In vivo development and single-cell transcriptome profiling of human brain organoids

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Abstract

Objectives: Human brain organoids can provide not only promising models for physiological and pathological neurogenesis but also potential therapies in neurological diseases. However, technical issues such as surgical lesions due to transplantation still limit their applications.

Materials and methods: Instead of applying mature organoids, we innovatively developed human brain organoids in vivo by injecting small premature organoids into corpus striatum of adult SCID mice. Two months after injection, single-cell transcriptome analysis was performed on 6131 GFP-labeled human cells from transplanted mouse brains.

Results: Eight subsets of cells (including neuronal cells expressing striatal markers) were identified in these in vivo developed organoids (IVD-organoids) by unbiased clustering. Compared with in vitro cultured human cortical organoids, we found that IVD-organoids developed more supporting cells including pericyte-like and choroid plexus cells, which are important for maintaining organoid homeostasis. Furthermore, IVD-organoids showed lower levels of cellular stress and apoptosis.

Conclusions: Our study thus provides a novel method to generate human brain organoids, which is promising in various applications of disease models and therapies.
1 | INTRODUCTION

The human brain is a highly organized structure that consists of various cell types including neurons, glia, and vascular cells. It is challenging to model human brain development because it requires coordinated regulation of different biochemical and physical events. Human brain organoids are stem-cell-derived three-dimensional cultures that recapitulate the cellular and architectural features of a human brain, making them a promising source for transplantation therapies. Indeed, previous studies have demonstrated that upon transplantation into adult mouse cortex, human cerebral organoids could exhibit progressive neuronal maturation and function, as well as robust host-mediated vascularization. These findings have brightened the prospect of using brain organoid transplantation for the treatment of neurological disorders.

Previous studies have evaluated the integration and therapeutic effects of in vitro human cerebral organoids (hCOs) by transplanting them into the rodent cortex, but challenges still remain in the current brain organoid transplantation techniques which hinder further clinical applications. Mature cerebral organoids usually form large aggregates (up to 3–5 mm in diameter) that require larger surgical cavities for organoid transplantation, which may bring more surgical lesions to healthy tissues along the entire route of delivery, especially when transplanting organoids into deep brain regions. Thus, the surgical injury itself may lead to loss of brain function. Therefore, there is an urgent need for new transplantation techniques that minimize injuries to the host brain.

Here instead of transplanting mature brain organoids, we injected small premature organoids into mouse brain and left them growing and differentiating to generate IVD-organoids. Using single-cell transcriptome analysis, we identified neuronal cells expressing striatal markers in IVD-organoids. Compared with in vitro cultured human cerebral organoids, we also found more pericyte-like and mature choroid plexus (ChP) cells in IVD-organoids, which were beneficial for maintaining brain homeostasis. Importantly, IVD-organoids showed reduced levels of cellular stress and apoptosis. Our observations thus suggest promising applications of IVD-organoids in cell therapies for neurological diseases without the risk of surgical lesions.

2 | MATERIALS AND METHODS

2.1 | Mice

The SCID mice were purchased from Shanghai LinChang Biotech. 4- to 5-week-old female mice were used for transplantation. All animal experiments were approved by the Institutional Animal Care and Use Committee (IACUC) of Institute of Biochemistry and Cell Biology, Shanghai Institutes for Biological Sciences, Chinese Academy of Sciences. All animals were housed in specific-pathogen-free animal facility and were given ad libitum access to food and water.

2.2 | hPSC culture

The human induced pluripotent stem cells (hPSC) line was generated from a 66-year-old healthy female and characterized by iXCells Biotechnologies (catalog no. 30HU-002). Cells were cultured on Matrigel (BD Biosciences) coated dishes in mTeSR1 medium (Stem Cell Technologies). Cultures were passaged every 5–7 days with Gentle Cell Dissociation Reagent (Stem Cell Technologies). All cells were tested negative for mycoplasma.

2.3 | Generation of premature organoids for transplantation and mature cerebral organoids

STEMdiff™ Cerebral Organoid Kit (Stem Cell Technologies) was used for generation of premature organoids. Briefly, three different batches of hPSCs were first dissociated with Gentle Cell Dissociation Reagent (Stem Cell Technologies) and transduced by incubating with lentivirus (FUGW-GFP) for 1 h in mTeSR1 medium containing 4 μg/ml polybrene. These GFP-labeled hPSCs were then resuspended in EB Seeding Medium, and 9,000 cells were added into a well of 96-well round-bottom ultra-low attachment plate. On day 5, EBs were transferred into a 24-well ultra-low attachment plate containing Induction Medium and cultured for an additional 2 days before being used for transplantation. To generate mature cerebral organoids, these premature organoids were embedded in Matrigel, transferred to Expansion Medium, and cultured for 3 days. On day 10, organoids were transferred to Maturation Medium and cultured on an orbital shaker. The medium was changed every 3 days during the maturation process.

2.4 | Stereotaxic surgery

To perform injections, SCID mice were anesthetized with 125 mg/kg tribromoethanol (MedChemExpress) and placed in a stereotaxic apparatus. Holes were drilled on the skull to expose the dura, and three premature organoids or dissociated cells from equal number of premature organoids were injected into the striatum using a 5-μl Hamilton syringe with a 22-gauge needle. The coordinates were: AP: +0.5 mm, ML: +2 mm for premature organoids and −2 mm for dissociated cells. After injection, the needle was left in place for 2 min before withdrawal to avoid reflux.

2.5 | Immunofluorescence staining

For cultured organoids, residual Matrigel was removed by washing with PBS. Organoids and mouse brains were fixed with 4% PFA in PBS overnight at 4°C and submerged in 30% sucrose for 2–3 days until samples sank. Then, organoids or mouse brains were embedded in OCT and cryosectioned at 20 or 30 μm, respectively. Antigen retrieval was performed with Citrate Antigen Retrieval Solution (Beyotime) at 95°C for 20 min. Slides were incubated

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2.5 mm, ML: −2 mm for

2.4 mm, respectively.
overnight at 4°C with primary antibodies diluted in PBS containing 0.3% Triton X-100 and 3% donkey serum. After three washes, slides were incubated with secondary antibodies at room temperature for 1 h. Primary antibodies used were as follows: MAP2 (rabbit, Millipore AB5622, 1:500), GFAP (rabbit, DAKO 2033401, 1:1000), STEM121 (mouse, Takara Y40410, 1:500), Iba1 (rabbit, Wako 019-19741, 1:500), CD31 (rabbit, Beyotime AF0099, 1:200), PDGFβ (rat, Thermofisher 14-1402-81, 1:500), Zbtb20 (rabbit, Proteintech 23987-1-AP, 1:200), TTR (mouse, R&D MAB7505, 1:500), NESTIN (mouse, Millipore MAB5326, 1:500), DCX (goat, Santacruz sc-8066, 1:200), LH2X (rabbit, Abcam ab184337, 1:500), TBR1 (rabbit, Abcam ab31940, 1:500), Brychury (rabbit, Abcam ab20680, 1:500), NGN2 (rabbit, Cell Signaling Technology 13144, 1:200), MASH1 (rabbit, Abcam ab211327, 1:100), FoxG1 (rabbit, Abcam ab18259, 1:100), CA12 (rabbit, Abcam ab195233, 1:100), and cleaved caspase 3 (rabbit, Cell Signaling Technology 9661, 1:200). Secondary antibodies used were as follows: donkey anti-rabbit cy3 (Jackson ImmunoResearch 711-165-152, 1:1000), donkey anti-mouse Alexa 647 (Jackson ImmunoResearch A31571, 1:1000), donkey anti-mouse cy3 (Molecular Probes 715-165-150, 1:1000), donkey anti-rat cy3 (Jackson ImmunoResearch 712-165-150, 1:1000), and donkey anti-goat cy3 (Jackson ImmunoResearch 705-165-147, 1:1000). All images were scanned using Olympus FV10i confocal microscope.

2.6 | scRNA-seq library preparation and sequencing

Two months after injection of premature organoids, mice were anesthetized with chloral hydrate and perfused with cold PBS. Brain was immediately removed from the skull and coronally sectioned with an ice-cold brain slicer matrix. GFP-positive regions in brain slices were rapidly dissected out using a fluorescence stereo microscope, digested for 30 min at 37°C with enzymatic dissociation medium (1 U/ml dispase, 2.5 U/ml papain, and 250 U/ml DNase I; all from Worthington), and mechanically triturated. The cell suspension was passed through a 40 μm cell strainer, and debris was removed by 22% Percoll (Sigma-Aldrich) density gradient centrifugation. The 3’ scRNA-seq library construction was performed on a Chromium Controller (10x Genomics) with the Single Cell 3’ v2 kit following the manufacturer’s instructions. The target capture rate was 8,000 cells per sample. Final libraries were sequenced on an HiSeq X Ten system (Illumina) as paired-end 150-bp reads.

2.7 | scRNA-seq data processing

Sequencing reads were aligned to mixed reference genome containing human (hg38), mouse (mm10), and GFP sequences using STAR in Cell Ranger 3.1.0 with default parameters. After removing poorly mapped reads and PCR duplicates, reads that were uniquely mapped to human transcriptome or GFP sequence and contained valid cell barcodes and unique molecular identifiers ( UMIs) were used to generate the gene-barcode matrix. Cells with less than 200 detected genes, or more than 20% mitochondrial transcripts, or no GFP reads were discarded. After filtering, gene expression in each cell was calculated as transcripts per 10,000 transcripts (TP10K), representing the fraction of each gene’s UMI count with respect to total UMIs in the cell and multiplied by 10,000. Only genes expressed in at least three cells were retained. The resulting digital expression matrix was log-transformed before downstream analyses with Seurat (v3.2.0). The top 2,000 highly variable genes were used to perform principal component analysis (PCA). Dimension reduction was implemented by uniform manifold approximation and projection (UMAP) using 1st to 25th principal components (PCs). Cell clusters were identified based on the Louvain clustering. Differentially expressed genes (DEGs) in each cluster were identified using the “FindAllMarkers” function with default parameters. Cell-type annotation was performed similarly to previous reports.

2.8 | Integration analysis with published organoid scRNA-seq data

Integration analysis of IVD-organoids and hCOs (GSE134049) was performed using the Seurat 3 integration pipeline. Canonical correlation (CC) analysis with the top 10 CC components was used to reduce the batch effects between two datasets. DEGs between two datasets in the corresponding clusters were identified using the “FindMarkers” function with the MAST algorithm. GO term analysis of DEGs was performed using the clusterProfiler package. PCA analysis was performed on ChP cells in IVD-organoids, together with mature ChP cells, immature ChP cells, neuronal cells, and stroma cells in ChP organoids (GSE150903). To estimate the levels of apoptosis (HALLMARK: M5902), glycolysis (GO: 0061621), hypoxia (GO: 0001666), and endoplasmic reticulum (ER) stress pathways (GO: 0034976), average expression levels of genes in these pathways were calculated. Two-sided Wilcoxon’s rank-sum test was used to determine the statistical significance.

2.9 | Statistical analysis

Statistical analyses were performed using GraphPad Prism 6 and R (3.6.1). Two-tailed unpaired Student’s t-test was used unless stated otherwise. Data were presented as a mean ± s.e.m. (standard error of mean). p-values in multiple tests were adjusted by false discovery rate (FDR) Benjamini–Hochberg method, and FDR <0.05 was defined as statistical significance.

3 | RESULTS

3.1 | In vivo development of human brain organoids in mouse brain

To trace the premature organoids after transplantation, we first performed lentiviral infection to obtain GFP-expressing hPSCs, which were then differentiated into premature organoids. After
7 days of in vitro culture, these premature organoids were transplanted into corpus striatum of immune-deficient mice by stereoinjection (n = 3). As a control, we also injected dissociated cells from equal number of premature organoids into the corpus striatum of the opposite side of the same mice (Figure 1A). Day-7 premature organoids were also analyzed by immunostaining for dorsal telencephalic marker NGN2, dorsal cortical marker LHX2, ventral telencephalic markers MASH1 and NKK2.1, preplate marker TBR1, forebrain marker FoxG1, neural stem/progenitor cell marker NESTIN, immature neuronal marker DCX, choroid plexus marker TTR, pericyte marker PDGFRβ1, and mesodermal marker Brachyury. We only observed NESTIN-positive cells, indicating the emergence of neural stem/progenitor cells instead of more differentiated cells in these day-7 premature organoids (Figure S1A). Two months after injection, we dissected the recipient mice and found that the GFP-labeled IVD-organoids exhibited robust survival and ventricle zone (VZ)-like structures, near which large cavities surrounded by GFP-positive ChP epithelium-like cells could often be found (Figure 1B). Such structures were not observed in the corpus striatum of the opposite side where dissociated cells were injected, suggesting that the organoid-like structures were specific to IVD-organoids (Figure 1C and Figure S1B). Nevertheless, neuronal and astrocytic differentiation could be robustly observed in both IVD-organoids and dissociated cells as indicated by neuron marker MAP2 and astrocyte marker GFAP (Figure 1D, E and Figure S1C, D). Interestingly, although IBA1-expressing microglia cells were also found in IVD-organoids and dissociated cells, they did not express GFP or human-specific marker STEM121, suggesting these microglia cells were migrated from host tissues (Figure 1F and Figure S1E). GFP-negative CD31-expressing microvasculature-like structures of host origin were also observed in the injection sites, indicating the formation of vascular network which may contribute to the survival and long-term development of IVD-organoids and dissociated cells (Figure 1G and Figure S1F).

**FIGURE 1** In vivo development of human brain organoids in mouse brain. (A) Experiment procedure for premature organoid generation and injection into mouse corpus striatum. (B) IVD-organoids were immunostained for GFP and DAPI. Left, coronal section showing IVD-organoids survived in mouse corpus striatum. Scale bar, 1000 μm. Middle, enlarged image showing the ventricle zone-like structure. Scale bars, 50 μm. Right, enlarged image showing the choroid plexus epithelium-like structure. Scale bar, 100 μm. (C) Comparation of grafted cells (left) and IVD-organoids (right). Scale bars, 500 μm. (D) Immunostaining of MAP2 in IVD-organoids. Scale bars, 20 μm. (E) Immunostaining of GFAP and STEM121 in IVD-organoids. Scale bars, 50 μm and 10 μm (high-magnification images). (F) Immunostaining of IBA1 in IVD-organoids. Scale bars, 50 μm. (G) Immunostaining of vascular marker CD31 in IVD-organoids. Scale bars, 50 μm.
3.2 | Single-cell transcriptome analyses of IVD-organoids

To gain further insight into the cell-type composition in IVD-organoids, we dissected out the GFP-positive IVD-organoids under a fluorescence stereo microscope two months after transplantation and performed single-cell transcriptome analysis. In total, we captured 6,131 GFP-expressing human cells, among which 8 subsets of cells were identified by unbiased clustering (Figure 2A, B). The majority of identified cell types in our IVD-organoids were also reported in stage-matched in vitro hCOs,18 including radial glial cells (RGCs), neurons, and astrocytes (Figure 2C). We also detected in IVD-organoids supporting cells that were beneficial for neurogenesis and cell survival,19 such as ChP cells and BMP signal-related cells (BRCs, GO:0030509, possibly be related to certain neural precursor cells during brain development) (Figure 2D).

Unexpectedly, we found a pericyte-like cluster expressing PDGFRb and COL3A1 (Figure 2E). Immunostaining further verified the existence of PDGFRb-positive cells in IVD-organoids, which were not detected in stage-matched in vitro hCOs and transplanted dissociated cells (Figure 2F). These results suggested that perivascular lineage differentiation, which is important for the neurovascular formation during brain development,20 could be induced in the grafted premature organoids. IVD-organoids thus exhibited increased cell-type diversity than hCOs, with more supporting cells generated.

Defined culture conditions guide the in vitro development of organoids representative for certain brain regions.3,21 In order to clarify the regional identity of IVD-organoids, we analyzed the expression profile of brain regional signature genes in IVD-organoids. In contrast to hCOs which mainly recapitulated the development of cortex,2,18 we found that cortical markers were hardly expressed in IVD-organoids (Figure S2A). Instead, we unexpectedly found that many cells in IVD-organoids expressed striatal markers ZBTB20 and STRN3. Consistently, we observed more ZBTB20-positive cells in IVD-organoids than in hCOs and dissociated cells by immunostaining (Figure 2G, H). These ZBTB20-positive cells were distributed sparsely in the IVD-organoids instead of in the lateral ganglionic eminence (LGE)-like pattern, probably due to the relative short period of time between transplantation and detection22–24 (Figure S2B). These results thus indicated that IVD-organoids exhibited a striatal identity, likely guided by their brain microenvironment.

3.3 | Efficient generation of ChP cells in IVD-organoids

In the IVD-organoids, ChP cells were one of the major cell subsets identified by our single-cell transcriptome analysis (Figure 2A). ChP is important for cerebrospinal fluid production, neuronal survival, and brain homeostasis,11 and ChP cilia play important roles in the regulation of brain development and neuroinflammation.25 Indeed, we observed expression of cilia markers such as FOXJ1, RSPH1, and ROPN1L in the ChP cluster (Figure S2C). Consistent with the single-cell transcriptome analysis, we observed by immunostaining ChP epithelium-like structures with transhyretin (TTR)-positive areas around ventricles of IVD-organoids, but not in hCOs and dissociated cells (Figure 3A and Figure S2D). Examination of regional markers revealed that most ChP cells expressed telencephalic markers instead of hindbrain markers, suggesting a telencephalic origin of the ChP cells in IVD-organoids (Figure S3A).

To further investigate the identity of ChP cells, we performed integrated analysis of single-cell transcriptome data of IVD-organoids and previously reported stage-matched in vitro hCOs.15 We found that ChP cells from IVD-organoids were co-clustered with cilium bearing cells (CBCs) in hCOs (Figure 3B). ChP cells in IVD-organoids expressed higher levels of mature ChP markers (TTR, CLIC6, and NME5) compared with CBCs in hCOs (Figure 3C), suggesting the generation of mature ChP cells in IVD-organoids. Indeed, we observed in IVD-organoids but not in hCOs robust expression of marker genes for tight junctions (TJP1, PATJ, CLDN1, and CLDN2) (Figure 3D and Figure S3C), a feature of mature ChP cells which is important for ChP to form selective brain barriers to protect central nervous system from harmful substances. Consistent with their functions in secreting cerebrospinal fluid (CSF), ChP cells in IVD-organoids also highly expressed CSF secretion associated transporters, including APOD (CSF lipoprotein), PARK7 (parkinsonism associated deglycase), and CA2/CA12 (CSF secretion enzymes) (Figure S3B–D). These observations were in agreement with the crucial roles of ChP cells in controlling nutrient transport and CSF production. We further performed integrated analysis of single-cell transcriptome data of IVD-organoids and previously reported human ChP organoids26 and found that ChP cells in IVD-organoids were most similar to the mature ChP cells in ChP organoids (Figure 3E). Although these differences might be partly owing to the variability among different organoid cultures, our findings at least indicated efficient generation of mature ChP cells in IVD-organoids.27,28

3.4 | Reduced metabolic stress and improved cell survival in IVD-organoids

It has recently been reported that metabolic pathways are often activated in organoids, leading to increased cellular stress in cultured hCOs.18 To compare metabolic stress between IVD-organoids and hCOs, we performed integrated analysis of the common neuronal and neural progenitor cell types (i.e., neurons and RGCs) identified among IVD-organoids and a few stage-matched hCOs.15,18,27,29 This analysis revealed that metabolic stress-associated pathways, such as glycolysis, ER stress, and hypoxia, were significantly less activated in IVD-organoids (Figure 4A, C, E). Indeed, we observed significant downregulation of key genes of these pathways in IVD-organoids, including glycolysis-related genes (ALDOA, ALDOC, and ENO2).
ER stress-related genes (CDK5RAP3, GORASP2, and TMX1), and hypoxia-related genes (BNIP3L, PDX1, and SFRP1) (Figure 4B, D, F). In addition to the analysis of neuronal and neural progenitor cells, we next compared ChP cells in IVD-organoids and those from in vitro ChP organoids26 and found that metabolic stress-associated pathways were also less activated in ChP cells in IVD-organoids (Figure S4A–C).

We next investigated the impact of in vivo microenvironment on the survival of IVD-organoids. Since apoptosis pathways have been reported to be activated in long-term cultured hCOs due to the lack of vasculature structures and brain microenvironment,30 we examined apoptosis signature genes neuronal and neural progenitor cells from either IVD-organoids or cultured hCOs and found that apoptosis pathway was significantly less activated in IVD-organoids.
(Figure 4G, H). Immunostaining of cleaved caspase 3 with IVD-organoids and hCOs further confirmed these findings (Figure 4I). Similar observations were also made when comparing ChP cells from IVD-organoids with those from in vitro cultured ChP organoids (Figure 5AD). Together, our results indicate that the in vivo environment generally reduces metabolic stress and apoptosis during the formation of brain organoids.

FIGURE 4 IVD-organoids exhibit improved cell survival. (A) Glycolysis signature levels in IVD-organoids and published hCOs. (B) Violin plots showing expression levels of glycolysis-related genes. (C) ER stress signature levels in IVD-organoids and published hCOs. (D) Violin plots showing expression levels of ER stress-related genes. (E) Hypoxia signature levels in IVD-organoids and published hCO. (F) Violin plots showing expression levels of hypoxia-related genes. (G) Apoptosis signatures between IVD-organoids and published hCOs. (H) Violin plots showing expression levels of apoptosis-related genes. (I) Immunostaining of cleaved caspase 3 in IVD-organoids and D60 hCOs. Scale bars, 200 μm. Quantification results are presented as mean ± s.e.m. (n = 3 for transplanted mice and cultured D60 hCOs). **p < 0.01, unpaired two-tailed t-test

4 | DISCUSSION

The high fidelity of human brain organoids to primary tissue is an advantage for potential transplantation therapy. However, organoids are usually large in size and they require much larger surgical cavities for transplantation than single cells. The needles used in transplantation surgeries may also cause more severe surgical injury to healthy tissues along the entire route of organoid delivery. Thus, due to their large size, the transplantation of hCOs is limited to outer regions of the brain (e.g., the cerebral cortex) by making an implantation cavity in the brain tissue. In this study, we generated human brain organoid in vivo by injecting small premature organoids with smaller needles, which was more suitable for the stereotaxic injection into deep brain structures with less surgical damage. Our results demonstrated that these premature organoids survived and continued to form organoid-like structures. Further single-cell transcriptome profiling also verified the cellular complexity in IVD-organoids. By balancing the advantages and disadvantages of organoid and cell transplantation, our approach enables developing brain organoids in various brain regions including deep structures.
Vascularization of brain organoids is critical for their long-term survival and functional maturation. It has been established that host-mediated vascularization can be observed in organoid graft.\(^4,5\) Similarly, host CD31-positive blood vessels were found in our IVD-organoids, suggesting that the integration of host vascular system improved IVD-organoids survival. In our single-cell transcriptome analysis, although we did not find any cell cluster containing CD31-positive human endothelial cells, premature organoid-derived cells with pericyte marker expression (PDGFR\(\beta\) and COL3A1) were detected. Similar to our observation, pericytes were also identified in vascularized organoids.\(^5\) The presence of pericytes may facilitate graft survival and integration by promoting angiogenesis process as well as supporting the microvascular structures.\(^31,32\) Although the most common developmental origin of pericytes is mesenchymal stem cells, studies also indicated their heterogenous origin which differs by tissues.\(^33-36\) Opposed to a mesodermal origin, our results suggest an unexpected neuroectodermal ancestry of pericytes. It is intriguing to explore whether these neuroectoderm-originated pericytes are functionally different from those derived from mesoderm. The ChP cells in IVD-organoids were more mature compared with those in cultured hCOS. Since ChP epithelial cells express and secrete a variety of supportive factors for neuronal growth and survival, it is reasonable to speculate that the combination of host vascular system and these human ChP cells in IVD-organoids can provide a more hospitable niche for organoid development and survival.\(^10-12\) Besides, compared with IVD-organoids, more apoptotic cells accumulate in large mature organoids after a long period of in vitro culture due to lack of the vascular system.\(^37,38\) Although these apoptotic cells are partially removed by microglia after transplantation, insufficient clearance (e.g., in elderly individuals) can lead to unwanted chronic immune response.\(^4,39,40\)

Corpus striatum is one of the most vulnerable regions in neurodegenerative diseases and lacks endogenous neurogenesis, and thus a desired target site for cell therapy.\(^41\) However, it locates much deeper in the brain than cortex, which hampers the transplantation of large-size hCOS. By generating IVD-organoids in corpus striatum, new approaches for cell therapy may be developed in future. The majority of cells in in vitro cultured cerebral organoids using unguided methodologies express signature genes of cortical areas, including SATB2 and CTIP2.\(^2,18,42\) However, these markers only expressed in a small minority of cells in IVD-organoids. Instead, striatal and ChP markers were widely expressed in IVD-organoids. It is reported that the environment of transplantation sites including striatum could influence the differentiation and distribution of grafted cells, suggesting that the identity of IVD-organoids may be regulated in a more complicated fashion, involving both the environmental cues (including but not limited to growth factors and neurotransmitters) from host striatum and intrinsic property of grafted cells.\(^33,44\) Further investigation on IVD-organoids in other host brain regions will lead to better understanding of the interplay between host brain and transplanted organoids.

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CONFLICT OF INTEREST
The authors declare no conflict of interest.

AUTHOR CONTRIBUTIONS
G.P., L.S., and S.H. supervised this study. S.H. and Y.Y. performed organoid culture and stereotaxic surgery. J.L. performed immunostaining experiments. H.Z. and J.C. performed scRNA-seq. F.H. and L.S. analyzed the scRNA-seq data. S.H., F.H., L.S., and G.P. wrote the manuscript.

DATA AVAILABILITY STATEMENT
The GEO accession number for datasets reported in this study is GSE165975.

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**SUPPORTING INFORMATION**

Additional supporting information may be found in the online version of the article at the publisher’s website.

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