Tissue Engineered Neural Constructs Composed of Neural Precursor Cells, Recombinant Spidroin and PRP for Neural Tissue Regeneration

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We have designed a novel two-component matrix (SPRPix) for the encapsulation of directly reprogrammed human neural precursor cells (drNPC). The matrix is comprised of 1) a solid anisotropic complex scaffold prepared by electrospinning a mixture of recombinant analogues of the spider dragline silk proteins – spidroin 1 (rS1/9) and spidroin 2 (rS2/12) - and polycaprolactone (PCL) (rSS-PCL), and 2) a “liquid matrix” based on platelet-rich plasma (PRP). The combination of PRP and spidroin promoted drNPC proliferation with the formation of neural tissue organoids and dramatically activated neurogenesis. Differentiation of drNPCs generated large numbers of βIII-tubulin and MAP2 positive neurons as well as some GFAP-positive astrocytes, which likely had a neuronal supporting function. Interestingly the SPRPix microfibrils appeared to provide strong guidance cues as the differentiating neurons oriented their processes parallel to them. Implantation of the SPRPix matrix containing human drNPC into the brain and spinal cord of two healthy Rhesus macaque monkeys showed good biocompatibility: no astroglial and microglial reaction was present around the implanted construct. Importantly, the human drNPCs survived for the 3 month study period and differentiated into MAP2 positive neurons. Tissue engineered constructs based on SPRPix exhibits important attributes that warrant further examination in spinal cord injury treatment.

Spinal cord injury (SCI) is one of the most difficult-to-treat conditions with an incidence ranging from 250 to 906 cases per million per year worldwide1. In spite of a large arsenal of neuro-rehabilitation tools and procedures that include reconstructive spinal surgery and mechanical rehabilitation, kinesiotherapy as well as magnetic and electric stimulation, the degree of spinal cord function restoration in severe spinal cord trauma (American Spinal Injury Association, ASIA A or B level) is still catastrophically low2,3.

One of the most promising experimental approaches for spinal cord regeneration is transplantation of neural stem cells or neural precursor cells (NPCs)4–7. The development of safe techniques to generate autologous NPCs, first suggested by the use of iPS technology8,9, and later by direct reprogramming of somatic cells10–13, opened up new therapeutic opportunities in the regenerative field. In particular, the use of autologous NPCs prepared from the patients’ bone marrow with the help of a cocktail of non-integrating transcription factors and small molecules

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rather uniformly in the differentiating neural organoids, while III-tubulin and especially MAP2-positive young MAP2-positive neuronal progenitors (Fig. 2A). Nestin- and GFAP-positive glial progenitors were distributed differentiating neural organoid-like structures, interconnected by processes of differentiating β

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activation of neural cell adhesion molecule (NCAM) expression and promotes neural differentiation due to

expression of multiple repeats of a neuron-specific GRGGL sequence recognized by neural progenitors25. Recombinant analogues of natural spidroin 1 and spidroin 2 (rS1/9 and rS2/12 proteins, respectively) are charac-

terized by high biocompatibility, absence of immunogenicity, low biodegradability and ability to promote animal tissue regeneration26,27.

Another scaffold component of interest in regeneration is platelet-rich plasma (PRP). A number of studies have shown that PRP promotes neurogenesis and axonal growth28, as well as enhancement of proliferation and migration of Schwann cells29 due to the presence of different growth factors (PDGF-AB, TGF-

NGF and GDNF) and platelet-released exosomes containing microRNA and other signaling molecules. These properties make PRP a prospective source of autologous growth factors providing the necessary biomimetic microenvironment for neuroregeneration40.

The aims of this study were to (1) develop a two-component matrix SPRPix, based on PRP and an anisotropic complex scaffold of recombinant spidroins and polycaprolactone (rSS-PCL) for the creation of 3D tissue engineered constructs with drNPC (2) to test the biocompatibility of the drNPC-SPRPix construct in the brain and spinal cord of the Rhesus macaque and 3) to examine the survival and behaviour of the implanted human drNPC cells.

Results

Immunophenotyping of drNPC. We initiated our study by characterizing the drNPCs obtained from direct reprogramming of human bone marrow cells13,15. Using flow cytometry and immunochemical analysis, we demonstrated high level expression of the neural stem cell marker SOX2 and neural markers βIII-tubulin and MAP2 in the drNPC culture (Fig. 1A). drNPC cultured on laminin-coated adhesive plastic in complete medium exhibited a characteristic "honeycomb" pattern and co-expressed βIII-tubulin and GFAP (Fig. 1B,C), SOX2 and Nestin (Fig. 1D). Upon switching to differentiation conditions, MAP2- and NF200-positive neurons emerged (Fig. 1E,F), as well as GFAP-positive astrocytes (Fig. 1E). Differentiated cultures did not express SOX2 (Fig. 1F), indicating complete differentiation. Thus, the flow cytometry and immunochemical analysis confirmed that drNPC cells have a human neural stem cell-like phenotype and were capable of differentiation into both neuronal and glial lineages.

Characterization of drNPC cultures in a PRP-based "liquid matrix". Having characterized the drNPC during active expansion and upon differentiation in adhesive plastic, we then characterized the behav-

ior of these cells upon culture in the new PRP-based "liquid matrix”; after 2–3 weeks of culture we observed an almost complete replacement of the PRP hydrogel by a three-dimensional neural network that consisted of differentiating neural organoid-like structures, interconnected by processes of differentiating βIII-tubulin and MAP2-positive neuronal progenitors (Fig. 2A). Nestin- and GFAP-positive glial progenitors were distributed rather uniformly in the differentiating neural organoids, while βIII-tubulin and especially MAP2-positive young neurons had a more central localization (Fig. 2B,C). The long processes of βIII-tubulin positive neurons were surrounded by nestin-positive progenitors (Fig. 2D). In conclusion, the PRP-based liquid matrix preserves the biological attributes of the cells, supporting proliferation and spontaneous differentiation into neuronal and glial lineages.

drNPC culturing on low-adhesive plastic coated with recombinant spidroin rS1/9. With the objective of testing the compatibility of drNPCs with spidroins, we then seeded drNPCs on a plastic dish coated with rS1/9 which contains the neural cell adhesion molecule (NCAM) specific amino acid sequence25. The cells rapidly adhered not only to the plastic but also to a glass coverslip in the center. Despite the use of complete medium, drNPCs plated on rS1/9 coated plastic exhibited intense spontaneous neuronal differentiation with the formation of βIII-tubulin positive neuroblasts and immature neurons (Suppl. Figs 1 and 2). This was accom-

panied by a high expression of BDNF as shown by immunochemical analysis (Suppl. Fig. 2). These data demonstrated that spidroin rS1/9 promoted not only drNPC adhesion to culture surfaces but also active neuronal differentiation, which we could partially replicate in laminin-coated plastic but only in the presence of differentia-
tion medium.

Construction and testing of various scaffolds. Having determined that spidroin rS1/9 had no neg-

ative impact on the biology of drNPCs, we then started working on constructing a spidroin-based scaffold. In
Figure 1. Immunophenotyping of drNPC. (A) Cytofluorometry of drNPC marker expression. 1. Negative control (isotypic immunoglobulins) 2. βIII-tubulin 3. MAP2 and 4. SOX2. (B–F) Immunocytochemical analysis for different markers (B) anti-βIII-tubulin. Bar = 100 µm (C,D) Undifferentiated drNPC coexpress βIII-tubulin (green) and GFAP (red) (C), as well as Nestin (green) and SOX2 (red) (D). (E,F) Differentiation of drNPC: MAP2-positive neurons (green) and GFAP-positive astrocytes (red) (E) NF200-positive (red) and SOX2 negative neurons (F). In panels (B–F), the blue channel corresponds to the Hoechst stained cell nuclei. In panels C–F Bar = 50 µm. Laser scanning confocal microscopy using Nikon A1 device.

parallel we also tested a commercially available collagen scaffold «Chondro-Gide»® (Geistlich Pharma AG, Switzerland) widely used for cartilage tissue regeneration, as a model system to develop the technique for creation of a two-component matrix based on PRP and a solid scaffold.
We tested a complex scaffold prepared by electrospinning a mixture of recombinant spidroins and polycaprolactone (PCL, a biocompatible and biodegradable compound with a MW ~ 400 kDa) – we called this scaffold rSS-PCL. Spiders are known to use a mixture of spidroins 1 and 2 to boost the resilience of the dragline silk of a web, the first of which confers enhanced strength and the second one, due to the presence of a GPGXX (X = G,Y,Q) motif, confers enhanced elasticity. In order to benefit from both properties, we utilized a mixture of rS1/9 and rS2/12 proteins. PCL was added to the mixture to improve its spinnability. We experimentally determined that a ratio of 6:1:1 (rS1/9:rS2/12:PCL) resulted in the optimal combination of scaffold stiffness and wettability which resulted in improved proliferation and neuronal differentiation of drNPC as shown in the corresponding results section.

Figure 2. drNPC cultured in the PRP based liquid matrix. (A) 3D reconstruction of cultured drNPCs-01 in PRP based hydrogel using laser scanning confocal microscopy. (B) Spontaneous differentiation of neural organoids formed in PRP based hydrogel with the appearance of βIII-tubulin neuronal progenitors (red) amongst the more common Nestin positive (green) NPCs (C) Neural organoids stained with MAP2 (green) and GFAP (red) further confirming glial and neuronal differentiation. (D) Low magnification image showing the overall morphology of the differentiating cultures with large numbers of Nestin positive (green) NPCs and βIII-tubulin positive neurons (red). The optical plane cuts through the center of the neural organoids. (B–D) bar = 100µm.
Physico-chemical characterization of the rSS-PCL scaffold. Scanning electron microscopy (SEM) demonstrated that the rSS-PCL scaffold was characterized by an anisotropic structure with a predominantly unidirectional orientation of parallel microfibrils; the thickness of each microfibril was about 1 µm (Fig. 3A,B). More detailed imaging with Atomic Force Microscopy (AFM) (Figs 3C and 5D) revealed that the microfibrils were not smooth, and their surface had a considerable roughness with a nanodimensional grain structure. Such a combination of micro- and nanotopography of the surface of spider silk-proteins is considered ideal for culturing eukaryotic cells. 

Figure 3. The spatial architecture and mechanical properties of the rSS-PCL scaffold: (A,B) SEM images, (C), 2D and (D), 3D – atomic force microscopy images, (E). typical Young's modulus distribution obtained by nanoindentation.
The Young’s modulus of the rSS-PCL scaffold in the hydrated state, measured by nanoindentation, was found to be $287 \pm 32$ kPa (Fig. 3E) and did not change after the scaffold was pressed under a 2 ton weight, which indicates an extraordinary ability of the material to sustain its spatial architecture and mechanical properties. Finally,

**Figure 4.** drNPC cultured in the rSS-PCL scaffold with or without the PRP-based liquid matrix (SPRPix matrix). (A–C) SPRPix (PRP + rSS-PCL scaffold); (A) Nestin (green), auto-fluorescence of the SPRPix matrix (red). (B) Spontaneous neuronal and glial differentiation revealed by MAP2 (green) and GFAP (red). (C) Intense proliferation of SOX2-positive (green) drNPCs, and βIII-tubulin-positive (red) neuronal progenitors. (D) The same concentration of drNPC cells seeded on rSS-PCL scaffold only. Nestin (green), βIII-tubulin (red). (E,F) high magnification examples of drNPCs cultured on rSS-PCL alone (E) or on SPRPix (F). Bar = 50 μm.
studies of the wetting properties of the rSS-PCL scaffold demonstrated good hydrophilicity, with an equilibrium contact angle of 63° ± 7°C.

Taken together, the physico-chemical properties of the rSS-PCL scaffold (unidirectionally oriented structure, developed micro- and nanotopography, appropriate elasticity and integrity) support its use as a carrier for a growing neuronal network.

**drNPC culture in Chondro-gide® and SPRPix matrix.** When drNPC were placed into Chondro-Gide® in complete medium alone, good cell adhesion to collagen fibers was observed; however, the proliferative activity and neural differentiation of the cultured drNPC was very low (Suppl. Fig. 2A). Seeding the same number of drNPC onto Chondro-Gide® in a mixture of PRP and complete medium, with the subsequent gel-clot formation on the scaffold surface, promoted a substantial increase in both proliferation and neural differentiation of drNPC in the gel phase of the two-component matrix (Suppl. Fig. 2B). Processes of differentiated neuronal progenitor cells were observed along the collagen fibers (which had a random spatial orientation).

In contrast to what was observed in Chondro-Gide®, drNPC cultured on the two component SPRPix matrix (PRP hydrogel embedded into the rSS-PCL scaffold) demonstrated a high level of adhesion and neuronal differentiation: drNPC completely covered the entire SPRPix matrix surface and formed long βIII-tubulin- and MAP2-positive processes oriented along the spidroin microfibrils (Fig. 4A–C,E). In contrast, omission of the PRP hydrogel from the culture led to the formation of drNPC “islets” with short, scarcely oriented processes (Fig. 4D,F).

When compared with Chondro-gide® plus PRP, the SPRPix matrix further increased both the proliferative activity of the seeded drNPC and the fraction of cells that underwent neuronal differentiation. Using the NIS Elements software (Nikon), we counted the total cell number (via the Hoechst channel) and the number of neuronal progenitors within the 640 × 640 × 25 μm (x × y × z) volume. The analysis of the ratio of Hoechst-positive cells (total number of cells) and βIII-tubulin positive cells (neuronal progenitors) showed that, in the case of the SPRPix matrix, the absolute numbers of cells/neuronal progenitors in the studied volume were 2600 ± 364/930 ± 148 after two weeks of culture, as compared to 1850 ± 296/174 ± 29 when cells were plated in rSS-PCL scaffold only (ie, without PRP). Thus, the total number of attached viable cells was 1.4 fold higher for the two-component SPRPix matrix than the single-component rSS-PCL scaffold (p < 0.05 by the Student’s t-test), while the fraction of neuronal progenitors was 3.8 times higher (9.4% vs 35.8% for the rSS-PCL scaffold and SPRPix matrix, respectively) (Suppl. Fig. 4). The absolute number of neuronal progenitors in the rSS-PCL scaffold and SPRPix matrix differed by more than 5 times (p < 0.01 by the Student’s t-test).

**Neurite alignment estimation in SPRPix matrix images by Fourier-transform statistical analysis.** In order to study neurite orientation in the SPRPix matrix, we analysed βIII-tubulin-positive cells using FFT statistics (summarized in Fig. 5). The first column shows original grayscale images used for statistical analysis. The second column shows the resulting FFT images (Fig. 5A). The intensity of these images is shown in a logarithmic scale linearly fitted to the [0,1] interval. The red lines in the plot show equivalent covariance ellipses. The third column shows a normalized angular distribution of FFT image sector sums fitted with a normal distribution. The results of directionality analysis are summarized in Supplemental Table 1. The SEM image of the rSS-PCL scaffold (Fig. 5A) shows high directionality with ε close to 1 and standard deviation σ below 30°. All images of neurite structures grown on anisotropic scaffolds demonstrate noticeable directionality with eccentricity form 0.46 to 0.79 and corresponding standard deviation between 40° and 50°. Higher standard deviation σ values were observed for the two-component SPRPix scaffold (Fig. 5B,C) compared to that of anisotropic rSS-PCL scaffold (Fig. 5D). The control image of drNPC without any scaffold shows the lowest eccentricity ε < 0.2. Taken together this data shows that the degree of neurite orientation is statistically higher on anisotropic scaffolds compared to no-scaffold control which shows almost isotropic growth (Fig. 5E).

**In vivo study of biocompatibility of SPRPix matrix in nonhuman primates.** As a final experiment, we implanted the SPRPix matrix seeded with human drNPC into the cerebral cortex and spinal cord of two NHPs (non human primates). The animals tolerated the surgeries well and no adverse events were detected during the in-life portion of the study, up to 12 weeks post implantation. Histopathology revealed no findings of microglial infiltration or gliomesodermal reaction in the host brain (Fig. 6A–C). In addition, there was insignificant astroglial response (Fig. 7) as reactive astrocytes were almost absent. Further immunohistochemical analysis with T-cell and microglial cell/macrophage markers revealed only a few infiltrating CD3-positive T-cells (Fig. 6D,E) and CD68 positive microglial cells (Fig. 6F–H) near the SPRPix matrix. The surrounding neural tissue demonstrated no signs of immune cell and microglial infiltration (Fig. 6G). Taken together, these data support biocompatibility of the drNPC-SPRPix combo in the NHP brain and spinal cord.

In order to reliably identify the implanted cells, two strategies were used. First, given that human female cells were implanted in male NHP recipients, we used antibodies against the macro-H2A.1 histone, which specifically labels the X-inactivated chromosome in the nuclei of female cells. Second, two human specific antibodies – one against mitochondria (h-Mito), and another against a cytoplasmic antigen (STEM-121) were used to reliably identify human cells in the NHP host.

We found numerous human female cells in the implanted NHP brain (Fig. 7). h-Mito positive (Fig. 7A,B) and STEM121 positive drNPCs (Fig. 7C,D) were detected in and around the SPRPix. Staining with βIII-tubulin indicated that the transplanted human cells differentiated along the neuronal lineage (Fig. 7A–C), while glial differentiation along the astrocytic lineage was rare (Fig. 7D). We further confirmed that the antibodies used to identify human cells did not react with naïve NHP brain (Fig. 7E,F).

Very similar observations were recorded in the spinal cord by both histopathology (Fig. 8) and immunostaining (Fig. 9). In the implanted spinal cord, Nissl staining around SPRPix demonstrated lack of infiltration by host
immune cells and histiocytes (Fig. 8A,C). No pathological changes were found neither in the white nor the gray matter directly adjacent to the implanted SPRPix matrix (Fig. 8A,C,E,F). In order to characterize the structure of the extracellular matrix surrounding the SPRPix, we performed an AFM study of the dried de-paraffinized sagittal spinal cord sections (Fig. 8G–H). This study showed that the SPRPix matrix was fully integrated within the surrounding tissue, and a loose, rather isotropic fine-fibered extracellular matrix (similar to that of normal
neural tissue), was observed in the direct proximity of the SPRPix. This extracellular matrix did not contain any tightly packed oriented collagen fibrils characteristic of a fibrotic matrix or scar. Nissl staining also revealed some cells that morphologically resembled interneurons around SPRPix (Fig. 8C); further staining confirmed that these

Figure 6. Histological and immunohistochemical (IHC) analysis of the endogenous immune response to the drNPC-SPRPix implanted into the brain cortex of a rhesus macaque. (A) Nissl stain; magnification $\times 100$ (B,C). H&E stain of the interface between the SPRPix and the surrounding host tissue; magnifications $\times 100$ (B) and $\times 200$ (C). (D,E). Representative images of CD3 staining showing low level of T cell infiltration in the brain tissue adjacent to the drNPC-SPRPix implant. (F–H) Representative images of CD68 staining showing examples of microglial/macrophages in the brain adjacent to the drNPC-SPRPix implant (G) and among the human cell graft (H). (E,H) depict the areas with the highest immune cell infiltration noted. Bar size 100 $\mu$m. Notations: rSS-PCL-spidroin-polycaprolactone scaffold; Ctx – brain cortex.
Figure 7. Immunofluorescence analysis of the brain of a rhesus macaque at the site of implantation of the SPRPix matrix with human drNPC. (A) Representative low magnification image of the site of drNPC-SPRPix implantation showing staining by human specific anti-mitochondria antibodies (h-Mito) and some co-expression with βIII-tubulin. (B) Higher magnification of h-Mito and βIII-tubulin staining. Arrow shows co-localization of h-Mito and βIII-tubulin indicating neuronal differentiation of transplanted human drNPCs. (C) Human specific staining by STEM-121 co-localized with anti-βIII-tubulin (arrow). (D) Staining with STEM-121 and anti-GFAP revealed no co-localization. (E–F) Staining of normal brain tissue shows absence of h-Mito (E) and STEM-121 (F) staining of non-transplanted NHP host tissue. Bar 100 µm. Laser scanning confocal microscopy.
Figure 8. Histological and immunohistochemical (IHC) analysis of the SPRPix matrix with human drNPC implant with AFM imaging in the projection of the posterior column of the spinal cord of the rhesus macaque. (A) Sagittal section of the spinal cord stained with H&E. (B) Same spinal cord fragment, stained with antibodies to macro-H2A.1 visualized by immunoperoxidase staining. Transplanted cells highlighted by arrows. (C) Enlarged H&E image of (A) demonstrating the cells adjacent to the SPRPix matrix; Bar size = 50 µm. (D) Same image as in (C), stained with antibodies to MAP2. (E,F) Sagittal sections of the spinal cord directly below the SPRPix matrix, Nissl (G) and H&E (H), stains. (G,H) 2D AFM image (overlay of two images from adjacent regions) and a 3D AFM image of the border between the SPRPix and the ECM of the surrounding tissue, showing full integration of the scaffold with the ECM. In panels (A,B,D–F) bar size = 100 µm. Notations: rSS-PCL-spidroin-polycaprolactone scaffold; GM – grey matter; WM – white matter; ECM – extracellular matrix.
cells were MAP2 positive (Fig. 8D) and some of them were double positive for h-Mito and βIII-tubulin (Fig. 9A, Suppl. Fig. 6) which suggests that at least some of transplanted human cells migrated into the spinal cord tissue and differentiated into neurons.

Finally, additional immunofluorescence analysis revealed absence of GFAP-positive reactive astrocytes around SPRPix (Fig. 9B,C), where the presence of numerous human nestin-positive cells was noted (Fig. 9C,D). We also detected βIII-tubulin positive human neuronal progenitors (Fig. 9A) as well as MAP2-positive human neurons between the SPRPix matrix and the spinal cord, and these neurons extended long processes in the spinal cord (Fig. 9D).

Discussion

Previous studies have demonstrated that transplantation of small (up to 500 µm) tissue engineered constructs outperform the transplantation of suspended cells alone based on a number of cell survival indices. Various factors are known to underlie this observation: transplantation of tissue engineered constructs eliminates cell detachment stress due to the use of proteolytic enzymes and preserves cell-cell contacts that are important for cell survival and function. Furthermore, transplantation of cell constructs allows pre-differentiation of cells in culture, ensuring that the right cell phenotype is administered, presumably enhancing transplantation kinetics and functional outcomes.

Our work focused on developing a complex matrix with two highly relevant materials for culturing human NPCs and for implantation into the CNS: PRP and spidroins. Due to its growth and survival-rich factor content, PRP has been regarded as a potential substrate for the formation of 3D neural tissue engineered constructs. PRP is also considered a biomimetic scaffold as it creates a natural microenvironment for NPCs, in which the cells, embedded in the hydrogel, may interact as they would in normal ontogenesis. Indeed, PRP-based biomimetic microenvironment is used to fill neural conduits during reconstructive surgery of peripheral nerve injuries. Silkworm-produced silk fibroin and proteins of the dragline silk of orb-weaver spiders’ web – spidroins – are characterized by unique biomechanical and adhesive properties. Many researchers consider them as a prospective material for creation of recombinant tissue engineered scaffolds, including those for neural tissue. Silk proteins have been mostly investigated for the development of scaffolds for regeneration of peripheral nerves and much less explored in the context of CNS regeneration. Interestingly, recombinant spidroin coating of cell culture surfaces has been shown to promote adhesion and neuronal differentiation. In our study, the two materials were combined for the creation of a two-component scaffold based on PRP and an anisotropically structured spidroin scaffold.

We first showed that the “liquid matrix” consisting of 25% PRP hydrogel dramatically increased proliferation and neuronal differentiation of drNPCs both when culturing on a non-adhesive plastic and within the structure of the two-component SPRPix matrix, demonstrating its critical value as part of an engineered construct aimed
at supporting regeneration of neural tissue. The observed stimulation of drNPC proliferation and neuronal differentiation in the SPRPix matrix is also likely due to the anisotropic rSS-PCL scaffold properties, including the presence of a large number of repeats (18 repeats) of the GRGGL sequence in rS1/9 that is a signal sequence for the growth of neural cells and serves as a ligand for binding with the NCAM receptors on the surface of neural cells. The other favourable factor is the large positive net charge of the rS1/9 and rS2/12 molecules (+28 and +36 at pH 7.0, respectively) which facilitates adhesion of neural cells to the scaffold via the NCAM receptors. The high stiffness of the prepared scaffolds is also of great importance, since it is known to play a notable role in the proper differentiation of neural cells and axon growth, and the rather high hydrophilicity (65%), in spite of the presence of a more hydrophobic PCL, is significant as well.

In addition, the nanotopographical relief of the microfibril surface formed in the process of electrospinning is of particular importance for enhancement of biocompatibility of the 3D anisotropic spidroin scaffold as it facilitates efficient adhesion of neural progenitors and their processes on the scaffold surface, while the oriented architecture of microfibrils promotes oriented axon growth. The calculated alignment of the extended neurites, based on the Fourier-transform analysis of the microscopic images, shows that the axon growth is more oriented on the SPRPix matrix as compared to the components of PRP or rSS-PCL scaffold alone. This is probably due to the enhanced adhesion of neurites to anisotropic rSS-PCL provided by the PRP-based hydrogel.

When designing scaffolds for use in CNS tissue engineering a highly desirable feature is biocompatibility with nervous tissue. Spider silk spidroins are biocompatible and biodegradable and typically induce only a mild inflammatory response that decreases within a few weeks post-transplantation. It has been previously shown that, due to their biodegradability, recombinant spidroins can be remodelled in the body allowing vascularization, innervation and tissue ingrowth with the eventual complete replacement by the native bone and connective tissue.

Biocompatibility of recombinant spidroin-1 (known as rS1/9) has been shown in rodent experiments and shown that the SPRPix matrix as compared to the components of PRP or rSS-PCL scaffold alone. This is probably due to the enhanced adhesion of neurites to anisotropic rSS-PCL provided by the PRP-based hydrogel. When designing scaffolds for use in CNS tissue engineering a highly desirable feature is biocompatibility with nervous tissue. Spider silk spidroins are biocompatible and biodegradable and typically induce only a mild inflammatory response that decreases within a few weeks post-transplantation. It has been previously shown that, due to their biodegradability, recombinant spidroins can be remodelled in the body allowing vascularization, innervation and tissue ingrowth with the eventual complete replacement by the native bone and connective tissue. Our study in two NHPs provided initial indications of biocompatibility and safety: the SPRPix matrix did not cause lymphocytic and microglial infiltration nor resulted in active astrogliosis and gliomesodermal reactions. Furthermore, the SPRPix matrix supported the survival of the transplanted human drNPCs within the engineered construct for at least 12 weeks post-transplantation. A subpopulation of the transplanted cells retained the expression of nestin and SOX2, while a significant portion of the cells became MAP2-positive neurons.

Our in vivo study did not employ the use of immunosuppressive drugs yet the human cells still survived in the NHP CNS. The absence of a notable reaction from the recipient immune system may be explained by low level of HLA-DR expression, weak antigen-presenting function, and/or the creation of a local immunosuppressive microenvironment. Indeed, iPS-derived neural stem/progenitor cells appear to exhibit these characteristics and the drNCP used in our study may behave in a similar way, with the additional advantage of being further protected from an initial host immune reaction due to their integration in a complex matrix. Future studies will investigate the mechanisms by which the SPRPix matrix promotes engraftment of drNPCs into the host CNS tissue without scar formation and minimal immune reactivity.

When addressing specifically the regeneration of the injured spinal cord, it is important to ensure not only the most favourable conditions for cell survival, but also for the promotion of oriented axonal growth. The SPRPix matrix appears to support both of these aspects in vivo upon implantation into the normal host spinal cord. This was achieved with minimal host immune response and no scar formation, demonstrating that the developed SPRPix matrix might be superior to other types of tissue engineered constructs like polysaccharide based hydrogel scaffolds and other polymeric scaffolds. Further studies will investigate the efficacy of drNPC-seeded SPRPix in spinal cord injury models.

Conclusion

The two-component SPRPix matrix described in this study, consisting of a “liquid” component based on a PRP-derived hydrogel within a rSS-PCL scaffold, dramatically stimulated proliferation and neuronal differentiation of drNPCs due to the complex action of the PRP components and creation of a 3D biomimetic environment with rSS-PCL that critically supported drNPC growth and differentiation. The rSS-PCL scaffold not only provided adhesion of drNPCs to the surface, but also functioned as a guiding scaffold, specifying a vector for axon growth resulting in aligned parallel neurite outgrowth. This study demonstrated the biocompatibility and safety of the two component SPRPix matrix in the NHP brain and spinal cord. Finally, our study provided preliminary evidence of integration and differentiation of the implanted NPCs into the host brain and spinal cord. The tissue engineered construct resulting from combining drNPC with the two-component SPRPix matrix exhibits important attributes that warrant further examination in spinal cord injury treatment.

Materials and Methods

Preparation of human neural precursor cells. Primary-reprogrammed human neural precursor cells (hereafter – directly reprogrammed neural precursor cells, drNPC) were provided by New World Laboratories Inc. Cryopreserved cells were thawed from liquid nitrogen and seeded onto laminin-coated plates in NeuroCult-XF Basal Medium (StemCell Technologies) with 1% Pen-Strep (Gibco), B-27 Supplement (50×) (Gibco), 20 ng/mL of bFGF and 20 ng/mL of EGF (complete medium). The cells were incubated at 37 °C in 5% CO2 and 5% O2; medium replacement was performed every 2 days. As needed, cells were passaged at a ratio of 1:3 when 80% confluence was reached.
For differentiation studies, a standard differentiation protocol was used: drNPC monolayers were switched to Neurobasal media ( Gibco, USA) supplemented by Glutamax (1 ×, Gibco, USA), B27 (1 ×, Gibco, USA), CultureOne (1 ×, Gibco, USA), BDNF (20 ng/ml), GDNF (20 ng/ml) and Ascorbic Acid (200 μM) for 14 days.

**Immunocytochemical analysis.** The expression of neural stem cell, neuronal and glial markers was performed by immunocytochemical analysis. The cells were fixed by adding 4% buffered formaldehyde solution containing 0.1% saponin. Primary antibodies to Nestin (R&D, 2 μg/ml), SOX2 (BD Biosciences, 5 μg/ml), βIII-tubulin (R&D, 2 μg/ml), MAP2 (Sigma-Aldrich, 5 μg/ml), GFAP (DAKO, 5 μg/ml), and NF-200 (Sigma-Aldrich, 5 μg/ml) were used together with Alexa Fluor 488 goat anti-mouse IgG (H + L) and Alexa Fluor 633 goat anti-rabbit IgG (H + L) secondary antibodies (all 1:400; Invitrogen, USA). The cell nuclei were labeled with Hoechst 33342 stain (Thermo Fisher Scientific). Immunofluorescence was analyzed using a Nikon A1 scanning laser confocal microscope (Nikon Co., Japan). All the staining studies were conducted in series, with 5 repeats in each series.

**Spidroin coating of low-adhesive surfaces.** drNPCs-01 were cultured on low-adhesive surfaces coated with spidroin (see biosynthesis details below) using Petri dishes with a cover glass bottom in the center (Confoical Petri Dishes, SPL Life sciences). For the cover glass coating with spidroin, 1% solution of the recombinant rS1/9 in 96% formic acid was poured into a sterile dish and incubated for 60 minutes, followed by the solution removal, washing of the dish with 96% ethyl alcohol and drying under UV-radiation in a laminar flow hood.

**Biosynthesis of the anisotropically structured complex spidroin matrix.** The yeast biomass growth of *S. cerevisiae* SCR-702T-1F9 and SCR-702T-2E12, and the extraction and purification of the recombinant spidroins rS1/9 and rS2/12, were conducted as previously described41. Briefly, the yeast cells were grown in a 3 L fermentor in 96% formic acid was poured into a sterile dish and incubated for 60 minutes, followed by the solution removal, washing of the dish with 96% ethyl alcohol and drying under UV-radiation in a laminar flow hood.

*Preparing the anisotropically structured complex spidroin scaffold.* rSS-PCL by electrospinning, a NANON – 01A apparatus equipped with a standard spinneret and a drum collector with a 100 mm width was used with a solution containing the recombinant spidroins rS1/9 (6%) and rS2/12 (1%), PCL (1%, "Biopolymer"), Ukraine) in the ratio of 6:1:1, in 1,1,1,3,3,3-hexafluoro-2-propanol (HFIP; Fluka Chemie GmbH, Germany). The solution was fed at a rate of 0.4 mL/hr. The process was conducted at a voltage of 9.5 kV, with a collector rotation rate of 1000 rpm at a 10 cm distance from the capillary to the collector. When the process was finalized, the prepared material was incubated in 96% ethanol for 15 min, and then dried for 18 hrs at room temperature in a laminar flow hood.

**Preparation of PRP.** All research involving human participants was performed in accordance with relevant guidelines/regulations. PRP was prepared from blood of three healthy male donors, after each donor had signed an informed consent for using their plasma for research purposes. The experimental protocol with the participation of blood donors was approved by the Local Ethics Committee of the Federal Research and Clinical Center of Specialized Medical Care and Medical Technologies of FMBA of Russia (Protocol No.10a of September 12, 2016). For preparation of PRP, we harvested 42 mL of donor blood into a syringe pre-filled with ACDA (acid citrate dextrose buffer Type A) (Hemocon) at a ratio of 1.4 mL per 10 mL. The harvested blood was centrifuged in an Eppendorf 5810 R centrifuge in 50 mL tubes (Corning) at 280 g and 20 °C for 15 min with acceleration of 3/3. Using a serological pipette, the upper 10–15 mL of plasma was collected above the erythrocyte layer and the Buffy coat, depending on the donor’s hematocrit, into a 50 mL tube. The sample was repeatedly centrifuged at 280 g and 20 °C, for 15 min; the upper two-thirds of the plasma was removed (Platelet poor plasma), the bottom one-third was re-suspended in the remaining volume, and the number of thrombocytes was estimated with a hemocytometer. We then selected PRP samples with the platelet count no less than 5 × 10^11/μL.

**Cultivation of drNPCs in PRP-based “liquid matrix”.** Monolayer drNPCs were removed from the surface of culture flasks by incubation with 2 mL of Trypsin-EDTA 0.25% ( Gibco) after washing with Dulbecco’s phosphate-buffered saline (D-PBS) ( Gibco). After the incubation, the trypsin-EDTA was inactivated with conditioned medium, and the suspended cells were transferred into a 50 mL tube, a cell count was performed using a TC20 automated cell counter (Bio-Rad), and the sample was centrifuged at 400 × g and 20 °C for 5 min. The supernatant was aspirated and the cell pellet re-suspended in 750 μL of the NeuroCult-XF Basal Medium (StemCell Technologies). 250 μL of the freshly prepared PRP was added to the cell suspension in NeuroCult-XF Basal Medium, resulting in a cell concentration of 4 million/ml that was then carefully re-suspended. To neutralize the sodium citrate (anti-clotting agent) in the Hemocon buffer, 2 μL of 10% CaCl2 per 100 μL of the medium was added and the solution immediately transferred into Petri dishes with a glass cover bottom (SPL Life sciences). The gel-clot formation occurred in the Petri dishes during 20 min at 5% CO2, 37 °C. Then, 1.5 mL of Complete medium was added to each Petri dish. The dishes were incubated at 5% CO2, 37 °C for 2 weeks, and the medium was replaced every 2 days.
Preparation of two-component matrices from PRP and anisotropically structured complex scaffold rSS-PCL (SPRPix). To create the two-component tissue engineered construct, the required size was cut with a scalpel from the sterile complex scaffold, prepared as described above, and placed in the center of a glass cover bottom Petri dish. 50 µL of the drNPC cell suspension (with a concentration of 4 million/mL), mixed with 25% PRP in the NeuroCult-XF Basal Medium containing CaCl₂, prepared as described above, was placed in layers onto the scaffold and incubated in 5% CO₂ at 37 °C until gel-clot formation. Then, 2.5 mL of Complete medium was added. The dishes were incubated at 37 °C in 5% CO₂ medium was replaced every day.

Five independent batches of SPRPix were made for each SPRPix batch, a total of five replicas were prepared. As a control, we also used a commercially available collagen scaffold «Chondro-Gide®» (Geistlich Pharma AG, Switzerland) which was seeded by the same number of drNPC with or without PRP (50 µL of the drNPC cell suspension with a concentration of 4 million/mL), following by culturing during 2 weeks before fixation and immunostaining.

SEM, AFM and mechanical studies. AFM images of the anisotropic complex scaffold. Were acquired on air in the PeakForce Tapping® Mode, using a MultiMode 8 atomic force microscope with a Nanoscope V controller and E scanner (Bruker). AFM imaging was conducted with RTESPA-300 probes (Bruker) with a nominal spring constant of 40 N/m, a nominal frequency of 300 kHz and a nominal tip radius of 8 nm. 10 × 10 µm images were obtained at a scan rate of 1 Hz and a 512 × 512 pixels resolution. The raw AFM images were processed using the NanoScope Analysis v.1.10 software (Bruker).

Mechanical studies. Of anisotropic scaffold microfibrils were performed using a PIUMA Nanoindenter (Optics11, Amsterdam, Netherlands), which included a controller, an optical fiber, and a spherical probe for the force-displacement curves' acquisition. The probe of a 26.5 µm radius was attached to a flexible cantilever with the spring constant of 0.44 N/m in contact with the optical fiber to measure its displacement.

Anisotropic scaffolds (4 samples in total) were studied in water over the area of 500 × 500 µm, with a 50 µm increment by each axis. To obtain the force-displacement curves, the probe was immersed for 5 µm into a scaffold sample at each point of measurement. The Young's modulus for each point was computed according to the Hertzian contact mechanics model for a spherical body indenting a flat surface, using the built-in PIUMA software.

Contact angle studies. In order to assess the wetting properties of the anisotropic scaffold, we conducted contact angle measurements. We applied a sessile drop technique at 25 °C and a relative humidity of 30% with an experimental optical unit described in Ref. 42. A 50 µL drop of distilled water, stained with Bengal Rose (Acros Organics, India), was placed onto a studied sample's surface. The contact angle measurements were performed in the continuous video regime for 300 seconds, with a total of 9 measurements.

Fourier-transform image analysis. For a quantitative estimation of neurites' alignment in the anisotropic rSS-PCL scaffolds we used a numerical analysis of the digital images obtained with SEM (for the scaffold alone) and with scanning laser confocal microscopy (for the tissue engineered constructs) using two-dimensional fast Fourier transform (FFT). The images were 8-bit grayscale TIFF files. All files were cropped to the same size of 825 × 825 pixels. A custom script in MATLAB (MathWorks Inc., USA) was written for the image analysis, allowing FFT of grayscale images to be generated by the $\text{fft2}$ function producing a complex-valued matrix of the same size as the original image. Modulus of FFT was treated as a probability distribution function in the frequency domain, and its asymmetry characterizes alignment of structures in the image⁴³. For a higher accuracy of statistical image analysis we reduced the noise of FFT applying a cut-off to the FFT matrix: all values below a preset limit were zeroed. The limit has been chosen based on the noise level estimated from the maximum values of 64 × 64 pixel subarrays in the corners of the whole FFT matrix. Second-order statistical moments were used to characterize the asymmetry of FFT distributions⁴⁴. Three independent second-order central moments determine an equivalent ellipse in the frequency domain. The shortest axis of the ellipse indicates the prevailing direction of the structure alignment in the original image (mean value $\mu$ of angular distribution). Eccentricity of the ellipse $\epsilon$ is a measure of alignment dispersion. For a randomly oriented structure eccentricity tends to 0; for a totally aligned structure eccentricity approaches 1. The main numerical measure of alignment is the angular standard deviation $\sigma$. In this experiment, $\sigma$ was calculated using a weighted sample variance of normal distribution fitting the dataset. These calculations were performed for each series of experiments.

In vivo study of biocompatibility on healthy primates. The experimental protocol for the study of biocompatibility of the neural tissue engineered SPRPix constructs containing human drNPC on healthy nonhuman primates was approved by the Local Bioethical Committee of the Research Institute of Medical Primatology of Russian Academy of Sciences (Protocol of July 13, 2016). The experiment was conducted on two healthy male rhesus macaques (Macaca mulatta) born in 2012 (Table 1) homed at the farm of the Research Institute of Medical Primatology of Russian Academy of Sciences in Sochi. One month prior to the study, the animals were relocated from the shared cage into individual cages with free access to water and food. One month post-quarantine, the surgical procedure of implanting the prepared neural tissue engineered constructs into the animals was performed.

Surgical protocol. The food supply and water were removed 12 and 4 hours, respectively, prior to the surgical procedure. In order to ensure their safe removal from the cages, animals received an intramuscular injection of Sedexin (1 mg/kg) and Ketamine (10 mg/kg). Their hair was then clean shaved from the back and head, the Tibial vein was catheterized, intubation was performed and the animals were transferred into a sterile operation room,
where they were connected to a ventilator with a semi-closed circuit with the following parameters: tidal volume 120–150 mL, respiratory rate 23–26 bpm, FiO₂, 0.4. Anesthesia was achieved via a bolus dosing of fentanyl and inhalation of isoflurane 1.2–2.0 vol. Prior to the surgery, the animals were also injected with ketonal (10 mg/kg), dexamethasone (1 mg/kg), Tranexam (20 mg/kg) for desensitization and multimodal anesthesia.

**Implantation into the brain.** The skin was scrubbed with iodine and alcohol. An incision of the cutaneous covering was performed followed by a layer-by-layer dissection of the aponeurosis. Using a neurosurgical burr, two milling holes with the diameter of 5 mm were applied in the projection of the parietal fissure, followed by the section of the dura mater and dull stratification of the pia mater with hemostasis. Using microsurgical forceps, 2–3 neural tissue engineered constructs sized 2 × 8 × 0.3 mm were inserted in the brain cortex at a depth of 5–7 mm. Hemostasis was controlled, then the dura mater, aponeurosis and head skin were consecutively sutured. Sterile bandage was applied to the wound.

**Implantation into the spinal cord.** The skin was scrubbed with iodine and alcohol. A 8–10 cm longitudinal midline skin incision was made to expose the T8–11 vertebrae, and paraspinal muscles were separated subperiosteally. A T8 interlaminectomy was performed, and the dura mater was exposed, which was dissected longitudinally in the projection of the corresponding segment. 2–3 tissue engineered constructs sized 2 × 15 × 0.3 mm were placed on the surface of the spinal cord in the projection of the posterior spinal column and slightly inserted by means of a careful dissection of fibers of the posterior spinal column. The plastic closure of the dura mater was performed with a fascial flap, and the dura mater suture was sealed with fibrin glue. The wound was sutured layer-by-layer and covered with an aseptic bandage.

**Termination and histology.** 12 weeks after the SPRPix matrix implantation, the animals were anesthetized by an IV injection of ketamine (20 mg/kg), followed by an overdose of propofol administered intravenously. After apnoea and atomic coma took place, the animals’ thorax and mediastinum were dissected, the right atrium auricle was opened for blood removal, the left ventricle was punctured, a cannula was placed transcardially and guided through the aortal valve up to the ascending aorta. Perfusion with 10% buffered formalin solution was carried out though the cannula in a volume equal to the body weight (up to 5L). Then, the brain and the entire spinal column were separated and fixed for 24h in the same solution at 4°C. The spinal cord in the projection of the implantation location (plus two segments above and below) was separated from the fixed preparations of the spinal column. The prepared spinal cord and brain samples were dissected along the medial line so that the implanted constructs were both on the right and on the left side. 50 µm-thick sagittal vibratome sections were made from the left preparations and used for immunofluorescent staining and confocal microscopy. Paraffin blocks and 5–6 µm-thick paraffin sections prepared from the other side were used for routine histology studies with cresyl violet (Nissl staining) and hematoxylin-eosin as well as immunohistochemical analysis.

**Immunohistochemistry.** Immunohistochemical analysis was performed using thick vibratome sections, as well as paraffin sections, by means of both fluorescent and immunoperoxidase detection. The spinal cord and brain sections were stained using antibodies to Nestin (R&D), SOX2 (BD Biosciences), βIII-tubulin (R&D), MAP2 (Sigma-Aldrich), GFAP (DAKO), NF-200 (Sigma-Aldrich,) macroH2A.1 (ab183041, Abcam), human-specific anti-mitochondria antibody (ab92824, Abcam), and STEM-121 (Y40410, TakaraBio), anti- CD3, CD31 and CD68 (Roche Diagnostics) (all as 5 µg/ml solutions in PBS). In the case of immunofluorescence detection with the use of vibratome sections, Alexa Fluor 488 goat anti-mouse IgG (H + L) and Alexa Fluor 633 goat anti-rabbit IgG (H + L) (all dilutions 1:400; Invitrogen, USA) were used as secondary antibodies, the fluorescence was detected with a Nikon A1 laser scanning confocal microscope (Nikon Co., Japan). Immunoperoxidase staining was performed on thin paraffin sections with a Benchmark ULTRA immunostainer (Ventana, USA) utilizing the manufacturer-recommended detection system.

**References**

1. Singh, A., Tetreault, L., Kalsi-Ryan, S., Nouri, A. & Fehlings, M. G. Global prevalence and incidence of traumatic spinal cord injury. *Clin. Epidemiol.* 6, 309–331 (2014).
2. Maynard, F. M. Jr. et al. International Standards for Neurological and Functional Classification of Spinal Cord Injury. *American Spinal Injury Association. Spinal Cord. 35*(5), 266–74 (1997).
3. Krucoff, M., Rahimpour, S., Slutsky, M., Edgerton, V. & Turner, D. A. Enhancing Nervous System Recovery through Neurobiologics, Neural Interface Training, and Neurorehabilitation, *Front Neurosci.* 10, 584 (2016).
4. Badner, A., Siddiqui, A. M. & Fehlings, M. G. Spinal Cord Injuries: How Could Cell Therapy Help? *Expert Opin Biol Ther.* 17(5), 529–541 (2017).
5. Yousefiard, M. et al. Neural stem/progenitor cell transplantation for spinal cord injury treatment; A systematic review and meta-analysis. *Neuroscience.* 322, 377–97 (2016).

| Animal identifier | Gender | Date of birth  | Weight at the study start (kg) | Duration of observation | Regions of implantation |
|-------------------|--------|----------------|------------------------------|------------------------|------------------------|
| 40400             | Male   | Nov. 20 2012   | 5.4                          | 12 weeks               | 1. Brain; Parietal lobe 2 cm from the sagittal suture bilaterally. |
| 40014             | Male   | Apr. 20 2012   | 5.6                          |                        | 2. Spinal cord: Lower thoracic T7 – T8 level, posterior surface of the spinal cord in the projection of the posterior spinal column. |

Table 1. Details and implantation locations in experimental animals.
6. Tabakov, P. et al. Functional regeneration of supraspinal connections in a patient with transected spinal cord following transplantation of bulbar olfactory ensheathing cells with peripheral nerve bridging. Cell Transplant. 23(12), 1631–55 (2014).
7. Muheremu, A., Peng, J. & Ao, Q. Stem cell based therapies for spinal cord injury. Tissue Cell. 48(4), 328–33 (2016).
8. Doulames V. M., Plant G. W. Induced Pluripotent Stem Cell Therapies for Cervical Spinal Cord Injury. Int J Mol Sci. 17(4) (2016).
9. Cefalo MG. et al. Human iPSC for Therapeutic Approaches to the Nervous System: Present and Future Applications. Stem Cells Int. 2016;4869071 (2016).
10. Zheng, J. et al. A combination of small molecules directly reprograms mouse fibroblasts into neural stem cells. Biochem Biophys Res Commun. 476(1), 42–8 (2016).
11. Mauksch, C., Jones, K. S. & Connor, B. Concise review: the involvement of SOX2 in direct reprogramming of induced neural stem/ precursor cells. Stem Cells Transl Med. 2(8), 579–83 (2013).
12. Velasco, J. et al. Concise review: Generation of neurons from somatic cells of healthy individuals and neurological patients through induced pluripotency or direct conversion. Stem Cells. 32(7), 2811–7 (2014).
13. Aflhors J.-E., Elayoubi R. Methods for reprogramming cells and uses thereof. United States Patent Application No. 61/256,967, 2009 Oct 31; International Patent Application No. PCT/CA2010/001727, 2010 Nov 1, United States Patent No. 9,453,205.
14. Aflhors J.-E. Direct Reprogramming of Human Adult Fibroblasts into Stable Neural Stem-Like Cells without Genetic Engineering. Presented at: Stem Cell Network Annual General Meeting, Montreal, Canada, November 1, (2009).
15. Nagesh, N. et al. Human Spinal Oligodendrogenic Neural Progenitor Cells Promote Functional Recovery After Spinal Cord Injury by Axonal Remyelination and Tissue Sparing. Stem Cells Transf Med. 7(11), 806–818 (2018).
16. Romanyuk, N. et al. Beneficial effect of human induced pluripotent stem cell-derived neural precursors in spinal cord injury repair. Cell Transplant 24(9), 1781–1795 (2015).
17. Fuhrmann, T. et al. Injectable hydrogel promotes early survival of induced pluripotent stem cell-derived oligodendrocytes and attenuates longterm teratoma formation in a spinal cord injury model. Biomaterials. 83, 23–36 (2016).
18. Guan X. et al. Development of hydrogels for regenerative engineering. Biotechnol J. 12(5) (2017).
19. Silva, J. et al. Fibrin functionalization with synthetic adhesive ligands interacting with αvβ1 integrin receptor enhances neurite outgrowth of embryonic stem cell-derived neural stem/progenitors. Acta Biomater. 51742–70661 (2017), 30441–5 (2017).
20. Goh K. L., Holmes D. F. Collagenous Extracellular Matrix Biomaterials for Tissue Engineering: Lessons from the Common Sea Urchin Tissue. Int J Mol Sci. 18(5) (2017).
21. Kuo, Y. C. & Hseuh, C. H. Neuronal production from induced pluripotent stem cells in self-assembled collagen-hyaluronic acid- alginate microgel scaffolds with grafted GRGDSP/Ln5-P4. Mater Sci Eng C Mater Biol Appl. 76, 760–774 (2017).
22. Sensharma, P., Madhumathi, G., Jayant, R. D. & Jaiswal, A. K. Biomaterials and cells for neural tissue engineering: Current choices. Mater Sci Eng C Mater Biol Appl. 77, 1302–1315 (2017).
23. Zweckberger K., Ahuja C. S., Liu Y., Wang J., Fehlings M. G. Self-assembling peptides optimize the post-traumatic milieu and synergistically enhance the effects of neural stem cell therapy after cervical spinal cord injury. Acta Biomater. 51742–70661 (2016).
24. Wu, J., Xie, L., Lin, W. Y. & Chen, Q. Biomimetic nanofibrous scaffolds for neural tissue engineering and drug development. Drug Discov Today. 13539–6446(17), 30313–9 (2017).
25. An, B. et al. Physical and biological regulation of neuron regenerative growth and network formation on recombinant dragline silks. Biomaterials. 48, 137–46 (2015).
26. Bogush, V. G. et al. A novel model system for design of biomaterials based on recombinant analogs of spider silk proteins. J Neuroimmune Pharmacol. 4, 1729 (2009).
27. Moiseenovich, M. M. et al. Tissue regeneration in vivo within recombinant spidroin 1 scaffolds. Biomaterials. 33(15), 3887–98 (2012).
28. Takeuchi, M. et al. Human platelet-rich plasma promotes axon growth in brain-spinal cord coculture. Neuroreport. 23(12), 712–6 (2012).
29. Zheng, C. et al. Effect of platelet-rich plasma (PRP) concentration on proliferation, neurotrophic function and migration of Schwann cells in Tissue Eng Regen Med. 10(5), 428–36 (2016).
30. Sánchez, M. et al. Platelet-rich plasma, a source of autologous growth factors and biomimetic scaffold for peripheral nerve regeneration. Expert Opin Biol Ther. 17(2), 197–212 (2017).
31. Hayashi, C. Y., Shapley, N. H. & Lewis, R. V. Hypotheses that correlate the sequence, structure, and mechanical properties of spider silk proteins. Int J Biol Macromol. 24, 271–5 (1999).
32. Thuber, A. E., Omenetto, F. G. & Kaplan, D. L. In vivo bioresponses to silk proteins. Biomaterials. 71, 145–157 (2015).
33. Gu, Y. et al. Chitosan/silk fibroin-based, Schwann cell-derived extracellular matrix-modified scaffolds for bridging rat sciatic nerve gaps. Biomaterials. 35(7), 2253–63 (2014).
34. Qu, J. et al. Electrospun silk fibroin nanofibers in different diameters support neurite outgrowth and promote astrocyte migration. J Biomed Mater Res Part A 101A, 2667–2678 (2013).
35. Lewicka, M., Hermanson, O. & Rising, A. U. Recombinant spider silk matrices for neural stem cell cultures. Biomaterials. 33(31), 7712–7 (2012).
36. Costanzo, C. & Pehrson, J. R. Histone macroH2A1 is concentrated in the inactive X chromosome of female mammals. Nature. 393(6685), 599–601 (1998).
37. Wang, Y. L., Gu, X. M., Kong, Y., Feng, Q. L. & Yang, Y. M. Electrospun and woven silk fibroin/poly(lactic-co-glycolic acid) nerve guidance conduits for repairing peripheral nerve injury. Neural Regen Res 10(10), 1635–1642 (2015).
38. Ozaki, M. et al. Evaluation of the immunogenicity of human iPSC cell-derived neural stem/progenitor cells in vitro. Stem Cell Res. 19, 128–138 (2017).
39. Gutnik, M. I., Weidner, N., Müller, R. & Blesch, A. Cell-seeded alginate hydrogel scaffolds promote directed linear axonal regeneration in the injured rat spinal cord. Acta Biomater. 27, 140–150 (2015).
40. Madigan, N. et al. Comparison of cellular architecture, axonal growth, and blood vessel formation through cell-loaded polymer scaffolds in the transected rat spinal cord. Tissue Eng Part A. 20(21–22), 2985–97 (2014).
41. Sidoruk, K. V. et al. Fermentation optimization of a Saccharomyces cerevisiae strain producing 1F9 recombinant spidroin. Appl. Biochem. and Microbiol. 54(7), 766–773 (2015).
42. Yuan, Y. & Lee, T. R. Contact Angle and Wetting Properties. Surface Science. Techniques 51, 3–34 (2013).
43. Ayres, C. et al. Modulation of anisotropy in electrospun tissue-engineering scaffolds: Analysis of fiber alignment by the fast Fourier transform. Biomaterials. 27(32), 5524–5533 (2006).
44. Jahne B. Digital Image Processing, 6th ed. Berlin-Heidelberg: Springer. Ch. 19 (2005).

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Author Contributions
V.P.B. and J.-E.A. planned the experiments, and wrote the paper; V.P.B. performed all immunostaining procedures, analyzed the results, prepared Figs 1, 2, 4, 6–9; V.G.B. and K.V.S. made rSS-PCL scaffold for in vitro and in vivo experiments; V.A.K. prepared NPCs for all experiments; N.N.S. prepared PRP for two-component matrix; E.M.S. and V.A.R. made differentiation assay of NPCs and prepared two-component matrix with cells; M.A.K. made flow cytometry. V.P.B., V.A.K., N.N.S. and M.A.K. conducted in vivo experiment; S.L.K. and P.S.T. performed AFM and biomechanics research and prepared Fig. 3; K.B.Y. made an analysis of anisotropy and prepared Fig. 5; A.V.T., J.-E.A., and A.V.A. coordinated the whole project, discussed the data and contributed to the writing of the manuscript. All authors reviewed the manuscript.

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