of CNS cell types. Cell 2008;135:749–62.

[31] Molyneaux BJ, Goff LA, Blettler AC, Chen HH, Hrvatin S, Rinni JL, et al. DeCoN: genome-wide analysis of in vivo transcriptional dynamics during pyramidal neuron fate selection in neocortex. Neuron 2015;85:275–88.

[32] Cambrowski MS, Wang L, Sugino K, Shields BC, Spruston N. Hipposeq: a comprehensive RNA-seq database of gene expression in hippocampal principal neurons. Elife 2016;5:e14997.

[33] Sugino K, Clark E, Schulmann A, Shima Y, Wang LH, Hunt LD, et al. The transcriptional logic of mammalian neuronal diversity. bioRxiv 2017;1:1.

[34] Rosenberg AB, Roco CM, Muscat RA, Kuchina A, Sample P, Yao Z, et al. Single-cell profiling of the developing mouse brain and spinal cord with split-pool barcoding. Science 2018;360:176–82.

[35] Zeisel A, Hochgener H, Lonnerberg P, Johnsson A, Memic F, van der Zwan J, et al. Molecular architecture of the mouse olfactory and auditory systems. Cell 2018;174:999–1014.e22.

[36] Marques S, Zeisel A, Codeluppi S, van Bruggen D, Mendanha Falcão A, Xiao L, et al. Oligodendrocyte heterogeneity in the mouse juvenile and adult central nervous system. Science 2016;352:1326–9.

[37] Zeisel A, Munoz-Manchado AB, Codeluppi S, Lonnerberg P, La Manno G, Jureus A, et al. Cell types in the mouse cortex and hippocampus revealed by single-cell RNA-seq. Science 2015;347:1138–42.

[38] Chen R, Wu X, Jiang L, Zhang Y. Single-cell RNA-seq reveals hypothalamic cell diversity. Cell Rep 2017;18:3227–41.

[39] Shekhar K, Lapan SW, Whitney IE, Tran NM, Macosko EZ, Kowalczuk M, et al. Comprehensive classification of retinal bipolar neurons by single-cell transcriptomics. Cell 2016;166:1308–23.

[40] Habib N, Li Y, Heidenreich M, Sweich L, Avraham-David I, Trombetta JJ, et al. Div-Seq: Single-nucleus RNA-Seq reveals dynamics of rare adult newborn neurons. Science 2016;353:925–8.

[41] Tasic B, Menon V, Nguyen TN, Kim TK, Jarlsk T, Yao Z, et al. Adult mouse cortical cell taxonomy revealed by single cell transcriptomics. Nat Neurosci 2016;19:335–46.

[42] Shin J, Berg DA, Zhu Y, Shin JY, Song J, Bonaguidi MA, et al. Single-cell RNA-seq with waterfalls reveals molecular cascades underlying adult neurogenesis. Cell Stem Cell 2015;17:360–72.

[43] Gokce O, Stanley GM, Treutlein B, Neff NF, Camp JD, Qin Y, et al. Cellular taxonomy of the mouse striatum as revealed by single-cell RNA-seq. Cell Rep 2016;16:1126–37.

[44] Saunders A, Macosko EZ, Wyssker A, Goldman M, Krienen FM, de Rivera H, et al. Molecular diversity and specializations among the cells of the adult mouse brain. Cell 2018;174:1015–30.

[45] Tasic B, Yao Z, Grayburg LT, Smith KA, Nguyen TN, Bertagnolli D, et al. Shared and distinct transcriptomic cell types across neocortical areas. Nature 2018;563:72–8.

[46] Moffitt JR, Bambah-Mukku D, Eichhorn SW, Vaughan E, Shekhar K, Perez JD, et al. Molecular, spatial and functional single-cell profiling of the hypothalamic preoptic region. Science 2018;362:eaa5524.

[47] Ximerakis M, Lipnick SL, Innes BT, Simmons SK, Adiconis X, Dionne D, et al. Single-cell transcriptomic profiling of the aging mouse brain. Nat Neurosci 2019;22:1696–708.

[48] Han X, Wang R, Zhou Y, Fei L, Sun H, Lai S, et al. Mapping the mouse cell atlas by microwell-seq. Cell 2018;173:1307–11.

[49] The Tabula Muris Consortium. Single-cell transcriptomics of 20 mouse organs creates a Tabula Muris. Nature 2018;562:367–72.

[50] Zhu Y, Sousa AMM, Gao T, Skarica M, Li M, Santpere G, et al. Spatiotemporal transcriptomic divergence across human and macaque brain development. Science 2018;362:eaa0877.

[51] Li M, Santpere G, Imamura Kawasawa Y, Evgrafov OV, Gulden FO, Pocheveddy S, et al. Integrative functional genomic analysis of human brain development and neuropsychiatric risks. Science 2018;362:eaa7615.

[52] Zhang Y, Sloan SA, Clarke LE, Caneda C, Plaza CA, Blumen- thal PD, et al. Purification and characterization of progenitor and mature human astrocytes reveals transcriptional and functional differences with mouse. Neuron 2016;90:37–53.

[53] Zhong S, Zhang S, Fan X, Wu Q, Yan L, Dong J, et al. A single-cell RNA-seq survey of the developmental landscape of the human prefrontal cortex. Nature 2018;555:524–8.

[54] Fan X, Dong J, Zhong S, Wei Y, Wu Q, Yan L, et al. Spatial transcriptomic survey of human embryonic cortical by single-cell RNA-seq analysis. Cell Res 2018;28:730–45.

[55] Nowakowski TJ, Bhaduri A, Pollen AA, Alvarado B, Mestaje-Radi MA, Di Lullo L, et al. Spatiotemporal gene expression trajectories reveal developmental hierarchies of the human cortex. Science 2017;358:1583–38.

[56] Darmanis S, Sloan SA, Zhang Y, Enge M, Caneda C, Shuer LM, et al. A survey of cell types in the immune system reveals rare and novel cell types. Cell 2018;174:999–1014.e22.

[57] Lake BB, Ai R, Kaeser GE, Salathia NS, Yung YC, Liu R, et al. Neuronal subtypes and diversity revealed by single-nucleus RNA sequencing of the human brain. Science 2016;352:1586–90.
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[58] Lake BB, Chen S, Sos BC, Fan J, Kaeser GE, Yung YC, et al. Integrative single-cell analysis of transcriptional and epigenetic states in the human adult brain. Nat Biotechnol 2018;36:70–80.

[59] Hodge RD, Bakken TE, Miller JA, Smith KA, Barkan ER, Graybuck LT, et al. Conserved cell types with divergent features in human versus mouse cortex. Nature 2019;573:61–8.

[60] Dave K, Janssens J, Koldere D, De Waegeneer M, Pech U, Kroft L, et al. A single-cell transcriptome atlas of the aging Drosophila brain. Cell 2018;174:982–98.

[61] Croset V, Treiber CD, Waddell S. Cellular diversity in the Drosophila midbrain revealed by single-cell transcriptomics. Elife 2018;7.

[62] Raj B, Wagner DE, McKenna A, Pandey S, Klein AM, DeAngelis J, et al. Single-cell RNA-seq reveals cell types and cell types in the vertebrate brain. Nat Biotechnol 2018;36:442–50.

[63] Pandey S, Shekhar K, Regev A, Schier AF. Comprehensive identification and spatial mapping of habenular neuronal types using single-cell RNA-seq. Curr Biol 2018;28:1032–65.

[64] Sousa AMM, Zhu Y, Raghanti MA, Kitchen RR, Onorati M, Tehenkamp ATN, et al. Molecular and cellular reorganization of neural circuits according to their function in the brain. Science 2017;358:1027–1032.

[65] ENCODE Project Consortium. An integrated encyclopedia of DNA elements in the human genome. Nature 2012;489:57–74.

[66] Liscovitch N, Chechik G. Specialization of gene expression during mouse brain development. PLoS Comput Biol 2013;9: e1003185.

[67] Masserottti G, Gascon S, Gotz M. Direct neuronal reprogramming: learning from and for development. Development 2016;143:2949–510.

[68] Rakic P. Evolution of the neocortex: a perspective from developmental biology. Nat Rev Neurosci 2009;10:724–35.

[69] O’Rahilly R, Muller F. Significant features in the early prenatal development of the human brain. Ann Anat 2008;190:105–18.

[70] Roth G, Dicke U. Evolution of the brain and intelligence in primates. Prog Brain Res 2012;195:413–30.

[71] Miller JA, Horvath S, Geschwind DH. Divergence of human and mouse brain transcriptome highlights Alzheimer disease pathways. Proc Natl Acad Sci U S A 2010;107:12698–703.

[72] Strand AD, Aragaki AK, Baquet ZC, Hodges A, Cunningham P, Holmans P, et al. Conservation of regional gene expression in mouse and human brain. PLoS Genet 2007;3:e59.

[73] Xiao PP, Zhao XM. The exploration of functional divergence between human and macaque brains based on gene networks. IEEE Inter Conf Bioinform Biomed (BIBM) 2016;2016:1879–84.

[74] Stuart JM, Segal E, Koller D, Kim SK. A gene-coexpression network for global discovery of conserved genetic modules. Science 2003;302:249–53.

[75] Havrylycz M, Ng L, Page D, Morris J, Lau C, Faber S, et al. Multi-scale correlation structure of gene expression in the brain. Neural Netw 2011;24:933–42.

[76] French L, Tan PP, Pavlidis P. Large-scale analysis of gene expression and connectivity in the rodent brain: insights through data integration. Front Neuroinform 2011;5:12.

[77] Bernier R, Golzio C, Xiong B, Stessman HA, Coe BP, Penn O, et al. Disruptive CHD8 mutations define a subtype of autism early in development. Cell 2014;158:263–76.

[78] Gofflot F, Chartoire N, Vasseur L, Heikkinen S, Dembele D, Le Merrer J, et al. Systematic gene expression mapping clusters nuclear receptors according to their function in the brain. Cell 2007;131:405–18.

[79] Zhang B, Horvath S. A general framework for weighted gene co-expression network analysis. Stat Appl Genet Mol Biol 2005;4:1544–6115.

[80] Oldham MC, Horvath S, Geschwind DH. Conservation and evolution of gene coexpression networks in human and chimpanzee brains. Proc Natl Acad Sci U S A 2006;103:17973–8.

[81] Miller JA, Nathanson J, Franjc D, Shim S, Dalley RA, Shapouri S, et al. Conserved molecular signatures of neurogenesis in the hippocampal subgranular zone of rodents and primates. Development 2013;140:4633–44.

[82] Iossifov I, Ronemus M, Levy D, Wang Z, Hakker I, Rosenbaum J, et al. De novo gene disruptions in children on the autism spectrum. Neuron 2012;74:285–99.

[83] Neale BM, Koy Y, Liu L, Ma’ayan A, Samocha KE, Sabo A, et al. Patterns and rates of exonic de novo mutations in autism spectrum disorders. Nature 2012;485:242–5.

[84] O’Roak BJ, Vives L, Girirajan S, Karakoc E, Krumm N, Cole BP, et al. Sporadic autism exomes reveal a highly interconnected protein network of de novo mutations. Nature 2012;485:246–50.

[85] Sanders SJ, Murtha MT, Gupta AR, Murdoch JD, Raubeson MJ, Willsey AJ, et al. De novo mutations revealed by whole-exome sequencing are strongly associated with autism. Nature 2012;485:237–41.

[86] De Rubeis S, He X, Goldberg AP, Poulten CS, Samocha K, Ciek AE, et al. Synthetic, transcriptional and chromatin genes disrupted in autism. Nature 2014;515:209–15.

[87] Dong S, Walker MF, Carriero NJ, DiCola M, Willsey AJ, Ye AY, et al. De novo insertions and deletions of predominantly paternal origin are associated with autism spectrum disorder. Cell Rep 2014;9:16–23.

[88] Fromer M, Pocklington AJ, Kavanagh DH, Williams HJ, Dwyer S, Gormley P, et al. De novo mutations in schizophrenia implicate synaptic networks. Nature 2014;506:179–84.

[89] Schizophrenia Working Group of the Psychiatric Genomics Consortium. Biological insights from 108 schizophrenia-associated genetic loci. Nature 2014;511:421–7.

[90] Betens K, Sleegers K, Van Broeckhoven C. Genetic insights in Alzheimer’s disease. Lancet Neurol 2013;12:92–104.

[91] Zhang B, Gaiteri C, Bodea LG, Wang Z, McElwee J, Podtelezhnikov AA, et al. Integrated systems approach identifies genetic nodes and networks in late-onset Alzheimer’s disease. Cell 2013;153:707–20.

[92] Ben-David E, Shifman S. Networks of neuronal genes affected by common and rare variants in autism spectrum disorders. PLoS Genet 2012;8:e1002556.

[93] Menashe I, Grange P, Larsen EC, Banerjee-Basu S, Mitra PP. Co-expression profiling of autism genes in the mouse brain. PLoS Comput Biol 2013;9:e1003128.

[94] Parikshak NN, Luo R, Zhang A, Won H, Lowe JK, Chandran V, et al. Integrative functional genomic analyses implicate specific molecular pathways and circuits in autism. Cell 2013;155:1008–21.

[95] Mahfouz A, Ziants MN, Rennert OM, Lelieveldt BP, Reinders MJ. Shared pathways among autism candidate genes determined by co-expression network analysis of the developing human brain transcriptome. J Mol Neurosci 2015;57:580–94.

[96] Okaty BW, Sugino K, Nelson SB. Cell type-specific transcriptomics in the brain. J Neurosci 2011;31:6939–43.

[97] Kirsch L, Liscovitch N, Chechik G. Localizing genes to cerebellar layers by classifying ISH images. PLoS Comput Biol 2012;8:e1002790.

[98] Li R, Zhang W, Ji S. Automated identification of cell-type-specific genes in the mouse brain by image computing of expression patterns. BMC Bioinformatics 2014;15:209.

[99] Zeng T, Li R, Mukkamala R, Ye J, Ji S. Deep convolutional neural networks for annotating gene expression patterns in the mouse brain. BMC Bioinformatics 2015;16:147.

[100] Duman D, Whinney A, Kestler L, Wibrands K, Basak O, Sasaki N, et al. Single-cell messenger RNA sequencing reveals rare intestinal cell types. Nature 2015;525:251–5.

[101] Tibshirani R, Walther G, Hastie T. Estimating the number of clusters in a data set via the gap statistic. J R Statist Soc B Part 2 2001;63:411–23.
Engelstoft MS, Egerod KL, Lund ML, Schwartz TW. Enteroneuroendocrine cell types revisited. Curr Opin Pharmacol 2013;13:912–9.

Haghighi L, Buttner M, Wolf FA, Buettner F, Theis FJ. Diffusion pseudotime robustly reconstructs lineage branching. Nat Methods 2016;13:845–8.

Rizvi AH, Camara PG, Kandror EK, Roberts TJ, Schieren I, Maniatis T, et al. Single-cell topological RNA-seq analysis reveals insights into cellular differentiation and development. Nat Biotechnol 2017;35:551–60.

Bendall SC, Davis KL, Amir el AD, Tadmor MD, Simonds EF, Chen TJ, et al. Single-cell trajectory detection uncovers progression and regulatory coordination in human B cell development. Cell 2014;157:714–25.

Quo X, Mao Q, Tang Y, Wang L, Chawla R, Pliner HA, et al. Reversed graph embedding resolves complex single-cell trajectories. Nat Methods 2017;14:979–82.

Marco E, Karp RL, Guo G, Robson P, Hart AH, Trippa L, et al. Bifurcation analysis of single-cell gene expression data reveals epigenetic landscape. Proc Natl Acad Sci U S A 2014;111: E5643–50.

Rashid S, Kotton DN, Bar-Joseph Z. TASIC: determining branching models from time series single cell data. Bioinformatics 2017;33:2504–12.

Lin C, Bar-Joseph Z. Continuous state HMMs for modeling time series single cell RNA-seq data. Bioinformatics 2019. https://doi.org/10.1093/bioinformatics/btz296.

Trappnell C, Caciarielli D, Grimsby J, Pokharel P, Li S, Morse M, et al. The dynamics and regulators of cell fate decisions are revealed by pseudotemporal ordering of single cells. Nat Biotechnol 2014;32:381–6.

Polioudakis D, Torre-Ubieta L, Langerman J, Elkins AG, Simonds EF, Chen TJ, et al. Single-cell trajectory detection uncovers progression and regulatory coordination in human B cell development. Cell 2014;157:714–25.

Rashid S, Kotton DN, Bar-Joseph Z. TASIC: determining branching models from time series single cell data. Bioinformatics 2017;33:2504–12.

Lin C, Bar-Joseph Z. Continuous state HMMs for modeling time series single cell RNA-seq data. Bioinformatics 2019. https://doi.org/10.1093/bioinformatics/btz296.

Trappnell C, Caciarielli D, Grimsby J, Pokharel P, Li S, Morse M, et al. The dynamics and regulators of cell fate decisions are revealed by pseudotemporal ordering of single cells. Nat Biotechnol 2014;32:381–6.

Polioudakis D, Torre-Ubieta L, Langerman J, Elkins AG, Stein JL, Vuong CK, et al. A single cell transcriptomic analysis of human neocortical development. bioRxiv 2018;40185.

Goel P, Kuceyeski A, LoCastro E, Raj A. Spatial patterns of genome-wide expression profiles reflect anatomy and fiber connectivity architecture of healthy human brain. Hum Brain Mapp 2014;35:4204–18.

Buckner RL, Andrews-Hanna JR, Schacter DL. The brain’s default network: anatomy, function, and relevance to disease. Ann N Y Acad Sci 2008;1124:1–38.

Wang GZ, Belgard TG, Mao D, Chen L, Berto S, Preuss TM, et al. Correspondence between resting-state activity and brain gene expression. Neuron 2015;88:659–66.

Richardi J, Altman A, Milazzo AC, Chang C, Chakravarty MM, Banaschewski T, et al. Correlated gene expression supports synchronous activity in brain networks. Science 2015;348:1241–4.

Whitaker KJ, Vertes PE, Romero-Garcia R, Vasa F, Moutoussis M, Prabhhu G, et al. Adolescence is associated with genetically patterned consolidation of the hubs of the human brain connectome. Proc Natl Acad Sci U S A 2016;113:9105–10.

French L, Paus T. A FreeSurfer view of the cortical transcriptome generated from the Allen Human Brain Atlas. Front Neurosci 2015;9:323.

Rizzo G, Veronese M, Expert P, Turkheimer FE, Bertoldo A. MENGA: a new comprehensive tool for the integration of neuroimaging data and the Allen Human Brain Transcriptome Atlas. PLoS One 2016;11:e0148744.

Gorgolewski KJ, Varoquaux G, Rivera G, Schwarz Y, Ghosh SS, Maumet C, et al. NeuroVault.org: a web-based repository for collecting and sharing unthresholded statistical maps of the human brain. Front Neuroinform 2015;9:8.

Rittman T, Rittman M, Azevedo T. Maybrain software package. https://github.com/RittmanResearch/maybrain.

Armatkevicuite A, Fulcher BD, Fornito A. A practical guide to linking brain-wide gene expression and neuroimaging data. Neuroimage 2019;189:353–67.

Ramsden HL, Surnelli G, McDonagh SG, Nolan MF. Laminar and dorsoventral molecular organization of the medial entorhinal cortex revealed by large-scale anatomical analysis of gene expression. PLoS Comput Biol 2013;11:e1004032.

Ponjavic J, Oliver PL, Lunter G, Ponjing CP. Genomic and transcriptional co-localization of protein-coding and long non-coding RNA pairs in the developing brain. PLoS Genet 2009;5:e1000617.

Qureshi IA, Mehler MF. Emerging roles of non-coding RNAs in brain evolution, development, plasticity and disease. Nat Rev Neurosci 2012;13:528–41.

Mercer TR, Dinger ME, Sunkin SM, Mehler MF,Mattick JS. Specific expression of long non-coding RNAs in the mouse brain. Proc Natl Acad Sci U S A 2008;105:716–21.

Ziats MN, Rennert OM. Identification of differentially expressed microRNAs across the developing human brain. Mol Psychiatry 2014;19:848–52.

Shin J, Ming GL, Song H. Decoding neural transcriptomes and epigenomes via high-throughput sequencing. Nat Neurosci 2014;17:1463–75.

Maze I, Shen L, Zhang B, Garcia BA, Shao N, Mitchell A, et al. Analytical tools and current challenges in the modern era of neuroepigenomics. Nat Neurosci 2014;17:1476–90.

Illingworth RS, Gruenewald-Schneider U, De Sousa D, Webb S, Merusi C, Kerr AR, et al. Inter-individual variability contrasts with regional homogeneity in the human brain DNA methylome. Nucleic Acids Res 2015;43:732–44.

Vermunt MW, Reinkin P, Korving J, de Bruijn E, Creyghton PM, Basak O, et al. Large-scale identification of co-regulated enhancer networks in the adult human brain. Cell Rep 2014;9:767–79.

PsychENCODE Consortium, Akbarian S, Liu C, Knowles JA, Vencevarino FM, Farnham PJ, et al. The PsychENCODE project. Nat Neurosci 2015;18:1707–12.