Axonal Extensions along Corticospinal Tracts from Transplanted Human Cerebral Organoids

Takahiro Kitahara,1,2 Hideya Sakaguchi,1,3,* Asuka Morizane,1 Tetsuhiro Kikuchi,1 Susumu Miyamoto,2 and Jun Takahashi1,2,*

1Department of Clinical Application, Center for iPS Cell Research and Application, Kyoto University, Kyoto 606-8507, Japan
2Department of Neurosurgery, Kyoto University Graduate School of Medicine, Kyoto 606-8507, Japan
3Present address: RIKEN BDR-Otsuka Pharmaceutical Collaboration Center, Kobe 650-0047, Japan

*Correspondence: jbtaka@cira.kyoto-u.ac.jp (J.T.), hideya.sakaguchi@riken.jp (H.S.)
https://doi.org/10.1016/j.stemcr.2020.06.016

SUMMARY

The reconstruction of lost neural circuits by cell replacement is a possible treatment for neurological deficits after cerebral cortex injury. Cerebral organoids can be a novel source for cell transplantation, but because the cellular composition of the organoids changes along the time course of the development, it remains unclear which developmental stage of the organoids is most suitable for reconstructing the corticospinal tract. Here, we transplanted human embryonic stem cell-derived cerebral organoids at 6 or 10 weeks after differentiation (6w- or 10w-organoids) into mouse cerebral cortices. 6w-organoids extended more axons along the corticospinal tract but caused graft overgrowth with a higher percentage of proliferative cells. Axonal extensions from 10w-organoids were smaller in number but were enhanced when the organoids were grafted 1 week after brain injury. Finally, 10w-organoids extended axons in cynomolgus monkey brains. These results contribute to the development of a cell-replacement therapy for brain injury and stroke.

INTRODUCTION

Treatment for neurological sequelae caused by brain injury or stroke is at present limited to medication and rehabilitation, which have some therapeutic effects but cannot provide fundamental cures. On the other hand, the reconstruction of lost neural circuits by a cell-replacement strategy is expected to be a curative therapy for neurological deficits after cerebral cortex injury (Grade and Gotz, 2017; Lindvall and Kokaia, 2011).

The transplantation of embryonic cerebral cortex tissues is an effective approach to reconstructing damaged neural circuits. Previous studies reported that the transplantation of cerebral cortex tissue of mouse embryo into the cerebral cortex of rodents provided axonal projections from the graft to the host brain and spinal cord, synaptic connections between the graft and the host, and functional recovery of the grafted host (Ballout et al., 2016; Gaillard et al., 2007; Peron et al., 2017). Other studies described the transplantation of human pluripotent stem cell (hPSC)-derived neural stem cells, neuronal progenitor cells, or neurons into rodent cerebral cortices to reconstruct neural circuits (Espuny-Camacho et al., 2013, 2018; Palma-Tortosa et al., 2020; Tornero et al., 2013, 2017).

Cerebral organoids are stem cell-derived structures generated in vitro and display the 3D architecture and physiology of the cerebrum (Eiraku et al., 2008; Kadoshima et al., 2013; Lancaster et al., 2013; Sakaguchi et al., 2019). They are potential tools for modeling cerebral-related disorders, such as Zika virus-related microcephaly, and for developing treatments (Cugola et al., 2016; Dang et al., 2016; Garcez et al., 2016; Qian et al., 2016; Watanabe et al., 2017). Since cerebral organoids have a developmental process similar to that of embryonic cerebral cortex, hPSC-derived cerebral organoids can be a cell source for the restoration of lost neural circuits via transplantation. The transplantation of hPSC-derived cerebral organoids into mouse cerebral cortex has been evaluated for the vascularization of the organoids, the rates of graft survival, and the neuronal differentiation after transplantation (Daviaud et al., 2018; Mansour et al., 2018).

Cerebral organoids recapitulate the process of neurogenesis in the development of the cerebral cortex (Heide et al., 2018; Quadrato and Arlotta, 2017; Suzuki and Vanderhaeghen, 2015). In vivo, cortical neurogenesis is initiated by rounds of asymmetric divisions of progenitor cells called radial glial cells (RGCs), which form the ventricular zone (VZ) in the apical side of the cerebral cortex. After an asymmetrical division of RGCs, one daughter cell remains as an RGC in the VZ, and the other becomes a neuron or an intermediate progenitor cell. Neurons that migrate out of the VZ form the cortical plate on the basal side of the VZ. Different types of neurons are thought to be generated temporally, and late-born neurons migrate to the basal side of early-born neurons, which results in the unique layer formation of the cerebral cortex with an inside-out pattern (Dwyer et al., 2016; Greig et al., 2013; Kwan et al., 2012; Toma and Hanashima, 2015). The deep layer of the cerebral cortex is formed earlier and contains subcerebral projection neurons (SCPNS), which extend axons to the outside of the cerebral cortex, such as the brainstem and the spinal cord. The upper layer is formed later and composed of callosal projection neurons.
(CPNs), which extend axons to other areas of the cerebral cortex (Leyva-Diaz and Lopez-Bendito, 2013; Molyneaux et al., 2007; Toma and Hanashima, 2015). In mouse embryo, SCPNs are generated at approximately embryonic days (E) 12.5–14.5, and CPNs are generated at approximately E13.5–E16.5 (Fame et al., 2011; Hevner et al., 2003; Kwan et al., 2012). Previous studies describing the transplantation of embryonic cerebral cortex generally used E14–E15 mouse cerebral cortex tissues as the grafts (Ballout et al., 2016; Gaillard et al., 2004, 2007; Peron et al., 2017).

Because the cellular compositions of embryonic cerebral cortex and cerebral organoids dramatically change along the time course of the development, differences in the developmental stages of the graft could greatly affect the results of the transplantation. However, the effect of these differences has not been compared in the transplantation of embryonic cerebral cortex and cerebral organoids. Furthermore, in cerebral organoid transplantation, the effect of the transplantation site, the effect of the host brain environment, and the applicability of the transplantation to primates have not been investigated.

In this study, to clarify which developmental stage of cerebral organoids is suitable for transplantation, we transplanted human embryonic stem cell (hESC)-derived cerebral organoids at early stage (the time of SCPN generation) and late stage (the time of CPN generation) into mouse cerebral cortex and compared the graft survival and axonal extension to the host. We also evaluated the effects of the transplantation site and host brain environment as well as the transplantation of cerebral organoids into the motor cortex of nonhuman primate brain.

RESULTS

Characterization of Cerebral Organoids at Different Developmental Stages

For the induction of cerebral organoids, we adopted the serum-free floating culture of embryoid body-like aggregates with a quick reaggregation method (Eiraku et al., 2008; Kadoshima et al., 2013) and our previous protocol (Sakaguchi et al., 2019) (Figure 1A). A total of 9,000 dissociated hESCs were put into each well of low-cell-adhesion-coated 96-well plates, where they formed aggregates efficiently after the initiation of differentiation. The formation of several cricoid neuroepithelium-like structures in the aggregates was observed in bright-field images (Figure 1B). Immunohistochemistry (IHC) analysis of the cerebral organoids at 6 weeks after the initiation of differentiation (6w-organoid) showed the expression of the telencephalic marker FOXG1 (Figure 1C). 6w-organoids contained the VZ, which was composed of RGC-like cells positive for the dorsal telencephalic progenitor marker PAX6, and cells positive for the neuronal marker MAP2 surrounded the VZ (Figure 1C). The proliferation marker Ki67 was expressed in many cells in the VZ, validating the proliferative activity of RGC-like cells in the VZ (Figure 1C). Cells positive for the SCPN marker CTIP2 had commenced to form the layer corresponding to the cortical plate while cells positive for the CPN marker SATB2 were not observed (Figure 1C).

In cerebral organoids at 10 weeks after the initiation of differentiation (10w-organoids), the size of neuroepithelium-like structures was larger compared with those of 6w-organoids (Figure 1B). IHC analysis of the 10w-organoids showed that the thickness of the total epithelium was also increased (6w, 142.1 ± 33.3 μm; 10w, 285.1 ± 28.8 μm) (Figure 1D). The increase in thickness was more prominent in the layer of CTIP2+ cells than in the layer of PAX6+ cells, which suggested the robust generation of SCPNs by asymmetric cell division (PAX6+ layer, 89.8 ± 17.0 μm [6w] and 96.9 ± 8.1 μm [10w]; CTIP2+ layer, 29.3 ± 8.5 μm [6w] and 127.7 ± 22.9 μm [10w]) (Figure 1D). The percentages of PAX6+ cells and Ki67+ cells in total cells of the organoid were lower in 10w-organoids than in 6w-organoids (PAX6, 44.7% ± 4.8% [6w] and 24.6% ± 7.1% [10w]; Ki67, 34.5% ± 5.6% [6w] and 12.5% ± 5.4% [10w]) (Figure 1E). In contrast, the percentage of CTIP2+ cells was higher in 10w-organoids than in 6w-organoids (6w, 25.5% ± 6.3%; 10w, 46.8% ± 9.3%) (Figure 1E). SATB2+ cells, which were absent in 6w-organoids, were evident in 10w-organoids (6.2% ± 5.1%) (Figure 1E). When dividing cells were labeled with 5-ethynyl-2′-deoxyuridine (EdU) in 6w-organoids, the EdU+ cells contained 5.9% ± 2.2% of CTIP2+ cells but no SATB2+ cells 1 week after the labeling. In contrast, when dividing cells were labeled in 10w-organoids, the EdU+ cells contained more SATB2+ cells (2.9% ± 1.0%) than CTIP2+ cells (1.8% ± 0.1%) (Figure S1).

These results showed that our culture system could recapitulate the cerebral cortical developmental process, and that 6w-organoids were at a developmental stage before the initiation of CPN generation (early stage) and 10w-organoids were at a developmental stage after the initiation of CPN generation (late stage). We used early- and late-stage organoids as grafts for the transplantations below.

Graft Overgrowth after Transplantation Was Obvious in 6w-Organoids

Next, to evaluate whether the difference in the developmental stage of the organoids and the difference in the site of transplantation affected engraftment, we transplanted 6w- or 10w-organoids into the brain of 7-day-old (P7) mice. The organoids at both stages were cut into pieces with 1 mm diameter, each containing 1.9 × 10⁵ cells on average, and the pieces were implanted into the bilateral frontal and parietal cortices (Figures 2A–2C).
At 12 weeks post-transplantation (wpt), IHC analysis with staining of human neural cell adhesion molecule (hNCAM) revealed the engraftment of 6w- and 10w-organoids in the mouse cerebral cortex (Figure 2D). 6w-organoids enlarged massively to protrude from the transplanted cavities and spread over the surface of the cerebral cortex. On the other hand, 10w-organoids did not cause massive overgrowth of the grafts (Figure 2D). Both 6w- and
10w-organoids showed high graft survival rates in both the frontal and parietal cortices (6w, 100%; 10w, 87.5% [frontal] and 93.8% [parietal]). The total graft volume in each mouse at 12 wpt was significantly larger in 6w-organoids than in 10w-organoids (6w, 34.7 ± 9.4 mm³; 10w, 4.9 ± 2.3 mm³) (Figure 2E), but with no difference in the graft volume depending on the site of transplantation (Figure 2F).

There was no difference in the volume of the host brain irrespective of the culture period of the organoids (6w, 190.5 ± 19.5 mm³; 10w, 196.8 ± 8.6 mm³), suggesting that the grafts grew independent of the developmental process of the brain. Both organoids included endothelial cell marker CD31-positive vessel-like structures, suggesting that the cerebral organoids were vascularized from the host environment after the transplantation (Figure 3A). There was no difference in the percentage of the CD31+ area to total graft area (6w, 0.69% ± 0.07%; 10w, 0.72% ± 0.15%) (Figure 3B).

In summary, cerebral organoids at both developmental stages had high rates of graft survival with vascularization, but the risk of graft overgrowth appeared higher in 6w-organoids.

Figure 2. Graft Overgrowth after Transplantation was Obvious in 6w-Organoids
(A) Bright-field images of 6w- and 10w-organoids before and after cutting into pieces. Scale bars, 1 mm.
(B) The number of cells contained in 1-mm pieces of 6w- and 10w-organoids. n = 6 for both organoids (n, number of aggregates). ns, not significant; Mann-Whitney test. Results are presented as the mean ± standard deviation.
(C) Schematic of the procedure for the transplantation of cerebral organoids into the bilateral frontal and parietal cortices of 7-day-old mice (one piece into each cavity).
(D) Immunohistochemistry for hNCAM in sagittal and coronal sections of mouse brains at 12 wpt. Upper left, lateral 1.08 mm; lower left, lateral 1.44 mm; right, Bregma +1.26 mm. Scale bars, 1 mm.
(E) The total volume of all survived grafts in each mouse at 12 wpt. n = 8 for each organoid (n, number of mice). ***p < 0.001, Mann-Whitney test. Results are presented as the mean ± standard deviation.
(F) The total volume of all survived grafts in the frontal or parietal cortices of each mouse at 12 wpt. n = 8 for each organoid (n, number of mice). ns, not significant; Wilcoxon test. Results are presented as the mean ± standard deviation.

See also Figure S5.
Cerebral Projection Neurons Were Observed in Engrafted Cerebral Organoids, but 6w-Organoids Contained a Higher Percentage of Proliferative Cells

Next, we evaluated whether the overgrowth of 6w-organoids after transplantation was related to their proliferative activity. In 6w-organoids at 12 wpt, the existence of PAX6+ cells was obvious, and some of these cells co-expressed KI67 (Figure 4A). In contrast, in 10w-organoids at 12 wpt, PAX6+ cells were only present sparsely, and few KI67+ cells were observed (Figure 4A). The percentages of PAX6+ cells and KI67+ cells in human nuclei (hNuclei)-positive cells were significantly higher in 6w-organoids than in 10w-organoids (PAX6, 18.5% ± 3.8% [6w] and 9.8% ± 1.8% [10w]; KI67, 5.5% ± 1.0% [6w] and 2.0% ± 0.6% [10w]) (Figures 4B, S2A, and S2B), indicating that 6w-organoids contained a significantly higher percentage of proliferative cells than 10w-organoids not only at the time of transplantation but also at 12 wpt.

To reconstruct host neural circuits by cerebral organoid transplantation, projection neurons are required. Therefore, we next evaluated whether projection neurons existed in the engrafted cerebral organoids. Both 6w- and 10w-organoids contained CTIP2+ cells and SATB2+ cells at 12 wpt (Figure 4C), and both contained more SATB2+ cells than CTIP2+ cells in hNuclei+ cells (Figures 4D and S2C). The percentage of CTIP2+ cells in hNuclei+ cells was significantly higher in 6w-organoids (6w, 15.3% ± 2.9%; 10w, 4.4% ± 1.3%). In contrast, the percentage of SATB2+ cells in hNuclei+ cells was significantly higher in 10w-organoids (6w, 48.8% ± 2.3%; 10w, 63.8% ± 8.0%).

In summary, both 6w- and 10w-organoids that engrafted in the cerebral cortex contained graft-derived projection neurons, including CTIP2+ SCPNs and SATB2+ CPNs, but 6w-organoids contained a higher percentage of proliferative cells, which might explain the graft overgrowth.

6w-Organoids Extended a Larger Number of Axons along the Host Corticospinal Tract

Since cerebral organoids that engrafted in the mouse cerebral cortex contained projection neurons, we investigated how the engrafted cerebral organoids extended axons to the host brain. Both 6w- and 10w-organoids in the frontal cortex extended hNCAM+ fibers to the surrounding cerebral cortex, the corpus callosum (CC), and the striatum (Str) of the host (Figure 5A). In addition, both organoids that engrafted in the parietal cortex extended hNCAM+ fibers to the surrounding cerebral cortex and CC of the host (Figure 5B).

When we evaluated the extension of graft-derived axons along the corticospinal tract (CST) of the host, hNCAM+ fibers from both organoids reached the internal capsule (IC), the cerebral peduncle (CP), and the cervical spinal cord (SC), confirming axonal projections along the host CST (Figure 5C). In Str, hNCAM+ fibers were extended along myelin basic protein-positive fasciculated bundles, in which SCPN axons are known to travel in rodents (Figure S3) (Sano et al., 2017). There was no obvious difference in the direction of axonal extensions between 6w- and 10w-organoids, but 6w-organoids provided more fibers along the host CST than 10w-organoids (Figure 5D). It is possible that the higher number of axons was due to a higher number of surviving cells in the grafts derived from 6w-organoids. Therefore, we normalized the number of axons to the number of hNuclei+ and CTIP2+...
cells in the grafts. Although the number of axons per hNuclei+ cell was higher in 6w-organoids (Figure S4A), there was no difference in the number of axons per hNuclei+CTIP2+ cell in the Str, CP, and SC (Figure S4B). These data suggested that both 6w- and 10w-organoids had the ability to provide axonal extensions along the host CST but the number of axons was higher in 6w-organoids, and that CTIP2+ cells in the graft contributed to the axonal extension along the CST.

6w-Organoids Extended More Axons but Caused Overgrowth in Injured Adult Brain

Above, we observed that cerebral organoids extended axons along the CST in the intact brains of P7 mice, but these effects were not confirmed in adult or injured brains. Therefore, we next transplanted 6w- and 10w-organoids into the frontal cortex of 6-week-old mice 1 week after lesioning (Figure S5A).

Figure 4. Cerebral Projection Neurons Were Observed in Engrafted Cerebral Organoids, but 6w-Organoids Contained a Higher Percentage of Proliferative Cells

(A) Immunohistochemistry for hNuclei, PAX6, and KI67 in 6w- and 10w-organoids engrafted in the cerebral cortex (Cx). Right column is a magnification of the white squares. Scale bars, 500 μm (left and middle columns) and 20 μm (right column).

(B) Percentages of PAX6+ cells and KI67+ cells in hNuclei+ cells in the engrafted 6w- and 10w-organoids. n = 8 for each organoid (n, number of mice). ***p < 0.001, Mann-Whitney test. Results are presented as the mean ± standard deviation.

(C) Immunohistochemistry for CTIP2, SATB2, and PAX6 in the engrafted 6w- and 10w-organoids. Right column is a magnification of the white squares. Scale bars, 500 μm (left column) and 20 μm (right column).

(D) Percentages of CTIP2+ cells and SATB2+ cells in hNuclei+ cells in the engrafted 6w- and 10w-organoids. n = 8 for each organoid (n, number of mice). ***p < 0.001, Mann-Whitney test. Results are presented as the mean ± standard deviation.

See also Figures S2 and S5.
Figure 5. 6w-Organoids Extended More Axons along the Host CST
(A) Immunohistochemistry for human hNCAM after the transplantation of 6w- and 10w-organoids into the frontal cortex (sagittal section) shows graft-derived axonal extensions to the cerebral cortex (Cx), corpus callosum (CC), and striatum (Str). Scale bars, 200 μm.
(B) Immunohistochemistry for hNCAM after the transplantation of 6w- and 10w-organoids into the parietal cortex (coronal section) shows graft-derived axonal extensions to the Cx and CC. Scale bars, 200 μm.

(legend continued on next page)
motor area, and a 1-mm piece of organoids was implanted 1 week later.

The graft volume at 12 wpt was significantly larger in 6w-organoids than in 10w-organoids (6w, 27.0 ± 17.7 mm$^3$; 10w, 6.7 ± 5.2 mm$^3$) (Figures S5B and S5C). So too were the percentages of Pax6$^+$ cells and Ki67$^+$ cells in hNucl$^+$ cells (PAX6, 13.6% ± 5.1% [6w] and 7.0% ± 3.4% [10w]; Ki67, 4.7% ± 1.1% [6w] and 2.6% ± 1.1% [10w]) (Figures S5D, S5E, and S6A). The percentage of CTIP2$^+$ cells in hNucl$^+$ cells was significantly higher in 6w-organoids, while that of SATB2$^+$ cells was higher in 10w-organoids (CTIP2, 21.2% ± 8.5% [6w] and 8.3% ± 2.9% [10w]; SATB2, 48.7% ± 9.5% [6w] and 66.6% ± 7.0% [10w]) (Figures S5F, S5G, and S6A). In addition, we observed extensions of hNCAM$^+$ fibers along the host CST in both groups (Figure S5H), but 6w-organoids provided a significantly larger number of axons to the host IC and CP (Figure S5I).

In summary, as in the case of transplantation into P7 mice, 6w-organoids extended a higher number of axons along the host CST but caused graft overgrowth with more proliferative cells in the injured brain of 6-week-old mice.

**Axonal Extensions from 10w-Organoids Were Enhanced by Delayed Transplantation after Lesioning**

In the transplantation of embryonic cerebral cortex, transplantation 1 week after lesioning in the host cerebral cortex promotes axonal extensions from the graft to the host brain rather than transplantation immediately after lesioning (Peron et al., 2017). We therefore considered that if the cerebral organoids had properties similar to embryonic cerebral cortex, then a 1-week delay could enhance axonal extensions even using 10w-organoids, which provided less axonal growth compared with 6w-organoids. To test this hypothesis, we prepared two groups of transplantation of 10w-organoids into 6-week-old mice: transplantation immediately after lesioning (no-delay group) and 1 week after lesioning (1w-delay group) (Figure 6A), and compared the results of the transplantation between the two groups.

The graft survival rate at 12 wpt was higher in the 1w-delay group than the no-delay group (no-delay, 66.7%; 1w-delay, 100%). Furthermore, the graft volume was significantly larger in the 1w-delay group than in the no-delay group (no-delay, 1.1 ± 0.6 mm$^3$; 1w-delay, 6.4 ± 1.8 mm$^3$) (Figures 6B and 6C). However, there was no significant difference in the volume of the host brain (no-delay, 219.2 ± 15.5 mm$^3$; 1w-delay, 210.9 ± 10.5 mm$^3$), in the percentage of the CD31$^+$ area to total graft area (no-delay, 0.62% ± 0.20%; 1w-delay, 0.76% ± 0.13%) (Figures 6D and 6E), in the percentages of Pax6$^+$ cells and Ki67$^+$ cells in hNucl$^+$ cells (PAX6, 9.7% ± 3.3% [no-delay] and 10.5% ± 3.5% [1w-delay]; Ki67, 2.7% ± 1.6% [no-delay] and 1.9% ± 0.9% [1w-delay]) (Figures 6F, 6G, and S6B), or in the percentages of CTIP2$^+$ cells and SATB2$^+$ cells in hNucl$^+$ cells (CTIP2, 11.3% ± 5.1% [no-delay] and 10.4% ± 4.5% [1w-delay]; SATB2, 70.8% ± 4.1% [no-delay] and 60.9% ± 6.1% [1w-delay]) (Figures 6H, 6I, and S6B). In both groups, the grafts extended axons along the host CST (Figure 6J), but there was a significantly larger number of hNCAM$^+$ fibers in the 1w-delay group than in the no-delay group (Figure 6K).

In summary, as in the case of transplanting embryonic cerebral cortex, a 1-week delay between lesioning and transplantation enhanced axonal extensions from the 10w-organoids to the host brain, suggesting that neural circuit reconstruction after cerebral organoid transplantation could be promoted by modulating the environment of the host brain.

**Cerebral Organoids Extended Axons in the Brains of Nonhuman Primates**

Finally, we investigated whether the transplantation of cerebral organoids could be applied to primates. From an ethical point of view, we transplanted hESC-derived 10w-organoids, which had a lower risk of overgrowth that causes compression of the host brain than 6w-organoids, into the cerebral cortex of cynomolgus monkeys. In order not to affect the higher brain functions of the monkeys, we transplanted cerebral organoids only into the bilateral precentral gyrus, namely the primary motor cortex (Figure 7A).

10w-Organoids survived in the cerebral cortex bilaterally in two out of four monkeys at 12 wpt. In the other two monkeys, no graft survival was observed. Using magnetic resonance imaging (MRI), we identified engrafted tissues as regions with high intensity on T2-weighted images and with low intensity on T1-weighted images. There was...
A. Time course of 10w-organoid growth in the brain of mice with no delay or 1w-delay.

B. Images showing the graft volume with no delay or 1w-delay.

C. Graph depicting the graft volume with no delay or 1w-delay.

D. Images of hNCAM, hNCAM CD31, CD31, CD31(z-stack) with no delay or 1w-delay.

E. Graph showing the CD31 area/graft area with no delay or 1w-delay.

F. Images of hNuclei, PAX6, Ki67 with no delay or 1w-delay.

G. Graph showing the ratio of marker* cells/hNuclei* cells with no delay or 1w-delay.

H. Images of PAX6, CTIP2, SATB2 with no delay or 1w-delay.

I. Graph showing the ratio of marker* cells/hNuclei* cells with no delay or 1w-delay.

J. Images of Str, IC, CP with no delay or 1w-delay.

K. Graph showing the number of hNCAM* fibers with no delay or 1w-delay.

(legend on next page)
no obvious overgrowth of cerebral organoids in any monkey (Figure 7B).

Using IHC analysis, we detected the engraftment of hESC-derived cerebral organoids using the human-specific cytoplasm marker STEM121 (Figures 7B and 7C). Engrafted tissues entirely expressed the neuronal marker TUJ1 (Figure 7C). As the case of transplantation into mouse cerebral cortex, the engrafted tissues contained more SATB2+ cells than PAX6+ cells, Ki67+ cells, and CTIP2+ cells in monkey primary cortex (PAX6, 9.4%; Ki67, 2.7%; CTIP2, 6.1%; SATB2, 60.6%) (Figures 7C and 7D). We observed CD31+ vessel-like structures in the graft, suggesting that the cerebral organoids obtained vascularization in the monkey brains (Figures 7E and 7F). IHC analysis with a mouse-specific CD31 antibody and a human/monkey-specific CD31 antibody suggested that the blood vessels in the grafts were derived from the host brain (Figure S7). Engrafted 10w-organoids extended STEM121+ fibers to the brains of the host monkeys. The STEM121+ fibers reached the neighboring cerebral cortical area as well as the subcortical tissue. There were also extensions of STEM121+ fibers in the subcortical tissue both toward the CC and Str (Figure 7G), but there were no fiber extensions in more distal parts, such as the IC or CP.

Taken together, hESC-derived cerebral organoids survived in the cerebral cortex of monkeys and provided axonal extensions along the callosal projection and the subcerebral projection, which suggested that cerebral organoid transplantation could be a useful approach for reconstructing neural circuits after cerebral cortex injury in primates.

**DISCUSSION**

In this study, we compared the graft volume and axonal extensions along the host CST after transplanting hESC-derived cerebral organoids at two developmental stages. One was the early stage (6w) before the initiation of CPN generation (SCPN-generating stage), and the other was the late stage (10w) after the initiation of CPN generation (CPN-generating stage). We found that the 6w-organoids extended a larger number of axons along the host CST, but that they also showed graft overgrowth. In addition, axonal extensions were enhanced in the brain 1 week after brain injury compared with just after the injury. Finally, cerebral organoids survived and extended axons after transplantation into monkey brain.

There are two reports regarding the transplantation of human cerebral organoids into animal brains, but axonal extensions from the organoids along the host CST have never been described. One report showed the vascularization of the organoids with integration to the host brain after transplantation in the mouse retrosplenial cortex (Mansour et al., 2018). The other compared 3D cerebral organoids and 2D culture-derived cells in terms of graft survival and neuronal differentiation in the mouse frontoparietal cortex (Daviaud et al., 2018). In this study, we demonstrated that human cerebral organoids grafted in the mouse frontal cortex extended axons along the CST, suggesting their potential for reconstruction of the host motor pathway.
Figure 7. Transplantation of 10w-Organoids into Monkey Cerebral Cortex

(A) Axial (left) and coronal (middle) T2-weighted images (T2WI) and coronal T1-weighted image (T1WI) immediately after the transplantation. Arrows mark the site of the transplantation. Scale bars, 5 mm.

(B) Coronal T2WI and T1WI at 12 wpt, and immunohistochemistry for STEM121 in coronal sections of monkey brain. Scale bars, 5 mm.

(C) Immunohistochemistry for STEM121, TUJ1, PAX6, CTIP2, and SATB2 in the engrafted tissue in monkey cerebral cortex. Scale bars, 1 mm (except inset) and 20 μm (inset).

(D) Percentages of PAX6+ cells, KI67+ cells, CTIP2+ cells, and SATB2+ cells in intragraft cells at 12 wpt. n = 2 (n, number of monkeys).

(E) Immunohistochemistry for TUJ1 and CD31 in the engrafted tissue and the host cerebral cortex. The right three columns show higher-magnification images of the boxed areas. Scale bars, 1 mm (leftmost column) and 100 μm (other columns).

(F) The percentage of the CD31+ area to total graft area. n = 2 (n, number of monkeys).

(G) Immunohistochemistry for STEM121 shows graft-derived axonal extensions to the cerebral cortex and in the subcortical tissues toward the corpus callosum and striatum. Scale bars, 5 mm (low magnification images) and 100 μm (high magnification images).

See also Figure S7.
In the cerebral cortex, SCPNs, which constitute the CST, express CTIP2 (Arlotta et al., 2005) but not SATB2 (Alcamo et al., 2008; Britanova et al., 2008; Fame et al., 2011). It is reported that in the transplantation of dissociated cerebral cortices from mouse embryos, graft-derived cells that extended axons along the host CST are also CTIP2+/SATB2- (Wuttke et al., 2018). In our study, 6w-organoids, which contained more CTIP2+ cells than did 10w-organoids at 12 wpt, extended more axons along the host CST, but there was no significant difference in the number of axons per graft-derived CTIP2+ cell in the Str, CP, or SC. These results suggest that CTIP2+ cells in the graft play a pivotal role in reconstruction of the CST. It remains unclear, however, what stage of cells in the organoids contributed to the axonal extensions. We suspect two candidates: one is projection neurons that already exist in the graft at the time of transplantation, and the other is immature cells that become projection neurons after the transplantation. In a previous report, graft-derived axons along the host CST were provided by cells that were already post-mitotic at the time of transplantation (Wuttke et al., 2018). In this study, compared with 10w-organoids, 6w-organoids contained less CTIP2+ cells at the time of transplantation, but provided more CTIP2+ cells with a larger number of axons after transplantation. This result might suggest that more post-transplant-generated CTIP2+ cells became projection neurons, which extended axons along the CST, compared with already-existing CTIP2+ cells.

In cell transplantation toward clinical application, graft overgrowth must be avoided because it causes compression or destruction of the host brain. However, the risk of graft overgrowth has not been investigated in cell transplantations aimed at cerebral cortex repair. We found that 10w-organoids, which are in the CPN-generating stage, have a lower risk of graft overgrowth. In previous studies describing the transplantation of embryonic cerebral cortex, E14–E15 mice, which corresponds to the CPN-generating stage, were used as the grafts (Ballout et al., 2016, 2019; Gaillard et al., 2004, 2007; Peron et al., 2017), which would explain why graft overgrowth did not become a problem in those studies.

The observed enhancement of the graft survival and axonal extensions of 10w-organoids by delaying the transplantation 1 week after the cortical resection suggests that adjusting the host brain environment is a possible approach to promoting axonal extensions from cerebral organoids. Our result is in agreement with a previous study that examined embryonic cerebral cortex (Peron et al., 2017). However, the mechanism through which the host brain 1 week after the resection supports cell survival and axonal extension is unknown. Previous studies suggest that the secretion of pro-angiogenic factors by cells surrounding the lesion (Skold et al., 2005) or the modulation of inflammatory responses (Ballout et al., 2019; Liauw et al., 2008) are candidates. It is reported that transplantation with 1-week delay provided a transient increase in graft vascularization 4 days after the transplantation (Peron et al., 2017). In our study, a 1-week delay did not cause an obvious difference in the graft vascularization at 12 wpt, but a transient increase in the graft vascularization might have contributed to the enhancement of axonal extensions.

There are substantial differences in the nervous systems between rodents and primates in terms of size and neuro-anatomical characteristics (Courtine et al., 2007). For example, in rodents, CST fibers descending in the spinal cord travel mostly in the dorsal columns and have no direct connections to motor neurons. In contrast, in primates, CST fibers in the spinal cord travel mostly in the lateral columns and have both direct and indirect connections to motor neurons (Courtine et al., 2007; Lemon, 2008). Such differences can limit the translation of the results obtained from rodents into treatment for human patients (Courtine et al., 2007). Therefore, when considering clinical application, primate transplantation models are necessary. One report more than 30 years ago transplanted allo-genic embryonic cerebral cortex tissues into the brain of rhesus monkeys, but no graft survival was observed in the cerebral cortex (Gopinath et al., 1989). The current study clearly showed that the transplantation of hPSC-derived cells into the primate cerebral cortex resulted in graft survival and axonal extensions. The axons, however, were observed only in the subcortical area, probably because a 12-week observation is too short when considering the size of monkey brains. Observation over longer periods is needed to confirm axonal extensions for longer distances.

Ethical issues need to be considered in experiments in which human cells are transplanted into the cerebral cortex of animals (Chen et al., 2019). In this study, to avoid the risk of manipulating the higher brain functions of monkeys by transplantation, we limited the transplantation site to the motor cortex. In addition, to avoid the risk of graft overgrowth compressing the monkey brain, only 10w-organoids were transplanted. Furthering this type of research will include ethical concerns, such as whether the animals can be humanized by creating brain chimeras and whether the cerebral organoids in vitro can have consciousness (Sawai et al., 2019).

Optimization of both the donor cells and host brain environment is critical for successful transplantations. We have shown that early-stage organoids extend more axons but cause graft overgrowth. Elimination of the proliferating cells by sorting may solve this problem (Samata et al., 2020). In parallel, the clarification and administration of supportive factors may enhance graft survival and axonal extensions. Stepwise solution of these issues will open...
the way to the realization of a cell-based therapy for brain injury and stroke.

**EXPERIMENTAL PROCEDURES**

**Animals**

All animal experiments described in this study were approved by the Institutional Animal Care and Use Committee of the Animal Research Facility at Kyoto University. All animals were cared for and handled in accordance with the Regulation on Animal Experimentation at Kyoto University.

SCID mice (C.B-17/lcrHsd-Ptkdscid, Shimizu Laboratory Supplies) 7 days (male and female, n = 16) and 6 weeks (male, n = 22) old, and purpose-bred male cynomolgus monkeys (*Macaca fascicularis*, Shin Nippon Biomedical Laboratories, n = 4) 3 years old, were used as the transplant recipients. All mice and monkeys were kept ad libitum access to food and water.

**Surgical Procedure for Mice**

All surgical procedures for mice were performed under anesthesia with isoflurane inhalation. In 7-day-old mice, cerebral organoids were transplanted into the bilateral frontal and parietal cortices immediately after making cavities in the cerebral cortices. Half of the mice (n = 8) received 6w-organoids and the other half (n = 8) received 10w-organoids. A skin incision was performed, and a 2 × 2 mm craniotomy windows were opened with bone flaps hinged on the lateral base. A cavity of 1 mm diameter and 1 mm depth was made by aspirating the cortical tissue in each craniotomy window. Cerebral organoids were cut into 1-mm-diameter pieces using micro-scissors (Bio Research Center, no. 16324319), and one piece was implanted into each cavity. The craniotomy window was closed by returning the bone flap, and the skin was sutured with 7-0 ETHILON (Ethicon). In 6-week-old mice, 1-mm pieces were transplanted into the right frontal cortex 1 week after (n = 16) or immediately after (n = 6) making the cavity.

**Surgical Procedure for Monkeys**

All surgical procedures for monkeys were performed under anesthesia with an intramuscular injection of ketamine (10 mg/kg) and xylazine (1 mg/kg). Bilateral precentral gyri were identified by preoperative MRI, and the coordinates of the targets were obtained. A midline skin incision was performed, and 10-mm burr holes were made above the bilateral precentral gyri based on the coordinates obtained from the preoperative MRI. The dura mater was incised, and a 2-mm cavity was made in the precentral cortex. 10w-Organoids were cut into 1-mm pieces, and 3, 5, or 11 pieces were implanted into each cavity. The dura mater and epicranial aponeurosis were sutured with 4-0 NUROLON (Ethicon), and the skin was sutured with 4-0 NESCOSUTURE (allresa).

An antibiotic (cefazolin sodium; Astellas Pharma) was administered for 3 days from the day of the operation (70 mg/body, intramuscularly). An immunosuppressant (Tacrolimus hydrate; Towa Pharmaceutical, prepared at 5 mg/mL in ethanol and polyoxyethylene hydrogenated castor oil 60) was administered from 1 day before transplantation until the day of euthanization (0.05–0.075 mg/kg, intramuscularly). The blood trough concentration of Tacrolimus was monitored and maintained at 10–25 ng/mL, with blood sampling once every week until the blood concentration reached within the target range followed by blood sampling at 4, 8, and 12 wpt.

**Data Acquisition**

Images were obtained using a confocal laser microscope (LSM710 and LSM800, Zeiss), except images for quantitative analyses and images of wide fields (Figures 2B, 5B, 6B, and 7B, left two images of 7C, and lower-magnification images of 7G), which were obtained using a confocal quantitative image cytometer (CQ1; Yokogawa).

**Statistical Analysis**

Statistical analyses were performed with PRISM software (GraphPad, version 8). Statistical significances in two-group comparisons were tested with the Mann-Whitney test (unpaired, nonparametric), except the comparison of graft volume in the mouse frontal and parietal cortices (Figure 2D), which was tested with the Wilcoxon test (paired, nonparametric). The results are expressed as the mean ± standard deviation.

**SUPPLEMENTAL INFORMATION**

Supplemental Information can be found online at https://doi.org/10.1016/j.stemcr.2020.06.016.

**AUTHOR CONTRIBUTIONS**

T. Kitahara, H.S., and J.T. designed the project. T. Kitahara and H.S. performed the research. A.M. and T. Kikuchi assisted with the experiments using monkeys. T. Kitahara and H.S. analyzed the data. T. Kitahara and H.S. wrote the original draft. J.T. reviewed and edited the manuscript. S.M. and J.T. supervised the research. H.S. and J.T. obtained grants for the research. All authors approved the final manuscript as submitted.

**ACKNOWLEDGMENTS**

The authors thank Takayoshi Matsubara (PerkinElmer Japan Co., Ltd. Informatics) for technical support in the data acquisition using CQ1 and in the data analysis using CellPathfinder; to Kei Kubota for technical support in experiments using monkeys; to Daiiuke Doi, Bumpel Samata, Sadaharu Torikoshi, Takafumi Shimogawa, and Ryosuke Tsuchimochi for taking care of the animals; to Towa Pharmaceutical for Tacrolimus hydrate; to Peter Karagiannis for critical reading of the manuscript; and to all members of the Takahashi lab for fruitful discussions. This work was supported by a grant from the Network Program for Realization of Regenerative Medicine from the Japan Agency for Medical Research and Development (AMED) (18bm0204004h0006) (to J.T.), Japan Society for the Promotion of Science (JSPS) KAKENHI grant no. 17H04302 (to J.T.), JSPS KAKENHI grant nos. 16H06897 and 18K15046 (to H.S.), and Grant-in-Aid for JSPS Fellows (JSPS PD research fellowship grant no. 17J10294) (to H.S.).

Received: December 12, 2019
Revised: June 16, 2020
Accepted: June 18, 2020
Published: July 16, 2020
REFERENCES

Alcami, A.E., Chirivella, L., Dautzenberg, M., Dobrevea, G., Farinas, L., Grosschedl, R., and McConnell, S.K. (2008). Satb2 regulates callosal projection neuron identity in the developing cerebral cortex. Neuron 57, 364–377.

Arlotta, P., Molyneaux, B.J., Chen, J., Inoue, J., Kominami, R., and Macklis, J.D. (2005). Neuronal subtype-specific genes that control corticospinal motor neuron development in vivo. Neuron 45, 207–221.

Ballout, N., Frappe, I., Peron, S., Jaber, M., Zibara, K., and Gaillard, A. (2016). Development and maturation of embryonic cortical neurons grafted into the damaged adult motor cortex. Front. Neural Circuits 10, 55.

Ballout, N., Rochelle, T., Brot, S., Bonnet, M.L., Francheteau, M., Prestoz, L., Zibara, K., and Gaillard, A. (2019). Characterization of inflammation in delayed cortical transplantation. Front. Mol. Neurosci. 12, 160.

Britanova, O., de Juan Romero, C., Cheung, A., Kwan, K.Y., Schwark, M., Gyorgy, A., Vogel, T., Akopov, S., Mitkovski, M., Agoston, D., et al. (2008). Satb2 is a postmitotic determinant for upper-layer neuron specification in the neocortex. Neuron 57, 378–392.

Chen, H.I., Wolf, J.A., Blue, R., Song, M.M., Moreno, J.D., Ming, G.L., and Song, H. (2019). Transplantation of human brain organoids: revisiting the science and ethics of brain chimeras. Cell Stem Cell 25, 462–472.

Courtine, G., Bunge, M.B., Fawcett, J.W., Grossman, R.G., Kaas, J.H., Lemon, R., Maier, I., Martin, J., Nudo, R.J., Ramon-Cueto, A., et al. (2007). Can experiments in nonhuman primates expedite the translation of treatments for spinal cord injury in humans? J. Neurosci. 37, 519–218.

Daviaud, N., Friedel, R.H., and Zou, H. (2018). Vascularization and engraftment of transplanted human cerebral organoids in mouse cortex. eNeuro 5, ENEURO.0219-18.2018.

Dwyer, N.D., Chen, B., Chou, S.J., Hippenmeyer, S., Nguyen, L., and Ghashghaei, H.T. (2016). Neural stem cells to cerebral cortex: emerging mechanisms regulating progenitor behavior and productivity. J. Neurosci. 36, 11394–11401.

Eiraku, M., Watanabe, K., Matsuo-Takasaki, M., Kawada, M., Yonemura, S., Matsumura, M., Watabe, T., Nishiyama, A., Muguruma, K., and Sasai, Y. (2008). Self-organized formation of polarized cortical tissues from ESCs and its active manipulation by extrinsic signals. Cell Stem Cell 3, 519–532.

Espuny-Camacho, I., Michelsen, K.A., Gall, D., Linaro, D., Hasche, A., Bonnefont, J., Bali, C., Orduz, D., Bilheu, A., Herpoel, A., et al. (2013). Pyramidal neurons derived from human pluripotent stem cells integrate efficiently into mouse brain circuits in vivo. Neuron 77, 440–456.

Espuny-Camacho, I., Michelsen, K.A., Linaro, D., Bilheu, A., Acosta-Verdugo, S., Herpoel, A., Giugliano, M., Gaillard, A., and Vanderhaeghen, P. (2018). Human pluripotent stem-cell-derived cortical neurons integrate functionally into the lesioned adult murine visual cortex in an area-specific way. Cell Rep. 23, 2732–2743.

Fame, R.M., MacDonald, J.L., and Macklis, J.D. (2011). Development, specification, and diversity of callosal projection neurons. Trends Neurosci. 34, 41–50.

Gaillard, F., Domballe, L., and Gaillard, A. (2004). Fetal cortical allografts project massively through the adult cortex. Neuroscience 126, 631–637.

Gaillard, A., Prestoz, L., Dumartin, B., Cantereau, A., Morel, F., Roger, M., and Jaber, M. (2007). Reestablishment of damaged adult motor pathways by grafted embryonic cortical neurons. Nat. Neurosci. 10, 1294–1299.

Garcez, P.P., Loiola, E.C., Madeira da Costa, R., Higa, L.M., Trindade, P., Delvecchio, R., Nascimento, J.M., Brindeiro, R., Tanuri, A., and Rehen, S.K. (2016). Zika virus impairs growth in human neurospheres and brain organoids. Science 352, 816–818.

Gopinath, G., Mahapatra, A.K., Bharadwaj, J.C., Banerji, R., Sharma, D.N., and Tandon, P.N. (1989). Transplantation of fetal neocortex in rhesus monkey. J. Biosci. 14, 255–260.

Grade, S., and Gotz, M. (2017). Neuronal replacement therapy: previous achievements and challenges ahead. NPJ Regen. Med. 2, 29.

Greig, L.C., Woodworth, M.B., Galazo, M.J., Padmanabhan, H., and Macklis, J.D. (2013). Molecular logic of neocortical projection neuron specification, development and diversity. Nat. Rev. Neurosci. 14, 755–769.

Heide, M., Huttner, W.B., and Mora-Bermudez, F. (2018). Brain organoids as models to study neocortex development and evolution. Curr. Opin. Cell Biol. 55, 8–16.

Hevner, R.F., Daza, R.A., Rubenstein, J.L., Stunnenberg, H., Olavarria, J.F., and Englund, C. (2003). Beyond laminar fate: toward a molecular classification of cortical projection/pyramidal neurons. Dev. Neurosci. 25, 139–151.

Kadoshima, T., Sakaguchi, H., Nakano, T., Soen, M., Ando, S., Eiraku, M., and Sasai, Y. (2013). Self-organization of axial polarity, inside-out layer pattern, and species-specific progenitor dynamics in human ES cell-derived neocortex. Proc. Natl. Acad. Sci. U S A 110, 20284–20289.

Kwan, K.Y., Sestan, N., and Anton, E.S. (2012). Transcriptional coordination of neuronal migration and laminar identity in the neocortex. Development 139, 1535–1546.

Lancaster, M.A., Renner, M., Martin, C.A., Wenzel, D., Bicknell, L.S., Hurles, M.E., Homfray, T., Penninger, J.M., Jackson, A.P., and Knoblich, J.A. (2013). Cerebral organoids model human brain development and microcephaly. Nature S01, 373–379.

Lemon, R.N. (2008). Descending pathways in motor control. Annu. Rev. Neurosci. 31, 195–218.

Leyva-Diaz, E., and Lopez-Bendito, G. (2013). In and out from the cortex: development of major forebrain connections. Neuroscience 254, 26–44.

Liuw, J., Hoang, S., Choi, M., Ergolu, C., Choi, M., Sun, G.H., Percy, M., Wildman-Tobriner, B., Bliss, T., Guzman, R.G., et al. (2008). Thrombospondins 1 and 2 are necessary for synaptic
plasticity and functional recovery after stroke. J. Cereb. Blood Flow Metab. 28, 1722–1732.

Lindvall, O., and Kokaia, Z. (2011). Stem cell research in stroke: how far from the clinic? Stroke 42, 2369–2375.

Mansour, A.A., Goncalves, J.T., Bloyd, C.W., Li, H., Fernandes, S., Quang, D., Johnston, S., Parylak, S.L., Jin, X., and Gage, F.H. (2018). An in vivo model of functional and vascularized human brain organoids. Nat. Biotechnol. 36, 342–441.

Molyneaux, B.J., Arlotta, P., Menezes, J.R., and Macklis, J.D. (2007). Neuronal subtype specification in the cerebral cortex. Nat. Rev. Neurosci. 8, 427–437.

Palma-Tortosa, S., Tornero, D., Gronning Hansen, M., Monni, E., Haji, M., Kartisvadze, S., Akty, S., Tsuyukov, O., Parmar, M., Deisseroth, K., et al. (2020). Activity in grafted human iPS cell-derived cortical neurons integrated in stroke-injured rat brain regulates motor behavior. Proc. Natl. Acad. Sci. U S A 117, 9094–9100.

Peron, S., Droguerre, M., Debarbieux, F., Ballout, N., Benoît-Marand, M., Francheteau, M., Brot, S., Rougon, G., Jaber, M., and Gaillard, A. (2017). A delay between motor cortex lesions and neuronal transplantation enhances graft integration and improves repair and recovery. J. Neurosci. 37, 1820–1834.

Qian, X., Nguyen, H.N., Song, M.M., Hadiono, C., Ogden, S.C., Hammack, C., Yao, B., Hamersky, G.R., Jacob, F., Zhong, C., et al. (2016). Brain-region-specific organoids using mini-bioreactors for modeling ZIKV exposure. Cell 165, 1238–1254.

Quadrato, G., and Arlotta, P. (2017). Present and future of modeling human brain development in 3D organoids. Curr. Opin. Cell Biol. 49, 47–52.

Sakaguchi, H., Ozaki, Y., Ashida, T., Matsuhara, T., Oishi, N., Kisara, S., and Takahashi, J. (2019). Self-organized synchronous calcium transients in a cultured human neural network derived from cerebral organoids. Stem Cell Reports 13, 458–473.

Samata, B., Takaichi, R., Ishii, Y., Fukushima, K., Nakagawa, H., Ono, Y., and Takahashi, J. (2020). L1CAM is a marker for enriching corticospinal motor neurons in the developing brain. Front. Cell. Neurosci. 14, 31.

Sano, N., Shimogawa, T., Sakaguchi, H., Iorio, Y., Miyawaki, Y., Morizane, A., Miyamoto, S., and Takahashi, J. (2017). Enhanced axonal extension of subcortical projection neurons isolated from murine embryonic cortex using Neuropilin-1. Front. Cell. Neurosci. 11, 123.

Sawai, T., Sakaguchi, H., Thomas, E., Takahashi, J., and Fujita, M. (2019). The ethics of cerebral organoid research: being conscious of consciousness. Stem Cell Reports 13, 440–447.

Skold, M.K., von Gertten, C., Sandberg-Nordqvist, A.C., Mathiesen, T., and Holmin, S. (2005). VEGF and VEGF receptor expression after experimental brain contusion in rat. J. Neurotrauma 22, 353–367.

Suzuki, I.K., and Vanderhaeghen, P. (2015). Is this a brain which I see before me? Modeling human neural development with pluripotent stem cells. Development 142, 3138–3150.

Toma, K., and Hanashima, C. (2015). Switching modes in corticogenesis: mechanisms of neuronal subtype transitions and integration in the cerebral cortex. Front. Neurosci. 9, 274.

Tornero, D., Watananait, S., Gronning Madsen, M., Koch, P., Wood, J., Tatarishvili, J., Mine, Y., Ge, R., Monni, E., Devaraju, K., et al. (2013). Human induced pluripotent stem cell-derived cortical neurons integrate in stroke-injured cortex and improve functional recovery. Brain 136, 3561–3577.

Tornero, D., Tsuyukov, O., Granmo, M., Rodriguez, C., Gronning-Hansen, M., Thelin, J., Smozhanik, E., Laterza, C., Watananait, S., Ge, R., et al. (2017). Synaptic inputs from stroke-injured brain to grafted human stem cell-derived neurons activated by sensory stimuli. Brain 140, 692–706.

Watanabe, M., Buth, J.E., Vishlaghi, N., de la Torre-Ubieta, L., Taxi-dis, J., Khakh, B.S., Coppola, G., Pearson, C.A., Yamauchi, K., Gong, D., et al. (2017). Self-organized cerebral organoids with human-specific features predict effective drugs to combat Zika virus infection. Cell Rep. 21, 517–532.

Wuttke, T.V., Markopoulous, F., Padmanabhan, H., Wheeler, A.P., Murthy, V.N., and Macklis, J.D. (2018). Developmentally primed cortical neurons maintain fidelity of differentiation and establish appropriate functional connectivity after transplantation. Nat. Neurosci. 21, 517–529.