Disorders of the Nervous System

Vascularization and Engraftment of Transplanted Human Cerebral Organoids in Mouse Cortex

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Visual Abstract

Neural stem cells (NSCs) hold great promise for neural repair in cases of CNS injury and neurodegeneration; however, conventional cell-based transplant methods face the challenges of poor survival and inadequate neuronal differentiation. Here, we report an alternative, tissue-based transplantation strategy whereby cerebral organoids derived from human pluripotent stem cells (PSCs) were grafted into lesioned mouse cortex. Cerebral organoid transplants exhibited enhanced survival and robust vascularization from host brain as compared to transplants of dissociated neural progenitor cells (NPCs). Engrafted cerebral organoids harbored a large NSC pool and displayed multilineage neurodifferentiation at two and four weeks after grafting. Cerebral organoids therefore represent a promising alternative source to NSCs or fetal tissues for transplantation, as they contain a large set of neuroprogenitors and differentiated neurons in a structured organization. Engrafted cerebral organoids may also offer a unique experimental paradigm for modeling human neurodevelopment and CNS diseases in the context of vascularized cortical tissue.

Key words: cerebral organoid; CNS injury; neural stem cell transplant; vascularization of intracerebral graft

Significance Statement

Neural stem cells (NSCs) hold great promise for neural repair, but conventional cell-based transplant methods face the hurdles of poor graft survival and inadequate neural differentiation. Here, we transplanted cerebral organoids derived from human pluripotent cells into lesioned mouse cortex. We report enhanced survival, robust vascularization, and multilineage differentiation of engrafted human cerebral organoids in host mouse brain. Cerebral organoid transplantation therefore represents an alternative, tissue-based transplantation strategy for neural repair and for modeling human neurodevelopment and CNS diseases in the context of vascularized cortical tissue.
CNS injury or degeneration results in devastating neurologic deficits from irreversible loss of neurons. Functional compensation from surviving neural networks and reparative efforts from endogenous neural stem cells (NSCs) are limited in their efficacy. Cell replacement therapy has thus been explored for reconstruction of neural circuits since the early 1970s (Das and Altman, 1972), and the interest has been rekindled with the advent of induced pluripotent stem cells (iPSCs; Takahashi and Yamanaka, 2006) and improved neuronal differentiation protocols (Thompson and Björklund, 2015; Grade and Götz, 2017). Grafted NSC can undergo neuronal and glial differentiation, and display neurite outgrowth (Snyder et al., 1997). However, engraftment rate, long-term survival, and neuronal differentiation remain limited. Hence, novel strategies are needed to advance cell replacement therapy as a viable treatment option for CNS injury or degeneration.

Contrary to the poor engraftment rate of implanted dissociated neural progenitor cells (NPCs), mouse embryonic cortical tissue transplants survive well and show successful integration in adult mouse brain with establishment of substantial connectivity with host targets (Gaillard et al., 2007). The proof-of-principle that neural tissue replacement can work in human has been provided by the success of intrastriatal transplantation of human fetal mesencephalic tissue for Parkinson’s disease (PD) patients (Kordower et al., 1995). Clinical trials have since been conducted for fetal tissue transplant in PD patients, demonstrating long-term safety and clinical benefits (Lazic and Barker, 2003; Barker et al., 2013). Therefore, a shift from cell-based to tissue-based transplantation represents a promising strategy, but the scarcity of fetal tissues and ethical concerns limit the development of such an approach (Lindvall et al., 2004).

Recently, a novel three dimensional (3D) culture method has been developed to differentiate human PSCs into cerebral organoids (Lancaster et al., 2013; Lancaster and Knoblich, 2014). Cerebral organoid culture taps into the enormous self-organizing capacity of embryonic stem cells (ESCs)/iPSCs to form complex tissue structures under defined feeder cell-free conditions, with no addition of exogenous patterning cues or morphogens, an important safety point for transplantation. After 30 d of culture in Matrigel droplets under rotary condition, cerebral organoids adopt a predominantly dorsal forebrain regional specification, containing fluid-filled ventricle-like structures that are aligned with Sox2+ neuroprogenitors in a ventricular/subventricular-like zone (VZ/SVZ) and doublecortin (DCX)+ neuroblasts in an outer layer. Rudimentary cortical stratification takes place after longer culture periods, with cortical neurons differentiating into pyramidal identities displaying glutamatergic receptor activity and efferent long-range axons in a stereotypical inside-out stratified layout (Lancaster et al., 2013; Lancester and Knoblich, 2014). Cerebral organoids thus provide a new experimental platform to study human brain development and to model CNS disorders such as microcephaly (Lancaster et al., 2013; Li et al., 2017), autism spectrum disorders (Forsberg et al., 2018), and Zika virus infection (Qian et al., 2016; Watanabe et al., 2017).

We hypothesized that transplantation of cerebral organoids derived from human ESC (hESC)/iPSC may enhance graft survival, neural differentiation, and integration in host brain as compared to conventional cell-based transplantation for the following reasons. First, the 3D cellular arrangement of the cortical plate-like tissue in cerebral organoids may provide a protective shield against hostile elements at the graft site. Second, a neuroprecursor pool in the cerebral organoids residing in protected stem cell niches in the VZ/SVZ may serve as a source for neurogenesis and stem cell-derived trophic factors. Third, a rudimentary 3D cortical structure is already in place at the time of transplant, enabling intrinsic patterning cues to direct organized neuronal differentiation and prevent aberrant proliferation and lineage progression.

Here, we performed comprehensive side-by-side comparisons of transplantation of dissociated NPC versus cerebral organoids, both derived from identical hESC cultures. Postnatal day (P)8–P10 mice were used as recipients and frontoparietal cortex as transplant site. We compared graft survival, vascularization, NSC population, and neurodifferentiation at two and four weeks after transplantation. We found enhanced survival of cerebral organoid transplants as compared to grafted NPC, and robust vascularization of the organoid grafts from host vessels. There were also abundant neuroprogenitors and evidence for multilayer differentiation in the engrafted cerebral organoids. A recent study by Mansour et al. (2018) also tested intracerebral grafting of hPSC-derived brain organoids. In that study, adult nonobese diabetic/severe combined immunodeficient (NOD-SCID) mice were used as recipients and retrosplenial cortex was selected as transplant site based on the rich vascularized surface overlaying this area. They reported that organoid grafts showed successful engraftment with robust vascularization from host brain, and in long-term analysis of up to eight months after transplant, progressive neuronal differentiation and maturation, long-range axon projections,
and functional graft-to-host synaptic connectivity were observed. Our data thus echo those of Mansour et al. (2018) in demonstrating the practicality of transplantation of hiPSC-derived cerebral organoids as a promising alternative for cell replacement therapy for CNS injury and neurodegeneration. Transplantation of cerebral organoids also provides a unique experimental paradigm to study human neurodevelopment and to model CNS diseases in the context of vascularized cortical tissue.

Materials and Methods

Animal care

P8–P10 CD1 mice of either sex (Charles River Laboratories) were used as transplant recipients for analysis up to four weeks after grafting without immunosuppressive treatment, as immunosuppression was only mandatory to achieve engraftment beyond two months (Espuny-Camacho et al., 2013). Mice were group-housed and kept in a 12/12 h light/dark cycle with free access to food and water ad libitum. All animal procedures were performed in accordance with the IACUC committee of Icahn School of Medicine at Mount Sinai animal care committee’s regulations.

hESC culture

Human pluripotent ESCs were provided by WiCell (H9 hES cells, WAe009-A). For hESC culture, six-well plates were coated with diluted Matrigel (growth factor reduced; 1:100, BD Biosciences) for 20 min at 37°C, and cells were plated and cultured in mTeSR1 media (STEMCELL Technologies) supplemented with 2 μM ROCK inhibitor Thiazovivin (Millipore) for 24 h. Cells were cultured with media changed every day until ready to passage or harvest in mTESR1 media (without ROCK inhibitor).

GFP labeling of hESCs

H9 hES cells were infected with lentivirus EF1a-GFP-IRES-Puro, followed by puromycin selection (1 μg/ml, Thermo Fisher Scientific). For lentivirus preparation, the pEGIP lentivirus plasmid (Addgene plasmid #26777) was transfected into 293T cells together with envelope plasmid pMD2.G and packaging plasmid psPAX2 (Addgene #12259 and #12260) with X-tremeGENE 9 DNA transfection reagent (Roche). Lentiviruses were concentrated from culture media supernatant 72 h after transfection by ultracentrifugation.

Generation of hNPC from hESCs

hES cells were cultured in low attachment 96-well plates for 4 d to generate embryoid bodies (EBs). EBs were then transferred to Matrigel-coated six-well plates for attachment and further cultured in neural induction medium (STEMdiff, STEMCELL Technologies) for 4–5 d. Next, cells were plated on laminin-coated (10 μg/ml, Thermo Fisher Scientific) six-well plates and cultured in human NSC medium (NeuroCult, STEMCELL Technologies) supplemented with 20 ng/ml of epidermal growth factor (EGF) and 10 ng/ml of basic fibroblast growth factor (bFGF; Peprotech) for 7 d for NPC maturation, which were then used for transplant.

For proliferation and differentiation assays, 12-mm glass coverslips were pre-coated with 50 μg/ml poly-D-lysine (PDL; Sigma Aldrich) and 10 μg/ml laminin (Thermo Fisher Scientific). NPC were seeded on coverslips at 12 000 cells/cm² density and cultured in human NSC medium (NeuroCult, STEMCELL Technologies) supplemented with 20 ng/ml of EGF and 10 ng/ml of bFGF (Peprotech). For differentiation assays, cells were cultured in the same media, but with withdrawal of mitogens (EGF, bFGF) for 5 d. Cells were fixed with 4% paraformaldehyde (pH 7.4, Acros Organics) in PBS at 4°C for 15 min and analyzed by immunofluorescence.

Cerebral organoid generation from hESCs

Human cerebral organoids were generated as described (Lancaster et al., 2013; Lancaster and Knoblich, 2014), with modifications as follows: human ES cells were detached using 50 μM EDTA (Thermo Fisher Scientific) and plated in round bottom ultra-low attachment 96-wells plate (CLS7007, Corning) at a density of 9000 cells per well in mTESR1 media (STEMCELL Technologies) supplemented with 1% antibiotics (penicillin streptomycin, Thermo Fisher Scientific) for a total of 6 d. During the first 4 d of the culture, media were supplemented with 10 μM Thiazovivin. Half of the media was changed every day. After 6 d of culture or when embryonic bodies (EBs) reached ~500–600 μm in diameter and when surface tissue began to brighten and formed smooth edges, media were switched to a neural induction media (Stemdiff, STEMCELL Technologies). Half of the media was changed every day for 3–4 d. After neuroepithelium emerged (usually at approximately day 9–10), organoids were embedded in Matrigel droplets (25 μl, BD Biosciences) and cultured in 6 cm Petri dishes (Falcon) for 4 d in cerebral organoid differentiation media consisting of 1:1 DMEM-F12 and Neurobasal media (Gibco), with addition of 0.5% N2 supplement (Life Technologies), 0.5% ml MEM-NEAA (Gibco), 1% Glutamax (Gibco), 1% B27 supplement without vitamin A (Life Technologies), 0.1 μM of 2-mercaptoethanol (Millipore), 2.6 μg/ml insulin (Sigma Aldrich), and 1% pen/strep antibiotics (Gibco). After 4 d, the organoid Matrigel droplets were cultured with addition of vitamin A on an orbital shaker (VWR) at 85 rpm for four additional weeks, and then used for transplant.

Grafting of NPC or cerebral organoid into mouse cortex

Before transplantation, a quality control of cerebral organoids was performed by brightfield microscopy to select the organoids that displayed an appropriate differentiation/maturation phenotype without massive cyst formation or premature differentiation. Then, a measurement of organoid size was performed by brightfield microscopy and organoids of similar sizes were selected for transplant. To estimate the cell numbers in organoids at time of transplant, organoids were dissociated by 15-min trypsin incubation, and cells were counted by Trypan blue exclusion. P8–P10 mice were anesthetized with isoflurane and secured on a stereotactic frame (Stoelting). Scalp was opened on the left hemisphere. Using a restricted depth
stab knife, an ~1 × 1 mm craniotomy window was opened, with the bone flap hinged on anterior base. A cortical lesion was made by removing ~1 mm³ piece of the frontoparietal cortex. Using a pair of forceps, one cerebral organoid at 42 d in vitro culture was implanted into the lesioned mouse cortex. The craniotomy window was closed by returning the bone flap to the original position and sealed with fibrin glue (Evicle, Fibrin Sealant), followed by skin closure with sutures.

**Intracerebral implantation of NPC**

NPC differentiated from H9 hES cells were dissociated with trypsin, counted, and assessed for viability by Trypan blue exclusion. Cells were washed twice with DMEM, and then suspended in DMEM at a concentration of 5 × 10⁵/µl. Using a 5-µl Hamilton syringe (Hamilton glass syringe 600 series RN) and a 26-gauge needle, 1 × 10⁵ NPC (2 µl) were implanted into the left frontoparietal hemisphere. To be consistent with the organoid transplant plant, the coordinates used for the intracerebral injection were 1 mm posterior to the bregma, 1 mm lateral to the sagittal suture (left hemisphere), and 1 mm below the dura.

**Histologic analyses**

Two or four weeks after transplantation, animals received a lethal dose of pentobarbital (390 mg/ml) and transcardial perfusion was performed with 50-ml cold PBS, followed by 50-ml cold 4% PFA/PBS. Brains were removed and post-fixed overnight in 4% PFA/PBS and cryoprotected for 48 h in 30% sucrose. Brains were then snap-frozen in isopentane (Sigma Aldrich) and embedded in O.C.T. compound (Tissue-plus, Fisher Healthcare) and stored at −80°C. Brain tissues were sectioned into 14-µm coronal sections with cryostat (CM1850, Leica). For immunofluorescence analysis, non-specific binding sites were blocked with 4% BSA in PBS (Fisher Bioreagents), 0.2% Tween (Tween 20, Acros Organics), and 10% normal donkey serum (Jackson ImmunoResearch) for 1 h at room temperature, and slices were then incubated with the following antibodies diluted in 4% BSA/PBS, 0.2% Tween: rabbit anti-activated caspase 3 (AC3; Abcam, ab2302, 1:100); rat anti-mouse CD31 (BD Biosciences, 553370, 1:200); rat anti-mouse CD45 (BD Biosciences, 550539, 1:200); guinea pig anti-DCX (Millipore, AB2253, 1:500); mouse anti-GalC (Millipore MAB342, 1:200); rabbit anti-GFP (Invitrogen 180063, 1:200); chicken anti-Neurofilament H (NF-H, Abcam, ab110258, 1:100); rabbit anti-Nanog (Abcam, ab109250, 1:300); rabbit anti-Oct4 (Abcam, ab194665, 1:300); rabbit anti-Olig2 (Millipore AB9610, 1:500); rabbit anti-SOX2 (Millipore AB5603, 1:200); rabbit anti-TBR1 (Abcam ab31940, 1:500); rabbit anti-TBR2 (Abcam ab23345, 1:500), and mouse anti-tubulin β-III (TuJ1, R&D Systems MAB1195, 1:100). Slides were then washed in PBS with 0.1% Tween and detection was performed with Alexa Fluor-coupled secondary antibodies (Invitrogen and Jackson ImmunoResearch) and DAPI nuclear counterstain (Invitrogen).

Quantifications were performed on at least two brain slices per animal and three independent mice for each condition. The two slices were separated by ~150 µm. We selected the sections that were in the middle of the transplant area. Grafts were outlined by GFP or hMito (human mitochondria marker) immunofluorescence. For each selected 14-µm graft-bearing brain slice, we quantified different markers within the entire GFP or hMito-labeled area by Fiji software (Schindelin et al., 2012) unless otherwise specified.

**Statistical analysis**

For each experiment, the number of mice used in each cohort, and the number of images analyzed from each animal are listed in figure legends. Data are presented as mean value with SEM, unless otherwise stated. At least three independent grafts in three different animals were tested and at least two representative images from each animal were quantified. Cluster-based summary statistics using within-subject averaging were performed whenever possible. GraphPad Prism 7 was used for statistical analyses. Differences between conditions were determined using a two-way ANOVA test, followed by a Tukey post hoc test. For individual comparison within the group, normality of data was assessed using a Shapiro–Wilk test followed by a Student’s t test. Results were considered significant for p < 0.05 (Table 1).

**Results**

**hESC-derived NPC and cerebral organoids for intracerebral transplantation**

For side-by-side comparisons, we differentiated the same batch of hESC into either NPC or cerebral organoids and performed stereotactic transplantation into left frontoparietal cortex of immunocompetent P8–P10 mice (Fig. 1A). We compared graft survival, vascularization, NSC pool, and neurodifferentiation at two and four weeks after transplantation (Fig. 1B). To ensure that the organoid grafts remained inside the lesion cavity following transplantation, we optimized the technique of opening a small craniotomy window with a bone flap hinged at the anterior base, and after transplantation, the bone flap was returned to original position, sealed with fibrin glue, followed by scalp closure (Fig. 1C).

To facilitate visualization of donor cells in host cortical tissue, we generated a hESC line expressing green fluorescent protein (GFP) ubiquitously (Zou et al., 2009; Fig. 2A). After lentiviral infection and puromycin selection, GFP-expression of hESC was assessed, while their pluripotency was verified by immunocytochemistry with uniform expression of stemness markers Oct4 and Nanog (Fig. 2B). GFP-positive hESC were then differentiated into hNPC by way of EB formation followed by neural induction (Fig. 2C,D). To confirm multipotency of hNPC, differentiation condition was applied for 5 d by mitogen withdrawal, and immunocytochemistry revealed expression of TUJ1 (neuronal marker), GFAP (astrocyte marker), and GalC (oligodendrocyte marker) while most cells re-
mained proliferative as shown by the proliferation marker Ki67 (Fig. 2E). In parallel, the same batch of GFP-expressing hESC were used to generate cerebral organoids (Lancaster et al., 2013; Lancaster and Knoblich, 2014). GFP expression was confirmed at each stage of organoid development (Fig. 2F,G). After 42 d of in vitro culture, with the last 28 d under rotary condition embedded in Matrigel droplet, cerebral organoids adopted a predominantly dorsal forebrain specification, containing multiple ventricle-like structures aligned with SOX2+ neu-

| Table 1. Statistical analyses |
|-------------------------------|
| Data structure | Type of test | Condition | Power |
| **Fig. 3B** | Dataset with more than two groups | Two-way ANOVA - Tukey post hoc test | Two weeks: organoids vs two weeks: NPCs | *p = 0.0393 |
| | | | Four weeks: organoids vs four weeks: NPCs | n.s., p = 0.7729 |
| **Fig. 3D** | Dataset with more than two groups | Two-way ANOVA - Tukey post hoc test | Two weeks: organoids vs two weeks: NPCs | n.s., p = 0.1836 |
| | | | Four weeks: organoids vs four weeks: NPCs | n.s., p = 0.9634 |
| **Fig. 5B, left** | Dataset with more than two groups | Two-way ANOVA - Tukey post hoc test | Two weeks: organoids vs two weeks: NPCs | n.s., p = 0.1059 |
| | | | Four weeks: organoids vs four weeks: NPCs | *p = 0.0293 |
| **Fig. 5B, right** | Dataset with more than two groups | Two-way ANOVA - Tukey post hoc test | Two weeks: organoids vs two weeks: NPCs | n.s., p = 0.6113 |
| | | | Four weeks: organoids vs four weeks: NPCs | n.s., p = 0.7485 |
| | | | Four weeks: organoids vs two weeks: organoids | n.s., p = 0.2955 |
| **Fig. 6B, top** | Dataset with more than two groups | Two-way ANOVA - Tukey post hoc test | Two weeks: organoids vs two weeks: NPCs | **p = 0.0014 |
| | | | Four weeks: organoids vs four weeks: NPCs | *p = 0.0108 |
| | | | Four weeks: organoids vs two weeks: organoids | n.s., p = 0.9757 |
| **Fig. 6B, bottom** | Dataset with more than two groups | Two-way ANOVA - Tukey post hoc test | Two weeks: NPC vs two weeks: Organoid | n.s., p = 0.2141 |
| | | | Four weeks: organoids vs two weeks: organoids | n.s., p = 0.998 |
| **Fig. 6D, top** | Dataset with more than two groups | Two-way ANOVA - Tukey post hoc test | Two weeks: organoids vs two weeks: NPCs | n.s., p = 0.1664 |
| | | | Four weeks: organoids vs four weeks: NPCs | n.s., p = 0.7125 |
| | | | Four weeks: organoids vs two weeks: organoids | n.s., p = 0.3625 |
| **Fig. 6D, bottom** | Dataset with more than two groups | Two-way ANOVA - Tukey post hoc test | Two weeks: organoids vs two weeks: NPCs | n.s., p = 0.9555 |
| | | | Four weeks: organoids vs four weeks: NPCs | n.s., p = 0.9037 |
| | | | Four weeks: organoids vs two weeks: organoids | n.s., p = 0.2857 |
| **Fig. 8B** | Dataset with more than two groups | Two-way ANOVA - Tukey post hoc test | Two weeks: organoids vs two weeks: NPCs | *p = 0.0311 |
| | | | Four weeks: organoids vs two weeks: NPCs | n.s., p = 0.1724 |
| **Fig. 8D** | Dataset with more than two groups | Two-way ANOVA - Tukey post hoc test | Two weeks: organoids vs two weeks: NPCs | *p = 0.0089 |
| | | | Four weeks: organoids vs two weeks: NPCs | n.s., p = 0.0697 |
| | | | Four weeks: organoids vs four weeks: NPCs | ***p = 0.0006 |
| | | | Four weeks: organoids vs two weeks: organoids | **p = 0.0041 |
| **Fig. 8F** | Dataset with two groups | Shapiro–Wilk followed by a Student’s t test | Two weeks: organoids vs four weeks: organoids | n.s., p = 0.3437 |
roprogenitors in the VZ/SVZ zone and DCX+ neuroblasts in the outer layer (Fig. 2H).

Enhanced survival of cerebral organoid transplants

We grafted one single hESC-derived cerebral organoid into the lesion cavity in the left frontoparietal cortex in P8–P10 mice. For NPC transplantation, dissociated NPC were implanted into identical cortical region by stereotactic injection. Experimental timeline. P8–P10 immunocompetent mice were used as recipients for transplantation of dissociated human NPC or human cerebral organoids. Histologic analyses of the grafts were performed two or four weeks after grafting. C, Schematic diagrams (top) and intraoperative photographs (bottom) of cerebral organoid transplantation. Briefly, from left to right, once scalp is reflected, a small craniotomy window was opened with the bone flap hinged at the anterior base, and a small piece of the frontoparietal cortex (~1 mm³) was removed. A single cerebral organoid was then implanted into the prelesioned mouse cortex, followed by return of the bone flap to close the craniotomy window, sealed with fibrin glue, followed by scalp closure.

To compare engraftment rate, we first measured the size of the graft areas as demarcated by GFP-labeled grafted cells. Consistent with earlier reports of poor survival of transplanted NSC in dissociated state (Johann et al., 2007), NPC grafts displayed significant shrinkage from two- to four-week time periods after transplant ($p = 0.039$), whereas the size of cerebral organoid grafts remained stable between two and four weeks after transplantation (Fig. 3A,B). In some instances, organoids grafts appeared fragmented, likely a result of technical difficulty (transplanting a relatively large human cerebral organoid into a small host mouse brain) or cell death from trauma, hypoxia, and inflammation before engraftment took place. Taken together, our finding of an enhanced survival of...
organoid grafts supports the hypothesis that structured cellular arrangement in tissue-based transplants provides better protection of donor cells from the hostile graft microenvironment.

We next examined the extent of apoptosis in NPC versus organoid transplants by immunohistochemistry for AC3. We found no significant differences in the average number of AC3+ cells per unit GFP+ graft area between the two graft types at two weeks (p = 0.18) or four weeks (p = 0.96) after transplantation, although there were high variabilities (Fig. 3C,D). Notably, similar to the finding by Mansour et al. (2018), the number of apoptotic cells were...
much lower in organoid grafts than in stage-matched cultured organoids (Fig. 3E), which may reflect phagocytic clearing in vivo.

To better understand the time course of organoid engraftment, we performed additional short-term post-grafting analyses. We found that by 3 or 5 d after transplantation, organoid grafts had not yet been firmly integrated into host brains. Dashed white lines delineate the graft areas; \( p < 0.05 \); n.s., non-statistically significant. Two-way ANOVA followed by a Tukey post hoc test; \( n = 3 \) mice for each time point and two images from each mouse.

Figure 3. Engraftment and survival of NPC and cerebral organoid transplants. A, Representative immunofluorescence images of NPC transplant (left) and cerebral organoid transplant (right) at the indicated time points post-grafting. Enlarged images of the boxed area are shown at the bottom. D: dorsal, V: ventral, M: medial, L: lateral. B, Quantification showing the relative size of GFP-positive area of NPC and organoid transplants at two and four weeks after grafting. A significant decrease of the graft size of the NPC transplants was detected at four weeks as compared to two weeks after grafting. C, Representative immunofluorescence images for hMito (human mitochondria) and AC3 in NPC transplants (left) or in cerebral organoid transplants (right) at the indicated time points after transplantation. D, Quantification showing the number of AC3-positive apoptotic cells per unit area of hMito-positive grafts at the indicated time points. E, Representative immunofluorescence images showing a high number of apoptotic cells (AC3+) in stage-matched cerebral organoids in culture at six and eight weeks. F, Representative immunofluorescence images of NPC transplant (left) and cerebral organoid grafts (right) 3 or 5 d after transplantation. At these early time points, grafts had not yet been firmly integrated into host brains.
activated microglia/macrophages. We detected low number of Iba1+ cells in both NPC and organoid transplants after two weeks; but after four weeks, the Iba1+ cells were increased and exhibited hypertrophied morphology in NPC transplants and adjacent host brain tissues, suggesting enhanced phagocytic activity. In contrast, Iba1+ cells remained relatively scant and small in size in organoid transplants after four weeks (Fig. 4A). The Iba1+ cells in the organoid grafts did not colocalize with hMito (human mitochondria marker), indicating host origin of the microglia within the grafts, similar to the data shown by Mansour et al. (2018). Similar results were observed using leukocyte common antigen CD45. At the transplant sites, we detected only scant CD45hi cells in organoid grafts, and slightly more CD45hi cells in NPC grafts, possibly indicating increased phagocytosis of dead donor cells within the NPC transplants (high regional variability precluded accurate quantification; Fig. 4B).

**Successful vascularization from host into the grafted cerebral organoids**

Vascularization of the graft is a key determinant for successful integration into host tissue. We therefore compared the extent of graft vascularization by immunohisto-
We first quantified the number of blood vessels in the entire graft area as labeled by hMito immunostaining. We detected robust vascularization of the organoid transplants at both two and four weeks after grafting (Fig. 5A). Remarkably, CD31+/PECAM1 microvasculatures were present not only at the periphery, but also at the center of the grafts. Closer inspection revealed no apparent overlap between

Figure 5. Vascularization of engrafted cerebral organoids. A, Representative immunofluorescence images demonstrate penetration of host CD31-positive blood vessels into hMito+ NPC grafts (left panels) or cerebral organoid grafts (right panels) at the indicated time points post-transplantation. Notice that CD31-positive endothelial cells inside the grafts are hMito-negative (white arrowheads). White arrows: host vasculature. Dashed white lines delineate the graft areas. Enlarged images of the boxed area are shown on the right. D: dorsal, V: ventral, M: medial, L: lateral. B, Quantifications of the number (left) and the average length (right) of CD31-positive blood vessels in hMito-labeled grafts demonstrate a higher number of vasculatures in the engrafted cerebral organoids compared to NPC transplants at four weeks after transplantation, but no significant difference in the average vascular length; +p < 0.05; n.s., non-statistically significant; n = 3 mice for each cohort, and at least two images analyzed from each mouse. Two-way ANOVA followed by a Tukey post hoc test.
CD31 and hMito immunostaining, indicating host origin of the graft vasculature. From two to four weeks following transplantation, the number of microvasculatures in the engrafted organoids remained stable (Fig. 5B). In the NPC transplants, we also observed abundant CD31+/microvasculatures within the graft after two weeks; but after four weeks, a lower number of CD31+ blood vessels were detected (Fig. 5A). Quantification confirmed the presence of significantly more CD31+ vessels in organoid transplants than in NPC transplants at four weeks after grafting (p = 0.029). To verify that the higher number of blood vessels found in organoid transplants was not due to shorter or more branched vasculatures, we measured average vascular length, which was comparable between the two graft types (Fig. 5B). Notably, donor cells appeared disorganized in relation to vasculatures in NPC grafts, which may reflect their dissociated state at the time of transplant as compared to the tissue-like cellular arrangement in organoid transplants.

Neuroprogenitor proliferation and neurodifferentiation in engrafted cerebral organoids

We next examined proliferation of donor cells in the transplants using the proliferation marker Ki67. A significantly lower density of Ki67+ cells per unit GFP+ graft area was detected in NPC transplants as compared to organoid transplants at both two and four weeks after grafting (p = 0.0014 and p = 0.0108, respectively), while there was no significant difference in organoid transplants between the two time points (p = 0.97; Fig. 6A,B). The percentage of Ki67+/DAPI+ cells at two weeks after transplantation was also much lower in NPC transplants (~2.5%) than in organoid transplants (~9%; p = 0.038), but the difference at four weeks became more variable, thus not statistically significant (~4.5% in NPC grafts and 9% in organoid grafts, p = 0.21; Fig. 6B). Of note, in NPC cultures, proliferative cells were abundant, even after 5 d under differentiation condition (shown in Fig. 2E); in contrast, in cultured organoids, Ki67+ cells were restricted to VZ/SVZ (Fig. 2H). Therefore, the starting number of proliferative cells in NPC transplants may in fact be higher than in organoid transplants. There was no significant difference in the percentage of Ki67+/DAPI+ cells in the organoid grafts between two and four weeks after grafting (p = 0.99). It is also notable that unlike in stage-matched cultured organoids where Ki67+ proliferative cells were detected predominantly in the progenitor zones aligning the ventricle-like structures (Fig. 7A), the Ki67+ cells in transplanted organoids appeared scattered throughout the grafts, reflecting disintegration of the ventricular structures following transplantation.

We also assessed neuroprogenitor populations in the two graft types. The Sox2+ neuroprogenitors remained abundant in both graft types, with comparable density at two and four weeks after transplantation. In regards to the percentage of Sox2+/DAPI+ cells, the results were also comparable for NPC and organoid transplants with ~30% cells expressing Sox2 at both time points (Fig. 6C,D). However, as the surviving NPC grafts markedly shrunk in size from two to four weeks after grafting, the total number of Sox2+ neuroprogenitors in the NPC grafts became markedly smaller; in contrast, there was a stable and sizable NSC pool in the organoid transplants at both time points (Fig. 6C). Similar to the Ki67 results, the Sox2+ cells also became scattered in the grafts, unlike in stage-matched cultured cerebral organoids where Sox2+ cells resided in the VZ/SVZ aligning ventricle-like structures (Fig. 7A). It is also worth mentioning that the host cortex also contains numerous Sox2+ oligodendrocyte precursor cells (OPCs).

We next compared neurodifferentiation in the two graft types. We detected significantly more DCX+ neuroblasts per unit area of GFP+ organoid transplants than NPC transplants at four weeks after transplantation (p = 0.0006), although at two weeks, the expression levels of GFAP per unit graft area were comparable between the two graft types (Fig. 8C,D). Temporal comparison showed a significant increase of GFAP expression within organoid transplants between two and four weeks after grafting (p = 0.004). It is worth noting the central location of GFAP+ cells within the grafts and the colocalization of GFAP and hMito immunofluorescence, which support donor origin of the astrocytes (as opposed to infiltrating host reactive astrocytes). Finally, the presence of the oligodendrocyte lineage was examined by Olig2 immunofluorescence. We observed an average of 10.8% and 12.4% of cells expressing Olig2 in organoid grafts at two and four weeks after transplantation, respectively (Fig. 8E,F). In stage-matched cultured cerebral organoids, there were abundant DCX+ neuroblasts in organoids at both six and eight weeks of culture, scant GFAP+ astrocytes at eight weeks of rotary culture, but no detectable Olig2+ cells at either six or eight weeks (Fig. 7B).

Further characterization of neuronal differentiation in the engrafted cerebral organoids showed the presence of cells expressing TBR2 (also known as EOMES), a marker for intermediate progenitor cells (Englund et al., 2005), which appeared more abundant at two weeks as compared to four weeks after grafting (Fig. 9A). Notably, there were no TBR2-expressing cells in the host cortex. We also observed cells expressing CTIP2, a marker corresponding to deep layer neurons, particularly at four weeks after grafting (Fig. 9B). These data are in agreement with the temporal pattern of lineage progression of neuronal differentiation; however, no layer organizations of CTIP2-positive neurons were detected in the organoid transplants, unlike in stage-matched cerebral organoid cultures (Fig. 7A).

To further assess the presence of mature neurons in different transplants and examine axonal growth and pro-
Figure 6. Cell proliferation and NSC pool in cerebral organoid transplants. **A**, Representative immunofluorescence images of NPC (left panels) and cerebral organoid transplants (right panels) stained for GFP and proliferation marker Ki67. D: dorsal, V: ventral, M: medial, L: lateral. **B**, Quantification (top) showing a higher density of Ki67+ cells per unit GFP+ area in cerebral organoid than in NPC transplants at both two and four weeks after transplantation. Bottom quantification: percentage of Ki67+/DAPI+ cells within the GFP+ cerebral organoid and NPC grafts. **C**, Representative immunofluorescence images of NPC (left panels) and cerebral organoid transplants (right panels) showing abundant engrafted cells (GFP+) expressing stem cell marker Sox2 (red). White arrows: Sox2+...
jections, we performed immunostaining for the neurofila-
ment heavy chain (NF-H). While no NF-H immunosignals
were detected at two or four weeks after NPC transplanta-
tion (data not shown), there was colocalization of NF-H and
hMito immunosignals in organoid transplants at both time
points, some with long projections (Fig. 9C). However, no
organized projection pattern was detected, which may re-

Figure 7. Stage-matched in vitro cerebral organoid characterization. A, Representative immunofluorescence images of cultured cerebral organoids at six or eight weeks of maturation show layered organization of cortical-like tissue with proliferating cells (Ki67+) and NPC (SOX2+) mainly localized in the VZ/SVZ and neurons (CTIP2+) localized in the outer layer. B, Representative immunofluorescence images of cerebral organoids after six or eight weeks of in vitro maturation. Left, A low number of astrocytes (GFAP+) was present in the organoids after eight weeks of maturation (white arrow), while no Olig2+ cells were detected. Right, Abundant neuroblasts (DCX+) were found at both time points while low level of expression of NF-H was detected at eight weeks of maturation.
Figure 8. Neurodifferentiation of the engrafted C-organoids. A, Representative immunofluorescent images of DCX-positive neuro-
blasts in NPC (left panels) and cerebral organoid transplants (right panels) at the indicated time points. D: dorsal, V: ventral, M: medial, L: lateral. B. Quantification of DCX immunointensity per unit area of GFP+ grafts. A significant stronger staining intensity of DCX was measured in organoid compared to NPC transplants after four weeks. C. Representative immunofluorescence images of GFAP+ cells in hMito-labeled NPC (left) and organoids grafts (right) at the indicated time points. White arrows denote colocalization of hMito and GFAP markers in transplants. D. Quantification of GFAP immunointensity per unit area of hMito+ grafts. A significant stronger staining intensity of GFAP was measured in organoid transplants after four weeks. E. Representative immunofluorescence images of Olig2+ cells in the engrafted cerebral organoids at the indicated time points. White arrows: Olig2+ cells in host cortical tissue. F. Quantification shows no significant difference in the percentage of Olig2+/DAPI+ cells in the organoid grafts between two and four weeks after grafting. Dashed white lines delineate the graft areas. Enlarged images of the boxed area are shown on the right: *p < 0.05, **p < 0.01, ***p < 0.001; n.s., non-statistically significant. Two-way ANOVA followed by a Tukey post hoc test and Student’s t test; n = 3 mice for each time point and two images from each mouse.

digm. In stage-matched cerebral organoid cultures, we also detected only low NF-H+ immunosignals (Fig. 7B).

Discussion

In conventional stem cell transplantation approaches, dissociated stem cells are deposited in suspension through intracerebral, intravenous, intraarterial, or transnasal routes, all of which expose the transplanted cells to immediate hostile elements in the host brain, resulting in poor survival (Bliss et al., 2007; Chen et al., 2011). Here, we conducted side-by-side comparison of transplanting human cerebral organoids versus dissociated human NPC that were derived from the same batch of hESCs and into identical cortical locations. We demonstrated enhanced survival of organoid grafts as compared to NPC transplants, thus supporting the notion that grafting donor cells in tissue-like cerebral organoids with 3D cytoarchitecture is an essential step to enhance graft viability by providing a protective shield against hostile host elements.

It is noteworthy of the technically challenges of the current study, which entails creating a large cavity in a small host mouse brain and grafting a relatively large human cerebral organoid; in comparison, stereotactic injection of dissociated stem cells is less traumatic. The use of adult mice as recipients may ease some of the technical difficulties, and a recent study successfully demonstrated intracranial transplant of human cerebral organoids into adult mice brains (Mansour et al., 2018). However, the different study design of the two studies and the different time frame of post-transplant analyses warrant careful comparison regarding the potential influences of transplant location, the age of the host mice, SCID versus immunocompetent recipient mice on the survival and development of organoid grafts. Despite these differences, many similar findings emerge from the two studies that echo each other. First, both studies found extensive vascularization from host brain into organoid grafts by 14 d after transplantation. Second, engraftment speed and expansion of the organoid grafts appeared comparable in both studies with overall reduction in graft size from day 0 to day 14 before vascularization takes place. Third, in both studies, human organoid grafts contained Iba1+ microglia that were of host origin. Four, both studies found limited programmed cell death in the organoid grafts in contrast to the massive apoptosis seen in stage-matched organoids in culture, which may be a result of robust vascularization and in vivo phagocytic clearing. Fifth, the time frames of neuronal and glial differentiation in engrafted cerebral organoids were similar in both studies, with scant axonal processes and low number of astrocyte differentiation by 14 d, which then increased in abundance over time.

A good vascularization is key for survival and development of cerebral organoids (Casper et al., 2003; Bates et al., 2009; Mansour et al., 2018). Remarkably, two weeks after transplantation, both NPC and organoid transplants have already attracted robust penetration of host vessels into the grafts, suggesting abundant vascular growth factors released from neuroprogenitors in the grafts. It also indicates that the poor survival of NPC transplants after four weeks cannot be simply explained by lack of vascularization. One major difference between the two graft types is the much more organized cellular arrangement and tissue-like cytoarchitecture in organoid transplants, in contrast to the random organization and dissociated state of donor cells in NPC transplants. Indeed, earlier studies showed that transplantation of tissue pieces results in better survival compared to cell suspension grafts (Clarkson et al., 1998; Sekine et al., 2011). 3D, cell–cell support from multiple cell types, and cell–matrix interaction all help to improve the success of tissue transplants (Tate et al., 2009). Hence, akin to tissue-based transplantation, cerebral organoid transplants likely confer better protection and trophic support for donor cells as a result of proper cytoarchitecture. Furthermore, donor cells in the organoid transplants continue to benefit from exposure to intrinsic patterning cues that are concordant with developmental stages, while the stem cell niches in the progenitor zone of cerebral organoids can provide proper biomechanical properties and spatial cues important for donor cell survival, proliferation, and lineage progression. This is reflected by a steady fraction of proliferative cells and enhanced multilineage neurodifferentiation in the organoid transplants. In addition, as we used P8–P10 mice as recipients and relatively large sized organoid transplants, the survival of our organoid grafts may have also benefited from spatial proximity to the lateral ventricles and the retrosplenial cortex with rich vascularized surfaces (Mansour et al., 2018).

NSC therapies have been shown to improve functional outcome in a variety of CNS disease models, and the benefits have largely been attributed to trophic support or immunomodulation (Pluchino and Martino, 2008; Feinstein et al., 2013). In cerebral organoid transplants, we...
detected a sustained NSC pool, which by itself may significantly contribute to tissue repair. It is worth noting that the spatial organization of neuroprogenitors along ventricles in the cerebral organoids was largely lost by day 14 following transplantation, as shown by the scattering patterns of Sox2+ cells and Ki67+ cells throughout the grafts. Regardless, compared to NPC transplanted in suspension, donor neuroprogenitor cells in organoid grafts would still benefit from a microenvironment with 3D and in vivo-like cytoarchitecture. Regarding neurodifferentiation, there is increased progeny cells of neuronal lineage in the organoid grafts with DCX+ neuroblasts and CTIP2+ cortical neurons. Astrocyte differentiation in the organoid transplants appears increased between two and four weeks after grafting, consistent with the developmental timeline of gliogenesis, which occurs after the main phase.

Figure 9. Differentiation of cerebral organoid transplants. A, Representative immunofluorescence images showing cells expressing intermediate progenitor marker TBR2 in the engrafted cerebral organoids at two and four weeks after transplantation. Notice no TBR2+ cells were observed in host cortical tissue and a slight decline of the number of TBR2+ cells in the hMito+ organoid grafts from two to four weeks after transplantation. B, Representative immunofluorescence images showing cells expressing deep layer neuronal marker CTIP2 in the engrafted cerebral organoids at two and four weeks after transplantation. Notice CTIP2+ cells in neighboring host cortex (white arrows), while in the hMito+ organoid grafts, there was an increase of CTIP2+ cells from two to four weeks after transplantation. C, Representative immunofluorescence images show expression of NF-H in the transplanted organoids after two and four weeks. White arrowheads denote colocalization of NH-H and hMito in organoid transplants. Dashed white lines delineate the graft areas. D: dorsal, V: ventral, M: medial, L: lateral.
of neurogenesis (Ge et al., 2012). The number of GFAP+ cells in organoid grafts appears more abundant than in stage-matched organoid cultures (Fig. 7B). This may reflect stronger glial differentiation signals at the transplant site in vivo than in cultured organoids. Concordantly, as compared to in vitro cerebral organoid cultures where oligodendrocytes were detected only at later stages (Renner et al., 2017; Matsui et al., 2018), we detected ~10% donor cells expressing Olig2 at both two and four weeks after grafting. Of note, during human neurodevelopment, Olig2 is expressed in OPC but also in a subset of neuroprogenitors (Jakovecvski and Zecvici, 2005), thus the Olig2+ cells in the organoid grafts may represent both populations.

Organoid transplantations have been examined for different target organs, including lung organoids (Dye et al., 2016), photoreceptor organoids (Santos-Ferreira et al., 2016), and liver organoids (Nie et al., 2018), all of which demonstrated good engraftment and maturation of transplanted organoids. In the current study, we limited the timeframe of our analyses to two and four weeks after transplantation, as immunocompetent P8–P10 mice were used with no immunosuppressive treatment. To investigate later time points, SCID mice or immunosuppression will be necessary. Mansour et al. (2018) recently detailed neurodevelopment in transplanted cerebral organoids up to 180 d after transplant, at which time extensive axonal growth and graft-to-host axonal projections, as well as functional synapses were observed. In both the current study and the study by Mansour et al. (2018), organoids were grown in culture for 40–50 d from hESCs (approximately four weeks in Matrigel routine condition) with largely dorsal forebrain specification, transplanting organoids of different maturity or of different regional specification by varying culture period or including different morphogen are worthwhile in future studies. Recent advancement allowed generation of region-specific neural organoids with features of neocortex (Lee et al., 2017), telencephalon (Watanabe et al., 2005), cerebellum (Muguruma et al., 2015), neural tube (Ranga et al., 2016), hippocampus (Sakaguchi et al., 2015), and neural retina (Kuwahara et al., 2015), among others (Wei et al., 2017). These specialized neural organoids open the door for further testing of homotypic transplants wherein region-specific cerebral organoids are transplanted into the corresponding region in host brain. In this regard, an earlier study showed substantially different set of efferent axon projections from transplants of homotypic embryonic motorcor-tex versus heterotopic embryonic visual cortical tissue into adult motor cortex (Gaillard et al., 2007). Additional studies are needed to control stem cell proliferation, as iPSC derivatives may have oncogenic potential even after differentiation (Blum and Benvenisty, 2008; Ghosh et al., 2011). Finally, ethical questions need to be discussed concerning the use and further development of more complex cerebral organoids (Lavazza and Massimini, 2018).

In summary, our study demonstrated enhanced survival and robust vascularization of human cerebral organoid transplants in lesioned frontoparietal cortex of mouse hosts. Cerebral organoid transplantations may offer a promising novel cell replacement strategy for repair of CNS injury and neurodegenerative disorders as they provide a large set of neural cell types with both neuroprogenitors and differentiated neurons in a structured organization similar to the targeted brain area.

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