Neurotrophic Factor-Laden Acellular Chondroitin Sulfate Scaffolds Promote Chronic Functional Recovery After Severe Traumatic Brain Injury

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ABSTRACT

Severe traumatic brain injury (sTBI) survivors experience permanent functional disabilities due to significant volume loss and the brain's poor capacity to regenerate. Chondroitin sulfate glycosaminoglycans (CS-GAGs) are key regulators of growth factor signaling and neural stem cell homeostasis in the brain. However, the efficacy of engineered CS (eCS) matrices in mediating structural and functional recovery after sTBI has not been investigated. We report that neurotrophic factor functionalized acellular eCS matrices implanted into the rat M1 region acutely post-sTBI, significantly enhanced cellular repair and gross motor function recovery when compared to controls, 20 weeks post-sTBI. Animals subjected to M2 region injuries followed by eCS matrix implantations, demonstrated the significant recovery of 'reach-tograsp' function. This was attributed to enhanced volumetric vascularization, activity-regulated cytoskeleton (Arc) protein expression, and perilesional sensorimotor connectivity. These findings indicate that eCS matrices implanted acutely post-sTBI can support complex cellular, vascular, and neuronal circuit repair, chronically after sTBI.

INTRODUCTION

Severe traumatic brain injuries (sTBI) caused by blunt force or penetrating trauma to the brain leads to extensive tissue loss and life-long disabilities¹. Although sTBIs account for only 10% of the approximately 1.7 million TBI cases reported in the United States annually, they are responsible for over 90% of all TBI associated costs¹. There are no clinical treatments to prevent cognitive impairments and tissue loss encountered after sTBI. Neural stem cells (NSCs) have long held a privileged position in neural repair strategies for their ability to mediate neuroprotective "bystander" signaling². However, inadequate control over NSC differentiation and the risk of immune rejection of xenografted NSCs pose serious limitations for their clinical application^{3,4}. Although autologous cell therapies could help mitigate some of these concerns⁵, issues related to poor survivability of transplanted cells⁶, and the risk of tumorigenesis in the case of human embryonic and pluripotent stem cells^{7,8}, pose significant barriers to clinical success.

When compared to cell therapies, the development of acellular brain-mimetic constructs that can potentiate neurotrophic factor mediated structural and functional brain repair has received little attention. Chondroitin sulfate proteoglycans (CSPGs) are major constituents of the brain extracellular matrix (ECM) and stem cell niche⁹⁻¹⁵. CSPG linked sulfated CS glycosaminoglycan (CS-GAG) side chains regulate growth factor signaling¹⁶⁻¹⁹, neuronal development²⁰, and neuroplasticity^{21,22}. Fibroblast growth factor (FGF2) and brainderived neurotrophic factor (BDNF) bind with high affinity to sulfated CS chains via specific sulfation motifs that act as molecular recognition sites²³. FGF-2 is expressed by NSCs and radial glial cells found in neurogenic niches of the brain²⁴, where it is known to promote neuroprotection²⁵, NSC migration and proliferation^{26,27}, and neurogenesis^{24,26,28}. BDNF is an important mediator of neuroplasticity at all stages of brain development²⁹, and is known to promote neuroprotection³⁰, and functional repair of the injured brain^{29,31,32}. FGF-2 and BDNF also promote angiogenesis after injury^{16,33-35}, and are critical for the maintenance of oxygen perfusion and tissue viability post-injury. In recent studies, acutely implanted 3D CS-GAG scaffolds with- and without encapsulated NSCs, enhanced the efficacy of endogenous NSCs and

mediated neuroprotection, four-weeks after sTBI³⁶. However, the chronic functional implications of acellular, neurotrophic factor functionalized eCS matrix implants post-sTBI, has so far not been assessed.

In this study, we conducted a systematic assessment of the long-term outcomes of implanting FGF-2 and BDNF-laden sulfated eCS matrices in rats subjected to sTBI. We performed magnetic resonance imaging (MRI) phase gradient analyses of regional cerebral blood flow (rCBF) and tissue vascularization in conjunction with longitudinal behavioral performance and functional recovery, followed by detailed terminal immunohistochemical assessments of tissue-specific biomarkers. We also quantified activity-related cytoskeletal associated protein (Arc) expression and volumetric vasculature tracing in cleared brain tissue, to localize newly formed functional clusters of neurons intra- and perilesionally in sTBI rats implanted with eCS scaffolds when compared to controls.

RESULTS

Acutely implanted eCS matrices promote chronic neuroprotection and recovery of gross motor function

In order to assess the chronic neuroprotective effects of neurotrophic factor-laden eCS matrix implants (Sup. Fig. 1), we performed controlled cortical impact (CCI)-induced sTBI in the caudal forelimb area (CFA) in rats and implanted eCS matrices 48h post-CCI in the lesion epicenter. We performed longitudinal assessments of motor function and MRI-based quantification of tissue volume loss, and conducted terminal histological quantification of injury volume (Fig. 1A).

While general locomotion was found to be similar across all groups (Sup. Fig. 2A), we observed that the performance of eCS matrix implanted rats was found to be comparable to Sham controls, whereas sTBI rats demonstrated significant motor deficits (Fig. 1B and Sup. Fig. 2B).

From results of longitudinal T2 weighted MRI at 4- and 20-week time points and lesion volume analyses (Figure 1C-D), as well as terminal Nissl staining of tissue and quantification at 20 weeks post-sTBI (Fig. 1E and F), we were able to detect the significantly greater tissue atrophy in the sTBI only animals when compared to eCS matrix implanted rats and Sham controls (Fig. 1D and 1F).

These results demonstrated that eCS matrix-implanted rats exhibited tissue and motor function recovery that was comparable to Sham control group animals chronically post-sTBI.

eCS matrices support enhanced proliferation of endogenous NSCs and neurotrophic factor expression

eCS matrices have been demonstrated to promote FGF-2-mediated neural progenitor homeostasis and neuroprotection four-weeks post-sTBI ^{17,37}. Considering these previous observations, we quantified the effects of neurotrophic factor-laden eCS matrices on the chronic maintenance of proliferating NSCs in the lesion site using Ki67 and Sox-1 biomarkers (Fig. 2A &C), and CS mediated FGF-2 retention

(Fig.2B,D&E). Endogenous NSC proliferation in brain tissue from eCS matrix-implanted rats was found to be comparable to Sham controls (Fig. 2C and Sup. Fig. 3A, C&D), but was significantly decreased in sTBI-only rats. Although a significant decrease in Ki67+ cells in tissue from eCS treated animals was observed when compared to Sham and TBI controls (Sup. Fig. 3A, B&E), the significantly increased percentage of Sox-1+ Ki67+ cells in eCS matrix treated animals when compared to TBI controls (Fig. 2C), suggests that NSCs do not contribute to the observed overall decrease in Ki67+ cells. These findings were further corroborated by FGF-2 expression levels (Fig. 2D), which followed a similar trend to Sox-1+ staining. CS56 antibody staining of implanted eCS matrices and endogenously produced CS-GAGs in the lesion site revealed no significant differences in CS56 staining between sTBI and eCS groups, but indicated a significantly greater CS56+ staining in both groups when compared to Sham control (Fig. 2E). Interestingly, qualitative differences in locational presence of CS56+ staining was observed throughout the lesion sites in eCS and sTBI-only animals. When compared to sTBI-only animals, which predominantly demonstrated CS56+ staining only in the lesion penumbra and indicated a complete lack of cellular presence in the lesion volume, eCS matrix-treated animals indicated a uniform presence of CS56+ staining throughout the lesion volume. The CS56+ staining of eCS matrix-implanted brain tissue indicates the potential residual presence of eCS matrix along with what appears to be endogenously produced CS-GAGs/CSPGs, 20-weeks post-TBI. These results suggest that eCS matrix implants might enhance NSC proliferation by facilitating the sustained expression and presence of FGF2 in the lesion site, 20-weeks postsTBI.

We used laser capture microdissection to isolate perilesional tissue labeled by rat IgG staining (Fig. 2F), along with any tissue present in the lesion cavity, in order to specifically identify regional changes in expression of *BDNF*, *FGF2*, *CXCL12*, and *CXCR4* transcripts (Fig.2G). We identified the significant and tissue-specific increase in expression levels of transcripts encoding neuroplasticity (*BDNF*) and neural progenitor proliferation (*FGF2*) factors, along with NSC homing (*CXCL12*, and *CXCR4*) transcripts in brain tissue explanted from eCS matrix-treated animals when compared to Sham and TBI controls (Fig. 2G;

> 2 fold, p < 0.05). Fold differences in target gene expression in eCS matrix implanted animals was compared to that of sTBI rats after normalizing to Sham group expression levels and endogenous controls (*GADPH*, and *HPRT1*).

eCS matrices promote chronic neurogenesis and neuroplasticity

Since cell proliferation, neuronal differentiation, and synaptic plasticity are indicators of normal brain tissue homeostasis ^{38,39}, we considered the occurrence of these processes within the lesion site as measures of functional neuronal network activity, tissue maintenance, and recovery. We quantified the presence of newly formed neurons using markers for immature/migrating neuroblasts (doublecortin, DCX+) and dividing (BrdU+) neuronal cells (Fig. 3A&B). We found a significantly increased number of dividing cells in the eCS-implanted rats when compared to Sham and sTBI groups (Sup. Fig. 4A-D). A similar trend was found for DCX+ cells (Sup. Fig. 4E) and DCX+/BrdU+/DAPI+ co-labeled cells in eCS-treated animals (Fig. 3B), demonstrating a significant increase in these markers when compared to Sham and TBI groups. These results indicate that eCS matrix implants might promote the differentiation of proliferating cells into new neurons at the lesion site up to 20 weeks post-sTBI.

Since synaptic vesicle presence is indicative of normal functional neurons and synaptic plasticity⁴⁰, we evaluated the potential change in synaptic plasticity at the 20-week time point using antibodies against the synaptic vesicle marker Synaptophysin-I (Syn; Fig. 3C). We observed a significant increase in Syn1+ signal in both DAPI+ (Sup. Fig. 5A-D; p = 0.014) and NeuN+ cells (Fig. 3D) in eCS matrix-implanted animals when compared to Sham and sTBI only rats.

Despite a net reduction in Olig2+ cells in eCS-implanted animals (Sup. Fig. 6C&D), we observed a significantly higher percentage of Olig2+ cells co-localized with NeuN+ cells in eCS rats when compared to those in the Sham and sTBI groups (Sup. Fig. 6A-C&E). In contrast to these findings, the co-localization of NeuN+ and Olig2+ relative to DAPI+ was found to be significantly reduced in sTBI rats (Sup. Fig. 6B&F) when compared to Sham and eCS matrix-implanted animals.

Taken together, these results suggest that despite a significant reduction in the number of neurons following TBI, eCS matrix implants promoted cell proliferation, neuronal differentiation, synaptic plasticity and potential myelination of newly formed neurons present intra- and perilesionally.

eCS matrix implants attenuate the chronic presence of neuroinflammatory cells

Attenuated influx of neuroinflammatory cells chronically post-sTBI could prevent prolonged tissue damage and atrophy and is a marker of a favorable tissue response post-TBI⁴¹. In order to mark the chronic presence of activated macrophages and reactive astrocytes that are characteristic of a neuroinflammatory cellular response to brain injury, we quantified CD68+ activated macrophages, and glial fibrillary acidic protein labeled (GFAP+) reactive astrocytes (Sup Fig. 7A-C). We observed that eCS matrix-implanted rats had similar levels of CD68+ cells (Sup Fig. 7D) as well as GFAP+ cells (Sup Fig. 7E) as Sham controls, while sTBI rats showed a significant decrease in CD68+ cells compared to Sham and eCS matrix implanted rats. Additionally, we found that CD68+ and GFAP+ cells represented a significantly lower proportion of DAPI+ cells (Sup Fig. 7F) in sTBI and eCS groups, compared to Sham controls. Although the sTBI animals demonstrated a significant reduction in activated macrophage response, this is likely due to the significant tissue loss and absence of intralesional tissue in sTBI animals when compared to Sham controls and eCStreated animals. Overall, these results suggest the prevalence of an attenuated neuroinflammatory cellular response in sTBI and eCS animals at 20 weeks post-TBI.

eCS matrices enhance local vascularization and global blood flow

Since inadequate vascularization is often responsible for the failure of implanted biomaterials⁴², we investigated the extent of tissue neovascularization in eCS matrix implanted animals when compared to controls, using Collagen IV (Col-IV+) and rat endothelial cell antigen (Reca+) markers (Fig.4A&B). We also used MRI phase gradient imaging of cerebral blood flow as a measure of vascular function (Fig. 4C-F).

We found a significant increase in Reca-1+ (Sup. Fig. 8D) and Col-IV+ (Sup. Fig. 8E) percentage area expression in the eCS matrix-treated animals compared to both Sham and sTBI controls. Notably, Reca-1 and Col-IV co-localization was detected to be ~60% in Sham and eCS matrix-treated groups, which was significantly enhanced when compared to the sTBI group (Fig. 4B).

Since a significant reduction of cerebral blood flow (CBF) chronically is linked to cognitive dysfunction and poor prognosis in humans with sTBI^{43.47}, and is also associated with the lack of neuronal activity and loss of neuronal volume in humans and rats^{48,49}, we assessed CBF using MRI-based normalized phase gradient mapping of identified blood vessels in the TBI lesion (nPG; Fig. 4C). We observed a significantly enhanced overall cerebral blood flow (CBF) in eCS matrix-implanted rats (Fig. 4D and Sup. Fig. 9; estimation statistics) when compared to Sham animals, while sTBI control animals showed a nonsignificant decrease in measured CBF. Using ipsi- and contralesional region of interest measurements (Fig. 4E), we detected a specific increase in cortical CBF in eCS matrix implanted rats with a significantly higher CBF detected in the contra- vs ipsilesional side (Fig. 4F). Taken together, these results indicate that eCS matrix implanted rats demonstrate chronically enhanced neovascularization perilesionally, along with significantly enhanced CBF both contra- and ipsilesionally when compared to TBI only controls.

eCS matrices promote chronic forelimb-specific functional recovery and activation of activity-regulated cytoskeleton-associated protein (Arc) in RFA

We used a skilled reach task in order to investigate the circuit-specific implications of the neuroprotective, neurogenic, and angiogenic attributes of eCS matrix implants following sTBI. The skilled reach task was used as a forelimb-specific assessment of motor recovery followed immediately by terminal volumetric imaging of activity-regulated cytoskeleton-associated protein (Arc+). This made it possible to assess task-specific responses following a lesion of the M2 – reach-to-grasp region in the rostral forelimb area (RFA). We found that eCS matrix-implanted rats that trained for 2 weeks on the skilled reach task (Fig. 5A), demonstrated reach-to-grasp performance recovery from week 2 that was comparable to Sham animals, and

that persisted up to week 8 (Fig. 5B and Sup Fig. 10). sTBI control rats showed significant forelimb functional deficits throughout the 8 weeks of testing, with a transient improvement that lasted about 3 weeks (week 4 through 6) before significantly worsening at week 8. The quantification of ipsi- and contralesional Arc+ neurons (Arc+/NeuN+ co-expression; Fig. 5C) immediately post reach and grasp activity, showed strong activation within anterior and posterior RFA regions and low contralateral signal in Sham animals (Fig. 5D). sTBI control rats, in comparison, showed low to no ipsilesional RFA activity with notable contralesional activation when compared to eCS matrix-implanted rats, which demonstrated Arc+ signal within the RFA lesion in addition to contralesional activation.

Perilesional laminar recordings of multi-unit activity reveal the preservation of sensorimotor responses 10 weeks post-TBI in eCS matrix implanted rats

We stimulated the contralesional paw using low intensity electrical pulses to further investigate whether eCS matrix implants directly facilitated recovery of sensorimotor responses (Fig. 5E; Sup. Fig. 11A). We evaluated the spontaneous and evoked (Sup. Fig. 11B and C) electrophysiological responses from the intact M1 and M1/S1 regions using a 32-channel laminar electrode (Sup. Fig. 11D). We observed that following sTBI, a rapid response to electrical stimulation of the left paw reduced in occurrence rate and increased in time jittering in all cortical layers and in the two recording positions CFA and CFA/S1 (Sup. Fig. 11E). Notably, the late response recorded in the CFA region in response to paw stimulation showed an increased delay in sTBI animals (Sup. Fig. 11E left; mean jitter: 36.5 msec), which was found to be faster in eCS matrix implanted rats (mean jitter: 15 msec). This effect was not detected in CFA/S1 region (Sup. Fig. 11E right). We then looked at the region activation pattern in stimulated/recorded rats using Arc+ labeling as in Fig. 5D (Fig. 5F). We found that the level of activity recorded in CFA/S1 returned to Sham levels in eCS rats, while sTBI control rats showed a lower activity level following contralateral paw stimulation. The sTBI control rats also showed a sustained post-stimulation activity registered during non-stimulation recording periods when compared to the pre-stimulation epochs (Sup. Fig. 12), which was not observed in Sham and eCS matrix-implanted rats. Consistently with forelimb-specific circuit activation, we found that

paw stimulation induced activation of the CFA/S1 area and associated motor area CFA with TBI and eCS matrix implanted rats showing a marked reduction and increased activation, respectively. These results indicate that the response and activation of the perilesional circuitry associated with sensorimotor function of forelimb was reduced in TBI rats and partially recovered in eCS matrix-implanted rats.

RFA-specific re-vascularization explains neuronal presence and behavioral performance

We performed brain tissue clearing to specifically investigate whether vascular architecture could reveal a stronger correspondence between vasculature features (Fig. 6), neuronal presence, and behavioral performance.

Following Imaris-based vasculature tracing (Fig. 6A and B), we observed that the anterior RFA region in eCS rats showed an increased vessel segment density (Fig. 6C) and cumulative vessel length (Fig. 6D) when compared to TBI control rats, whereas both TBI and eCS matrix-implanted animals demonstrated a marked reduction in these features compared to Sham. Notably, only 3 out of 5 rats showed tissue presence in the RFAa ROI in TBI controls, whereas all animals in Sham and eCS matrix-implanted groups showed cellular and tissue presence. Tissue clearing and volumetric imaging revealed a larger propensity of tissue preservation in eCS matrix-implanted rats compared to sTBI controls (Sup. Fig. 13), which is consistent with lesion volume analysis obtained in Fig. 1. We also observed that the posterior RFA showed a decreased density of vasculature in both sTBI and eCS matrix-implanted animals.

Interestingly, we found that the mean vessel diameters of vasculature in sTBI controls and eCS matriximplanted animals were not significantly different from those in Sham rats (Fig. 6E). However, consistently with the increased CBF observed in eCS matrix implanted animals (Fig. 4 - 20 weeks), the distribution of both vessel length density and diameter in RFAa were enhanced in eCS matrix-implanted rats when compared to Sham and TBI controls, as shown in Q-Q plots (Sup. Fig. 14A and B).

Importantly, we found a significant correlation between NeuN+ cells and vessel density (Sup. Fig. 15A), and NeuN+ cells and forelimb performance (Sup. Fig. 15B). The correlation between vessel density and

skilled-reach task performance also returns a strong Pearson's R-value (Sup. Fig. 15C), although not significant. Taken together, these results indicate that eCS matrix implants promoted vascularization intralesionally (RFAa), which strongly correlated with neuronal presence and forelimb performance in rats.

DISCUSSION

Parenchymal volume loss of brain tissue is highly correlated with the level of TBI severity in humans, with patients sustaining severe lesions experiencing significantly greater volume loss when compared to those who sustained mild TBIs⁵⁰. As a result, functional losses are inevitable and neuroplastic changes are limited, leaving patients with chronic and debilitating impairments¹. Studies on the intracortical implantation of brain-mimetic CS and hyaluronic acid GAG scaffolds after TBI and stroke have demonstrated the ability to mediate complex structural and functional repair of brain tissue^{37,51,52}. The acute implantation of acellular eCS scaffolds that are functionally diverse and compositionally similar to brain ECM, could present a rational approach to mitigating the significant volume and functional losses encountered chronically after sTBI.

Using a materials design strategy that exploits the native functionality of CS, we demonstrate that neurotrophic factor-laden eCS matrix implants were neuroprotective, neurogenic, and significantly enhanced the peri- and intralesional presence of newly formed blood vessels chronically after sTBI when compared to controls. The role of a global reduction of CBF and hypoperfusion in chronic dysfunction and poor prognosis post-sTBI is well-established⁴⁴⁻⁴⁸. Our observations correlating enhanced vascular density to neuronal presence and reach-to-grasp function recovery, suggests that eCS matrix implants orchestrate complex vascular repair that directly contributes to neuronal function and task-specific recovery chronically post-sTBI. These effects are likely mediated by the enhanced presence and signaling of FGF-2 and BDNF, which is potentiated by CS binding of these factors as demonstrated previously^{17,51}. We nonetheless interpret the implications of observed rCBF increase cautiously in the absence of comparative real-time cerebral oxygenation studies.

The compensatory reorganization of the motor cortex has been observed in regions spanning the perilesional circuitry and the contralateral hemisphere post-TBI⁵³⁻⁵⁶. In our studies, the volumetric quantification of Arc+ activated neuronal populations following left-limb usage confirmed enhanced activity in both the perilesional and contralateral regions in sTBI-only rats compared to Sham controls. eCS

matrix-implanted rats showed a pattern of functional activity similar to sTBI-only rats, with the added activity of intralesional neurons within the implant. Our results also showed a close correspondence between Arc-dependent activation patterns, and both electrophysiological response and recovery of forelimb-specific sensorimotor functions. This evidence suggests that intracortical eCS implants mediated the reorganization of neuronal circuitry leading to chronic recovery of reach-to-grasp function post-sTBI.

In summary, our results demonstrate that rationally designed, brain-mimetic, acellular eCS implants possess native structure-function attributes required to mediate chronic tissue level repair and functional recovery post-sTBI. This study opens up new avenues for the design and application of synthetic sulfated GAG constructs for brain tissue repair, and informs future investigations of neuronal connectivity and electrophysiological responsiveness between the eCS matrix implanted tissue and perilesional cortical columns.

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Author Contributions

C.-F.L, M.B, and LK designed experiments.

C.-F. L., M.B, H.M., R.F. and M.S. performed all animal behavior and surgeries.

N.B. and L.K. performed LCM and qRT-PCR experiments.

M.B performed MRI imaging and G.S. performed phase gradient analysis.

C.-F.L., M.S. and A.A performed the tissue clearing, volumetric imaging and volumetric quantification.

C.-F.L., M.S., R. F., N.B and C.E.L. performed all samples cryosectioning, staining and imaging.

C.-F.L., R.M. and N.B performed IHC-based quantification, behavior analysis, qRT-PCR analysis and statistical analysis.

C.-F. L. and L.K. wrote the manuscript and C.-F.L., Q. Z., P. H., R.V.B and L.K. edited the manuscript.

DATA AVAILABILITY STATEMENT

Data presented and analyzed in this manuscript are available upon request from lohitash@uga.edu

MATERIALS AND METHODS

A – Animal procedures and preparation of eCS matrix

1. Animals

Sprague-Dawley rats were obtained for the 20 weeks (n = 33, Harlan Laboratories, 7 weeks old; males only) and the 10 weeks experiments (n = 21, Charles River Laboratories, 3 weeks old; males only), and randomly assigned to each treatment: control craniotomy with no injury (SHAM group; 20 weeks: n = 9; 10 weeks: n = 6), Controlled Cortical Impact (CCI; TBI group; 20 weeks: n = 9; 10 weeks: n = 6), and CCI implanted with eCS scaffolds (eCS group; 20 weeks: n = 15; 10 weeks: n = 9). A custom-designed controlled cortical impactor delivered the desired impact to the M1 motor cortex (20 weeks: Caudal Forelimb Area, CFA; AP: 0.5 mm; ML: 0.5 mm, relatively to bregma; 10 weeks: Rostral Forelimb Area, RFA; AP: 3.0 mm; ML: 2.5 mm, relatively to bregma) of the eCS matrix implanted animals and TBI-only controls.

All rats were single housed with *ad libitum* access to food and water and following a reverse 12h-light cycle (light OFF 7:00-19:00, Light ON 19:00-7:00) in a room maintained at 70% humidity and 23°C. All procedures on animals were approved by the University of Georgia Institutional Animal Care and Use Committee (IACUC), and protocols were performed in accordance with the Guide for the Care and Use of Laboratory Animals published by the National Institution of Health (NIH).

2. Surgical procedures

Prior to CCI injury, each animal was anesthetized with 5% isoflurane and buprenorphine was injected subcutaneously (0.3mg/ml, 0.05 mL/300g; Henry Schein). Animals were placed on a stereotaxic frame attached to a temperature controlled heating pad (37°C) with their scalp shaved and sanitized (70% EtOH and 3% Povidone-Iodine). Following, a sagittal incision, the periosteum was cleaned using Etch-gel (Phosphoric acid Etching; Henry Schein), and a craniotomy was performed using a 5-mm-diameter trephine bur fitted to an electronic drill. A 3-mm CCI tip was fitted onto the pneumatic piston and positioned in

contact with the surface of the dura (fully extended position), and then retracted to adjust for an impact depth of 2mm. A severe CCI injury was induced by programming the piston speed to 2.25 m/s and a dwell time of 250 ms, resulting in an initial 3-mm-diameter injury with a depth of 2 mm. Absorbable gelatin (Gel foam ®, Pfizer pharmaceutical) was applied to the injury site, and sterile cotton swabs were used to remove excess blood. The Gel foam was then removed, and the injury site was covered completely with a layer of 1% sterile SeaKem (Lonza) agarose. Skin flaps were sutured together, closing the wound. Triple antibiotic cream was applied on the sutured skin.

3. Preparation and intralesional delivery of eCS matrix

Injured animals were randomly selected to receive intralesional injections of eCS matrix functionalized with the neurotrophic factors fibroblast growth factor (FGF-2; $50\mu g/mL$; R&D Systems; 233FB025CF) and brain-derived neurotrophic factor (BDNF; $50\mu g/mL$; R&D Systems; 248BDB010CF), 48h post-sTBI. The eCS matrices were prepared using previously described methods ^{17,36} (22 uL: 8 μ L of 10% methacrylated-chondroitin-sulfate-A-glycosaminoglycan; 7.58 μ L neurobasal media; 4 μ L of 1% photocrosslinker Irgacure-2959, Sigma-Aldrich; 1.21 μ L of FGF-2; 1.21 μ L of BDNF), loaded in a 50 μ L Hamilton Syringe (fitted blunted 21 Gage tight lock stainless steel needle) and photo-crosslinked for 1min just prior to injection. Injection was performed following reopening of the scalp following the above described procedure, removing the 1% agarose and positioning the syringe (angle of 32°, depth: 2mm; speed: 2uL per min for 10 min). Buprenorphine was administered (0.3mg/ml, 0.05 mL/300g; Henry Schein) and the skin was sutured back.

B - Behavioral assessment of functional recovery

Chronic functional recovery was assessed using longitudinal gross motor recovery assays including the beam-walk test (BW)⁵⁷, rotarod test (RTT)⁵⁸ and open field test (OFT) for time points corresponding to week 1, 2, 4, 6, 8, 10, 12, 14, 16, 18, and 20 post-TBI. Chronic recovery of forelimb-specific function was assessed using the skilled reach task (SRT) following preference and forced-side training at 1, 2, 4, 6, and

8 weeks post-TBI. All tasks were performed during the dark cycle in a red-light-lit room held at constant 70% humidity and 23°C.

D - Magnetic Resonance Imaging (MRI)

1. MR imaging procedure

A 7T Varian Magnex MRI scanner and a 35mm-dualtuned surface coil were used to perform MRI of the brain in order to investigate the chronic (20-weeks) effects of TBI. All animals were placed directly under isoflurane (1.5%) anesthesia on the holder of the animal-tube assembly where a catheter was added to the tail of each rat intravenously with a line prefilled with 0.2ml heparin flush attached. The animal's head was securely positioned directly under the 20mm-dualtuned surface coil, and the animal-tube assembly was placed inside the 210 mm horizontal bore of the small animal scanner. For each imaging sequence, eight coronal brain slices with 1 mm thickness and zero gap between slices were acquired using a field of 40x40 mm and a matrix size of 256x256. The following four imaging sequences were acquired for each animal: (1) T₂-weighted fast spin echo (T2WI): TR = 2.0 s, TE = 32 ms, and 4 averages requiring an acquisition time of 4 min and 20 sec and (2) T₂*-weighted multi-gradient-echo (MGEMS): TR=600ms, 8 echoes (TE=10-45ms in 5ms increments), FA=25°, and 4 averages requiring an acquisition time of 10 min and 14 sec. Therefore, the total acquisition time for each animal was ~15 min.

2. Injury volume determination

T₂WI was processed using ImageJ (National Institute of Health, MD), and using the difference in mean gray area in the ipsilesional side compared to the contralesional side. As T2W images revealed the largest areas of injury and inflammation, T2W images were used to design a region of interest (ROI) library for each animal's injured cortex.

3. Phase-gradient mapping

The rate of flow in a blood vessel is inversely proportional to the change in phase between the vessel and

the surrounding tissue ⁵⁹. However, during MR acquisition, phase measurements are only recorded in the range of $\pm \pi$, and any values that extend outside of this range get "wrapped", or forced into this range, making calculating the change in phase difficult in areas where wrapping occurs. Since the change in phase is proportional to the change in the phase gradient, phase gradient maps (PGMs) can be used to avoid this wrapping problem ⁶⁰.

Changes in rCBF between groups was determined using the T2*-weighted MGEMS data using the following procedure ⁶¹. Phase gradient maps (PGMs) were generated for each slice by calculating the magnitude of the phase gradients determined in the readout and phase encoding directions ⁶⁰. From the PGMs, phase gradient differences between a vessel and the surrounding tissue were measured across all distinguishable vessels located within the cortex, sub-cortical and basal forebrain areas, and thalamus. The third echo (TE=20ms) was used for all measurements as this echo produced the best contrast for distinguishing vessels within the PGMs. To account for systematic phase variations in the MRI acquisition across rats (differences in coil tuning, gradient shimming, etc.), measurements were normalized using the phase gradient differences were calculated for all vessels located on both the contralateral and ipsilateral side of the brain to the injury for each region and group.

E – Brain tissue preparation and immunohistochemistry

1. Tissue collection at 20-week time point

20 weeks post-injury, animals were heavily sedated using ketamine (65mg/kg) and transcardially perfused with 250 mL 0.1 phosphate-buffered saline (PBS, pH 7.4) followed by 250 mL 4% paraformaldehyde solution. Brains were then extracted and cut in half (transversal cut), 7mm from the apex of the frontal cortex, flash frozen in liquid nitrogen and stored at -80°C. Coronal sections were obtained using a Leica Cryostat, starting from the center of the injury and moving caudally. A total of 48, 15-µm-thick, sections were collected per animal and distributed evenly over 10 charged slides and 2 membrane slides (4 sections

per slide). 2 slides per animal were placed in 4% paraformaldehyde containing 4M sucrose, and stained with 0.1% cresyl violet solution to label Nissl bodies in neurons. The remaining 8 charged slides were used for immunofluorescence assays, and the 2 membrane slides were used for qRT-PCR. For immunostaining, slides were dried and placed in 4% paraformaldehyde solution containing 4M sucrose, assigned to one of the primary antibody groups and incubated overnight at 4°C (Table 1). Slides were then incubated with blocking buffer consisting of 1:220 dilutions of appropriate secondary antibodies (Table 2), counterstained with cell nuclear stain, NucBlue ® (Life Technologies, NY), mounted with Fluormount-G (Southern Biotech, AL), sealed with coverslips and stored at -20°C until imaged.

2. Laser capture microdissection (LCM)

LCM was performed on two membrane slides per animal as previously described⁶². In brief, brain tissue sections were placed on a cold block and fixed for 2 minutes in ice-cold 75% ethanol solution then rinsed thrice with ice-cold nuclease free PBS (pH 7.4, Life Technologies). Brain sections were stained in a 1:220 dilution of goat anti-rat IgG + 500 U RNase inhibitor (Life Technologies). After 1-hour exposure, sections were rinsed three times with nuclease free PBS and dehydrated sequentially with 75%, 95%, 100% ethanol for 15 seconds each. Sections were placed on the ice block to dry for 5 minutes prior to LCM using a laser microdissection microscope (Leica).

3. qRT-PCR

Total RNA was extracted from LCM processed tissue using a picopure RNA isolation kit (ThermoFisher). and quantified using a nanodrop analyzer (ThermoFisher). 100 ng equivalent of pooled total RNA (5 slices per animal; 7 rats) was used to synthesize cDNA using an RT First Strand Kit (Qiagen). Pre-validated primers targeting rat *CXCR4*, *CXCL12*, *BDNF*, *FGF-2*, along with the housekeeping genes *HPRT1* and *GAPDH* (Qiagen) were used in qRT-PCR assays for 40 cycles in three steps: 95°C for 10 minutes, 40 cycles of 95 °C for 15 seconds, and 60 °C for 1 minute. Melting curve analysis was conducted to validate amplification specificity. All assays were conducted in triplicate on a 7900 HT instrument (Applied

Biosystems) and gene expression levels normalized to endogenous gene controls plotted as relative units using the $\Delta\Delta$ Ct method. Gene expression was analyzed with an absolute fold difference greater than 2. Gene expression was normalized against the average expression of SHAM controls.

4. Tissue collection and preparation for task and stimulation specific activation of immediate

early genes - 10 weeks

For the visualization of functional activity in the whole brain, we used antibody staining against the immediate early gene (IEG) encoded actin-regulated cytoskeleton-associated protein (Arc). Two groups of rats were used at 10 weeks post-TBI. The first cohort of rats performed a continuous 40 min of SRT (specific left forelimb activation with high repetition; n = 3-4 per group) before being immediately sacrificed for tissue collection. The second cohort of rats were used for intra-cortical recordings and perilesional electrophysiological response to the left paw stimulation (see electrophysiological recordings and paw stimulation section; n = 3-4 per group) before being sacrificed for tissue collection. These two cohorts were used to map the volumetric activation of neuronal networks in relation to either volitional control of the left forelimb (SRT cohort) or the sensorimotor response to (Paw Stim). All brains were collected after transcardial perfusion of paraformaldehyde as described above, and processed for iDisco+ tissue clearing.

5. iDisco+ tissue clearing and immunostaining

Following post-fixation, the frontal sections of the brains were treated for immune staining and tissue clearing using the iDisco+ method⁶³ with steps described briefly in Table 3.

6. Tissue imaging and quantification

Brain slices were imaged using either a Zeiss LSM7100 confocal microscope (Zen® software; Immunostained slices), epifluorescence Leica DM IRBE (Volocity® software; Cresyl/Nissl Stain slices) or Lavision Ultra II lightsheet microscope (iDisco+ cleared brains). Image quantification for immunostained slices was performed using MATLAB and custom scripts based on the image processing toolbox. For cresyl violet stained slides the region of interest (ROI) represented 10.494 mm², and four images were taken per animal per group (n = 7, 28 total images). ImageJ was used to calculate and analyze the total number of marked nissl bodies, thresholding to signal peak (~200) and using the subtraction tool to eliminate any noise (National Institute of Health, MD). For volumetric images, all image stacks were converted and processed using the software Imaris @ (Bitplane). Nuclear counts (NeuN+ and Arc+ Staining) and colocalization were performed using spot detection and spot colocalization. Vascular density (Reca1+), orientation was performed using a batch sequence of 1) surface mapping, 2) surface model-based masking, 3) masked image Gaussian filter and 4) filament tracing (closed-loop) for vasculature parameter extraction.

G – Electrophysiological recordings

1. Perilesional electrophysiological recordings

At week 8, rats that received RFA-targeted lesion were induced and maintained under anesthesia using a Ketamine/Xylazine cocktail (100mg/Kg) and placed on a stereotaxic frame. Following sagittal scalp incision, a craniotomy was performed caudally to the injury (AP: 0 mm, ML: 2 to 4mm, relatively to bregma), followed by durotomy to allow for the insertion of a 32 channel silicone probe (Neuronexus). The silicone probe was then inserted in the motor area (position1: CFA, AP: 0 mmm, ML: 2.5 mm) or sensorimotor area (position2: S1, AP: 0 mmm, ML: 3.5 mm) at a depth of 2 mm from the surface of the brain. All recordings were performed with a single shank, 32 channel linear probe with 50 um spacing between recording sites (177 mm²) and a maximum span of 1.6 mm (A1x32-6mm-50-177-CM32, 15um thickness, Gen4, lot# P994). The probe reference was used as main reference and a screw positioned above the right cerebellum was used as ground.

Neural data was digitalized and recorded at 20 kHz (unit gain) using a Multichannel systems (MCS) W2100 acquisition module and a wireless headstage (HS32-EXT-0.5mA, 16bit). For spike analysis, the broadband

electrophysiological data was real-time bandpass filtered (300-5000 Hz) and a baseline pre-calculated threshold was used as trigger to save spike waveforms and spike event timestamps (threshold: 5 standard deviation). All recordings were performed for 5 min following a 10 min of stabilization.

2. Paw Electrical Stimulation Evoked Neural Activity

Paw electrical stimulation were performed from a separate HS32-EXT-0.5mA headstage to provide isolated stimulation. The stimulating electrode was a stainless-steel needle inserted in the foot pad of the rat paw. Paw stimulations were performed at a fixed intensity (50uA), biphasic electrical pulse (1msec phase 1) optimized to minimize recording artefacts while eliciting triggered neural response. The Paw stimulation pulses were repeated 120 times at a frequency of 1Hz. Each stimulation was monitored using an LED output a synchronized TTL port to the recording system.

I - Statistical analysis

All statistical analyses were performed using SigmaPlot (SyStat Software, Inc., CA) or Matlab statistical toolbox (Mathworks, inc.). For multiple group comparison (e.g. immunohistochemistry quantifications, Volume or area quantification), we used a one-way ANOVA followed by a multiple comparison least significant difference (LSD) due to the low number of group (3 groups, SHAM, TBI, and eCS). For group comparison on the normalized phase gradient analysis (MRI) we used a 3-way ANOVA followed by multiple comparison using the LSD approach. For behavior data, we performed two-way repeated measure ANOVA using between factor group (three level: SHAM, TBI, eCS) and within factor time with a post-hoc multiple comparison based on the LSD. When necessary (failure of normality of sample distribution), non-parametric tests such as Kruskal-Wallis and Mann-Whitney U Ranksum test were used in place of one-way ANOVA and t-tests. For distribution comparisons, the measure of effect size (Matlab toolbox, dabest estimation statistics)⁶⁴ was used to avoid bias due to high sample number as well as quartile-quartile plot against SHAM distribution. For all tests, p<0.05 was considered significant.

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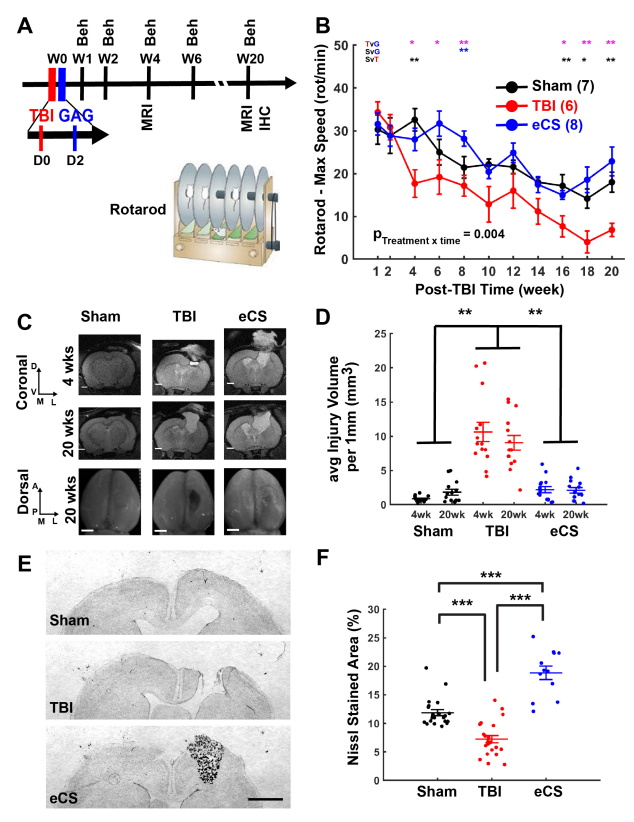


Figure 1: eCS implants reduce lesion volume and improve gross motor function 20 weeks post-TBI

A – Experimental schedule. All rats received a TBI at week 0 (D0). eCS implants were intracortically administered 48hour post-TBI (D2). Following one week of recovery, all rats underwent behavioral testing at week 1 and 2, and every other week therafter (4, 6, 8, 10, 12, 14, 16, 18. 20).

B – The Rotarod (RR) test was used a measure of balance and motor coordination. Two-way repeated measure ANOVA; $p_{Treatment} = 0.153$; $p_{Time} < 0.001$; $p_{TreatmentxTime} = 0.004$. Post-hoc multiple comparison using Dunn-Sidak correction are shown above each time point for Sham vs TBI (red), Sham vs eCS(blue) and TBI vs eCS(black).

C - Representative T2W MRI images (top panel) for each treatment group Sham, TBI and eCS (coronal section), scale bar: 500 μm; top view of the extracted brain (bottom panel), scale bar: 1 mm; D: dorsal, V: Ventral, M: middle, L: lateral, A: anterior and P: Posterior.

D – Average injury volume quantified for each 1mm slice around the injury and based on the T2W MRI images (mm3). Two-way repeated measure ANOVA; Treatment factor: p < 0.001; Time factor: p = 0.53; Treatment x Time: p = 0.0279.

E – Representative Nissl stained coronal sections of rat brain for the Sham, TBI and eCS groups. Scale bar:1 mm.

F – Lesion volume quantification using Nissl stain for Sham (6 rats, 4 images per rat), TBI (6 rats, 4 images per ras) and eCS (3 rats, 4 images per rat) groups. One-way ANOVA; Treatment: p < 0.001. Post-hoc LSD *, ** and *** are for p <0.05, p < 0.01 and p <0.001, respectively. Graphs show mean \pm s.e.m.

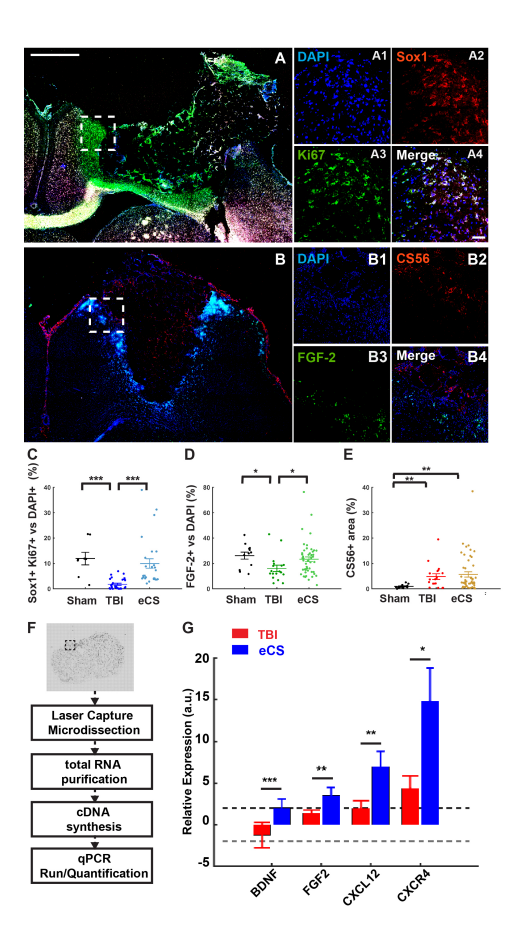


Figure 2: eCS implants promote NSC proliferation and neurotrophic factor expression

A – Representative tiled images of ipsilesional hemisphere (left coronal sections) of eCS rat brain tissue; scale bar = 1mm. A1-A4 – representative magnification of dashed white square shown in C for DAPI (A1), Sox1 (A2), Ki67(A3) and merged (A4); scale bar is 100 μ m;

B - Representative tiled images of ipsilesional hemisphere (left coronal sections) of eCS rat brain tissue;

scale bar = 1mm. B1-B4 - representative magnification of dashed white square shown in A for DAPI (B1),

CS56 (B2), FGF2 (B3) and merged (B4); scale bar is $100 \ \mu m$;

C – Co-localization Sox1+ cells with Ki67+ cells as percentage of Ki67+ cells for each treatment. Kruskal-Wallis, Treatment: p <0.001.

D – Co-localization FGF-2+ cells with DAPI+ cells as percentage of DAPI+ cells for each treatment. Kruskal-Wallis, Treatment: p <0.05.

E - CS56 + percentage area for each treatment. Kruskal-Wallis, Treatment: p < 0.001. Post-hoc LSD Mann-Whitney U test *, **, and *** are for p< 0.05, p< 0.01, and p< 0.001, respectively.

F – At 20-week time point, brains were flash frozen and tissue was laser microdissected. Total RNA was purified and used to synthesize cDNA. qRT-PCR was run using pre-validated primers of interest for later quantification of gene expression.

G – Relative expression of *BDNF*, *FGF2*, *CXCL12* and *CXCR4* transcripts in sTBI and eCS treatment groups. Two-tail t-test *, **, and *** are for p<0.05, p<0.01, and p<0.001, respectively. Graphs show mean \pm s.e.m.

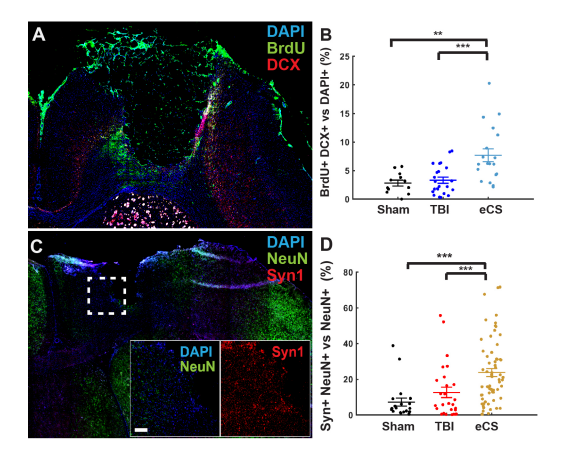


Figure 3: eCS implants promote neurogenesis and plasticity

A – Representative tiled images of ipsilesional hemisphere (left coronal sections) coronal sections with merged DAPI (Blue), DCX (red) and BrdU (green) staining of eCS rat brain tissue; scale bar = 1mm. B –Co-localization of DCX+ and BrdU+ cells with DAPI+ cells as percentage of DAPI+ cells for each treatments; Kruskal-Wallis, Treatment: p < 0.001.

C – Representative tiled images of ipsilesional hemisphere (left coronal sections) with merged DAPI (Blue), Syn1 (red) and NeuN (green) staining of eCS rat brain tissue; scale bar = 1mm. Inset shows representative magnification of dashed white square shown as DAPI and NeuN (merged; left) and Syn1 (right); scale bar is 100 μ m;

D – Co-localization Syn + cells with NeuN + cells as percentage of NeuN + cells for each treatment. Kruskal-Wallis, Treatment: p < 0.001. Post-hoc LSD Mann-Whitney U test *, **, and *** are for p<0.05, p<0.01, and p<0.001, respectively. Graphs show mean \pm s.e.m.

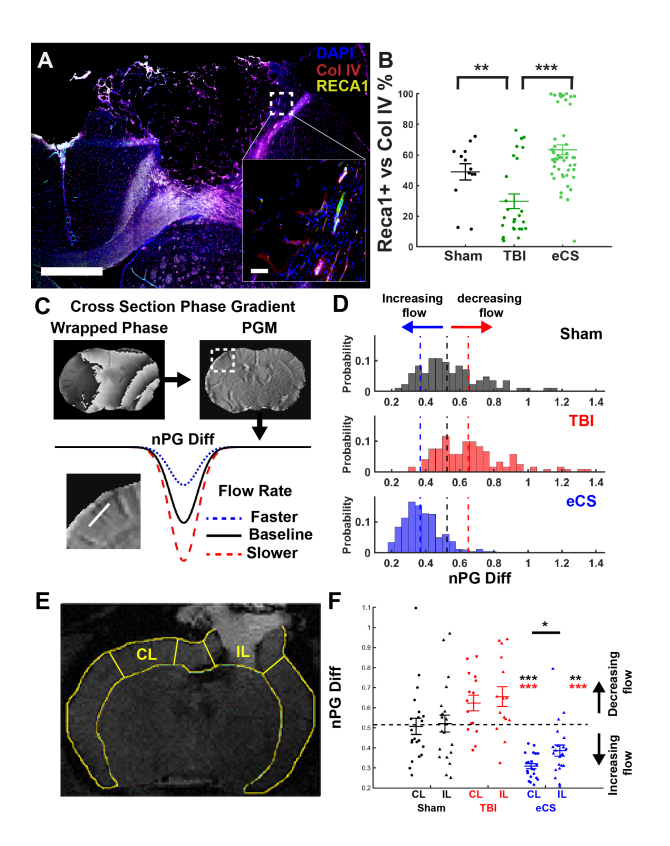


Figure 4: eCS promote vascularization and increased CBF 20 weeks post-TBI

A – Representative tiled image of ipsilesional hemisphere of the eCS group; scale bar = 1mm.

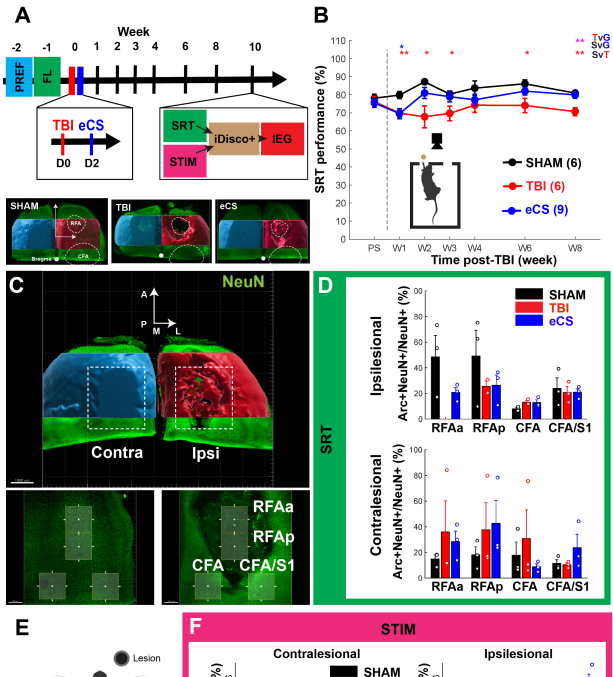
B – Co-localization Reca1 + cells with Col IV + cells as percentage of Col IV + cells for each treatment. Kruskal-Wallis, Treatment: p <0.001. Post-hoc LSD Mann-Whitney U test *, **, and *** are for p<0.05, p<0.01, and p<0.001, respectively.4

C – Phase gradient was estimated from MRI session. First the wrapped phase map (top left panel) was used to generate phase gradient maps (PGMs, top right panel) for each slice by calculating the magnitude of the phase gradients determined in the readout and phase encoding directions (see Material and Methods for details). (bottom panel) Normalized phase gradient (nPG) differences between a vessel and the surrounding tissue were measured across all distinguishable vessels at the coronal cross section (inset, white line). Note: The rate of flow in a blood vessel is inversely proportional to the change in phase between the vessel and the surrounding tissue (see methods).

D – Distribution of nPG for all identified blood vessel in each treatment groups Sham (n = 84), TBI (n = 121) and eCS (n = 432). Comparison using the glass delta measure effect size and 10,000 bootstrapping. Dashed line represent the median nPG for Sham (black), TBI (red) and eCS (blue). One-way ANOVA, F(2) = 230.56, p = 8.57e-51 / measure effect size p < 0.001.

E – Representative segmentation of MRI coronal section in sub-region for regional and hemispheric comparison in blood flow. IL: ipsilesional, CL: contralesional.

F - nPG for the perilesional region of interest and matching region of interest in the contralateral hemisphere. Two-way ANOVA, Treatment: p <0.0001. Post-hoc one-sample t-test; Post-hoc LSD *, **, and *** are for p<0.05, p<0.01, and p<0.001, respectively. Graph shows mean \pm s.e.m.





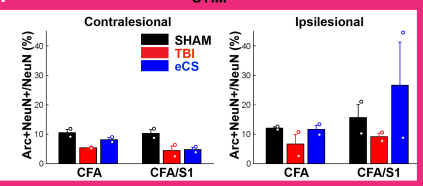


Figure 5: eCS implanted animals demonstrate enhanced recovery of Reach-to-Grasp-specific motor function 8 weeks post-TBI

A – Experimental schedule of skilled reach task. (Top panel) All rats received training for the skilled reach task starting from a preference training followed by a forced left limb training. A CCI at the rostral forelimb area (RFA) was performed followed by eCS implants (48h post-sTBI). Rats were tested weekly/bi-weekly on the skilled reach task. At week 10, Rats were terminally sacrifices and brain processed for immediate early gene expression following either an intensive skilled reach task assay or contralesional paw stimulation/brain recordings. (Bottom panel) Representative volumetric segmentation of lesion from iDisco+ cleared and NeuN+ stained brains. Representative images showing tissue presence for Sham (left), TBI (middle) and eCS (right) groups using surface mapping in Imaris. The contralesional and ipsilesional hemispheres are color coded in blue and red, respectively.

B – Overall performance for the skilled reach task in percentage of the Sham (n = 6), TBI (n = 6) and eCS (n = 9) groups post-TBI and treatment. The SRT performance score is calculated from the average between success hit (%), left-limb usage (%) and efficiency (%). Repeated measure ANOVA, Time: p <0.05, Group: p < 0.01, Time x Group: p = 0.1306; Post-hoc LSD * and ** indicate a p<0.05 and p<0.01, respectively. The color of stars indicates statistical test between the following pairs: Sham vs TBI (SvT, red), Sham vs eCS (SvG, Blue) and TBI vs eCS (TvG, magenta). RFAa: RFA anterior; RFAp: RFA posterior. SRT: Skilled reach task.

C – Region of Interest (ROI) definition for the analysis of Arc+ NeuN+ colocalization for reach-to-grasp function mapping. The ipsilesional and contrlaesional sides are marked in red and blue, respectively. All rats were sacrificed at week 10 post-sTBI, 30 min post SRT assay, transcardially perfused, and brain tissue processed for iDisco+ immunostaining. Imaris quantification of lightsheet images was used to determine the percentage of Arc+ expressing NeuN+ cells within each ROI. The contralesional and ipsilesional hemispheres are color coded in blue and red, respectively.

D – Ipsilesional (top panel) and contralesional (bottom panel) quantification of Arc+ NeuN+ colocalized cells post-SRT assay reveals RFA specific activation of forelimb circuitry in Sham as well as in eCS rats. TBI rats showed absence of cell or activity in the RFAa. Both TBI and eCS rats showed increased perilesional and contralesional activity 10 weeks post-TBI. Sham (n = 3), TBI (n = 3) and eCS (n=3). Line plots and bar graphs show mean \pm s.e.m.

E-Experimental setup for the paw stimulation combined with laminar electrophysiological recording. Rats were sacrificed after stimulation-recording to quantify neuronal activity-induced Arc expression in somatosensory regions.

F – Contralesional (left panel) and ipsilesional (right panel) quantification of Arc+ NeuN+ colocalized cells post-STIM assay for Sham (n = 2), TBI (n = 2) and eCS (n = 2). Bar graph show mean+/-s.e.m. Line plots and bar graphs show mean \pm s.e.m.

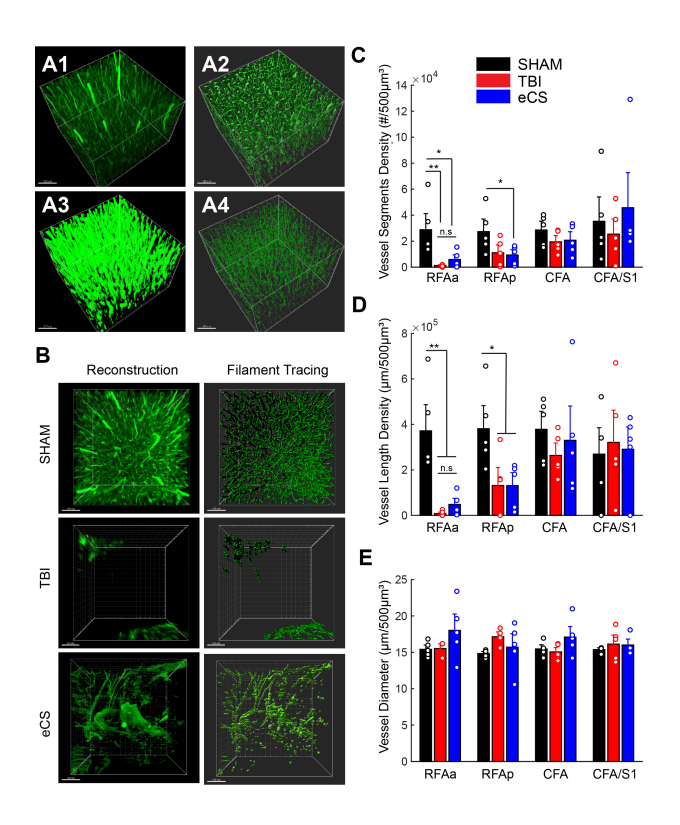


Figure 6: eCS implants promote changes in intra and perilesional vascular architecture 10 weeks post-TBI

A – Image processing step for vasculature tracing. Using Imaris software on volumetric imaging of iDisco+ cleared brains, we first used reconstructed volumetric images from Reca1+ staining (green - A1), performed surface mapping (A2), surface-based masking and Gaussian filtering (A3) and finally filament tracing (A4). Representative vascularization in eCS implanted region; original Reca1+ images (left) and vasculature traced ROIs (right)

B – Representative images of Sham (Top panel), TBI (middle panel) and eCS (bottom panel) rat brain tissue for the reconstructed original volume image (left) and vasculature tracing (right).

C – Quantification of blood segment vessel density per 500 μ m³ for Sham (n = 15), TBI (n = 15), eCS (n = 15) groups in all four ROIs as in figure 6.

D – Quantification of blood vessel length density per μ m/500 μ m³ for Sham (n = 15), TBI (n = 15), eCS (n = 15) groups in all four ROIs as in figure 6.

E – Quantification of mean blood vessel diameter per 500 μ m³ for Sham (n = 15), TBI (n = 15), eCS (n = 15) groups in all four ROIs as in figure 6. Post-hoc LSD Mann-Whitney U test *, ** and *** indicate p < 0.05, p < 0.01 and p < 0.001, respectively. Bar graphs show mean ± s.e.m. RFAa: RFA anterior; RFAp: RFA posterior/perilesional; Graphs show mean ± s.e.m.

TABLES

Table 1: Primary Antibody panel for immunostaining

Purpose	Antibody	Manufacturer	Cat#	Host	Dilution	Usage
Neuronal	NeuN	EMD	MAB377	Mouse	1:500	Slices
Presence/marker		Millipore				
Neuronal	NeuN	EMD	ABN91	Chicken	1:500	iDisco+
Presence/marker		Millipore				
Neuronal	Doublecortin	Abcam	ab18723	Rabbit	1:500	Slices
Proliferation	(DCX)					
Myelination	Olig2	Abcam	ab109186	Rabbit	1:100	Slices
Neural	Sox-1	Abcam	ab87775	Rabbit	1:1000	Slices
Progenitor	Ki67	Santa Cruz	sc-7846	Goat	1:500	Slices
Neuroinflamation	GFAP	DAKO	z0334	Rabbit	1:1000	Slices
	CD68	Bio-Rad	mca341R	Mouse	1:500	Slices
Growth Factor	FGF	Abcam	ab8880	Rabbit	1:200	Slices
presence						
Chondroitin	CS56	Sigma-	C8035	Mouse	1:200	Slices
Sulfate GAG		Aldrich				
presence						
Plasticity	Synapsin-1	Abcam	ab14692	Rabbit	1:500	Slices
Neuronal	Arc	Synaptic	156 003	Rabbit	1:1000	Slices
Activation		Systems				/iDisco+
Marker						

Vasculature	RECA	BioRad	MCA970R	Mouse	1:500	Slices
						/iDisco+
	Col IV	Abcam	Ab19808	Rabbit	1:500	Slices

Table 2: Secondary Antibody panel

Alexa-Fluor	Host	Reactivity	Isotype	Catalog #	Company
488	Goat	Mouse	IgG (H+L)	A11006	Life Technology
488	Goat	Chicken	IgY (H+L)	A11039	Life Technology
647	Goat	Rabbit	IgG (H+L)	A21244	Life Technology
488	Donkey	Goat	IgG (H+L)	A11055	Life Technology
594	Goat	Mouse	IgG (H+L)	A11005	Life Technology

Table 3: iDisco+ tissue clearing steps. For primary and secondary antibody concentration please refer to table 1 and 2. RT: room temperature. DCM: dichloromethane. DBE: Dibenzyl ether. PBS: phosphate buffered saline. PTx2: 0.05% Triton-X100 in 1x PBS. PTwh: 0.0.5% Tween-20, 0.5mL of 10mg/mL of Heparin in 1x PBS.

Process	Condition	Reagent	Solvent
Initial wash	15min x3 in each	Ptx2 and PTwh	1x PBS
Serial dehydration	1h each	20/40/60/80/100%	1x PBS
		methanol	
Defatting	Overnight	DCM (66% v/v)	Methanol
Bleaching	Overnight	H2O2 (5% v/v)	Methanol
Serial rehydration	1h each	20/40/60/80/100%	Methanol
		PBS	

permeabilization	4 days	Triton-X100, 20%	1x PBS
		DMSO	
Blocking	2 days	Goat serum (6%v/v)	Ptx2
		and DMSO $(10\% \text{ v/v})$	
Primary antibody	7 days, 37deg C, 100	Goat serum (3%v/v)	Ptx2
incubation	RPM	and DMSO $(5\% v/v)$	
wash	overnight		PTwh
Secondary antibody	6 days, 37deg C, 100	Goat serum (3%v/v)	Ptx2
incubation	RPM	and DMSO $(5\% v/v)$	
Final dehydration	1h each	20/40/60/80/100%	DiH2O
		methanol	
defatting	3h, RT	DCM (66% v/v)	Methanol
defatting	Overnight	DCM 100%	DCM
Final	30min – hold	100% DBE	DBE
defatting/clearing	indefinitely		