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Additional Information
Biohybrids of scaffolding hyaluronic acid biomaterials plus adipose stem cells home local neural stem and endothelial cells: Implications for reconstruction of brain lesions after stroke

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Abstract: Endogenous neurogenesis in stroke is insufficient to replace the lost brain tissue, largely due to the lack of a proper biological structure to let new cells dwell in the damaged area. We hypothesized that scaffolds made of hyaluronic acid (HA) biomaterials (BM) could provide a suitable environment to home not only new neurons, but also vessels, glia and neurofila- ments. Further, the addition of exogenous cells, such as adipose stem cells (ASC) could increase this effect. Athymic mice were randomly assigned to a one of four group: stroke alone, stroke and implantation of BM, stroke and implantation of BM with ASC, and sham operated animals. Stroke model consisted of middle cerebral artery thrombosis with FeCl3. After 30 days, ani- mals underwent magnetic resonance imaging (MRI) and were sacrificed. Proliferation and neurogenesis increased at the subventricular zone ipsilateral to the ventricle and neuroblasts, glial, and endothelial cells forming capillaries were seen inside the BM. Those effects increased when ASC were added, while there was less inflammatory reaction. Three-dimensional scaffolds made of HA are able to home newly formed neurons, glia, and endothelial cells permitting the growth neurofilaments inside them. The addition of ASC increase these effects and decrease the inflammatory reaction to the implant. © 2018 Wiley Periodicals, Inc. J Biomed Mater Res B Part B: 000: 000–000, 2018.

Key Words: biomaterials, hyaluronic acid, cell therapy, adipose stem cells, stroke

INTRODUCTION
In spite of the extensive pharmacological and neurorehabil-itation efforts made to recover the neurological deficits, stroke is still the main cause of disability in the occidental world. Ischemic stroke produces an area of necrosis (core zone) surrounded by a partially damaged zone (area of penumbra).1 Most of the therapeutic efforts up to now have been directed to reduce the extent and to improve the functional recovery of the penumbra zone.2 However, there is not a reliable way to repair the lost tissue.3

After an experimental stroke, there is an increase of the neurogenesis at the subventricular zone (SVZ), a main adult
neurogenic niche, especially at the ipsilateral hemisphere, and new neurons and supporting cells migrate massively to the area of infarction. However, most of these cells disappear from the lesion by the 5th week. We hypothesize that the main reason for this is the lack of a proper substrate where the cells can survive, reconnect, and recover their function due to the generation of an adverse environment after the stroke and/or to the absence of a material scaffold upon which new neurons and supporting structures such as glia and vessels can grow and organize.

The use of three-dimensional (3D) biomaterials (BM) could provide a proper scaffold for these newly formed cells, permitting the organization of neural tissue with neurons, glial cells and vessels, guiding the neurite prolongations and providing a more appropriate environment for neural regeneration. Among the multiple available bio-

compatible materials, we chose hyaluronic acid (HA) because this is an already used BM in other clinical applications and it is simple to make and manipulate.

Furthermore, mesenchymal stem cells have been shown to home endogenous stem cells close to the infarcted area, as well as to increase neovascularization at the same site and proliferation at the SVZ, and also to diminish the inflammatory reaction at the lesion site. We hypothesized that the addition of mesenchymal stem cells to the BM would increase their capacity to home endogenous stem cells, and possibly to facilitate their entry into the BM by inhibiting the scar reaction around the scaffold. Among the mesenchymal cells available, we have chosen adipose stem cells (ASCs) because of practical reasons: they are easily handled, they are widely available from lipoaspirate donation for clinical application in stroke.

The objectives of this study were to determine if 3D scaffolds of hyaluronic acid (HA) implanted at the penumbra zone of a vascular stroke lesion in rats were able to home local neural and endothelial cells and neurites, and also to compare these homing capacities between BM co-grafted with adult human ASC and BM without them, as well as to study the host’s reaction to the implant in both cases, both in terms of glial scar formation and neurogenetic production. Also, if this represents a viable and safe method in order to transfer this to the clinical practice.

MATERIALS AND METHODS

Preparation of HA porous scaffolds

HA porous scaffolds were made as described Rodríguez-Pérez et al. Briefly, HA sodium salt from Streptococcus equi (1.5-1.8 M Da; Sigma-Aldrich) was dissolved overnight to a 5% (w/v) in 0.2 M sodium hydroxide (NaOH, Scharlab). Then, divinyl sulfone (DVS, Sigma-Aldrich) was added as a crosslinker in a 9:10 DVS:HA monomeric units molar ratio. Once homogenized for 10 s more, the solution was vacuum-injected into molds made from sintered beads of poly (ethyl methacrylate; PEMA, Elvacite® 2043, Lucite International, Inc.). These beads had a diameter of between 180 and 250 μm. The crosslinking reaction of HA with DVS was left to complete for an hour and a half.

After that, PEMA beads template was removed by leaching reflux in a Soxhlet extraction apparatus using boiling ethanol as solvent for 48 h. Afterwards, HA scaffolds were immersed in distilled water for 3 h, frozen at −20°C for another 2 h and then lyophilized. All scaffolds were cut into cylinders with 0.8 mm diameter and 8 mm of length. Other tests were carried out to warrant the absence of pyrogens or endotoxins, to guarantee biomaterial’s innocuity.

Analysis of 3D structure of biomaterials

The HA scaffolds, previously sterilized by chemical means, were analyzed by scanning electron microscopy to analyze the 3D configuration of the scaffold, ensuring that pores are between 300 and 150 μm in diameter were present, so that ASC can easily enter the BM. For this purpose, they were processed in a critical dot dryer, followed by gold-palladium coating to provide electroconductivity to the sample and analyzed in a scanning electron microscope JEOL JSM 5410 with a voltage of 10 kV at different magnitudes (Figure 1).

Adipose stem cells

Adipose tissue samples were obtained from donors during routine abdominoplasty following informed patient consent and according to the guidelines set by the corresponding

FIGURE 1 Scanning electronic micrographs showing porous scaffolds made of hyaluronic acid at different magnifications (A–D). A–C: shows images of the biomaterial and the characteristics of the scaffolding and D–E: images of ASC cells colonizing the scaffold. Scale Bar A: 500 μm, B: 300 μm, C: 150 μm, D: 80 μm, and E: 30 μm.
Ethics Committee on Biomedical Research. The adipose tissue was transported in sterile bottles and immediately processed to obtain the ASC and process under GMP conditions (Histocell, S.L., Bilbao, Spain). Phenotype characteristics, clonic capacity and stem cell differentiation biomarkers as described by Gomez-Pinedo et al.23

The ASC were maintained until phase 7 in culture on the material to perform studies of telomerase and C-myc expression, to guarantee the stability of the cultured cells on the biomaterial. Before use, we proceeded to carry out the immunophenotype of the ASC cells to be considered mesenchymal stem cells and useful for our purpose.

Scaffold preloading
ASC were used in the fourth cell phase. Cells were resuspended in culture medium at a concentration of 5x10^4/4 μL. This concentration was inoculated inside the HA biomaterial (dimensions for the preclinical study of 1 x 1 mm) with the help of a Hamilton® microsyringe. The biomaterials inoculated with ASC were maintained in 24-well plates with the culture medium composed of DMEM-Glutamax + Antibiotic / 10% FBS and incubated at 37°C/5% CO₂ for 72 h, for subsequent implant in the model of athymic nude mice [Figure 1(D–E)].

FeCl₃ thrombosis model
Nude-Foxn1<sup>−/−</sup>/nu (Envigo-Harlan Laboratories) adult male mice were used for the experiments. At 7 weeks of age were housed in a temperature-controlled room with access to food and water ad libitum. They were kept in individual cages and under standard conditions in the animal facilities of the Hospital Clínico San Carlos. All procedures were carried out under animal welfare regulations in accordance with EC Council Directive November 24, 1986 and approved by the local ethics committee.

Mice were anesthetized with 80–100 mg/kg ketamine and 10 mg/kg xylazine. A small craniotomy was performed to access the middle cerebral artery. For the formation of the ischemia model, a FeCl₃ solution (20%) was applied to the surface of the artery for 1 min. In the control group (sham) 0.9% of saline solution was used.15,16

Experimental and control groups
Four groups were defined: Stroke (mice subjected to the stroke model), Stroke+BM (mice subjected to the stroke model to whom HA biomaterials alone were implanted at the core zone), Stroke+BM+ASC (mice subjected to the stroke model to whom HA biomaterials co-grafted with ASC were implanted at the core zone) and sham (animals undergoing the operation for the stroke model except the application of the thrombotic FeCl₃ solution). Six animals were assigned randomly to each group.

Seven days after the ischemic procedure, animals belonging to groups Stroke+BM and Stroke+BM+ASC were again anesthetized and the wound reopened. A burr hole was performed at the cranial vault just medial to the temporal crest.

![Figure 2](image.png)

**FIGURE 2.** Magnetic resonance imaging demonstrating the infarcted area in the control group (only stroke) and the implanted groups. In the latter, the biomaterial is fully integrated, and it is associated with less cavitation.
FIGURE 3. Analysis of the glial scar. Representative immunohistochemical image showing glial fibrillary acidic protein (GFAP), (A) stroke; (B) stroke + BM; (C) stroke + BM + ASC; (D) The graph shows that in the group of animals with stroke, in which ASC were administered in biomaterials, they showed less expression of the GFAP, in the other two groups (stroke and stroke + BM) the expression of GFAP was significantly higher (**p < 0.05 and *p < 0.01) Scale Bar: 100 μm.

at the same coronal plane at which the occluded MCA branch was located. A cannula was inserted to implant the biomaterials (with or without ASC) at a depth of 13 mm from the dural plane. (Supporting Information Figure 2).

Magnetic resonance imaging
The day of the sacrifice, the mice were anesthetized with isoflurane 1.5–2% to obtain a brain MRI. All the MRI experiments were performed on a BIOSPEC BMT 47/40 (Bruker, Ettlingen, Germany) spectrometer operating a 4.7 T, equipped with a 11.2 cm actively shielded gradient system, capable of 200 mT/m gradient strength and 80 μs of rise time. During the MRI procedure, the animals were kept at 37 °C and a MR compatible respiration sensor was used to control the animals. The MRI experiments consisted of three-dimensional T2WI, a series of DWI to calculate the ADC maps and a series of PWI to calculate CBF maps. A 7-cm birdcage radiofrequency coil was used for transmission and reception. Image analysis was performed using ParaVision 3.0.1 (Bruker, Ettlingen, Germany). This calculation was carried out using MATLAB 7.3 (MathWorks, Inc., Natick, MA).

BrdU administration. The BrdU is an analog of the thymidine that the cell incorporates in the cellular cycle when entering into cellular division. Two hours before the sacrifice of the animals an intraperitoneal pulse of BrdU (60 mg/kg, Sigma-Aldrich) dissolved in physiological saline was administrated to evaluate the proliferative activity of the endogenous stem cells in the SVZ, as well as implanted ASC cells.23

At 37 days after stroke surgery, mice in each group were anesthetized with a dose of pentobarbital (60 mg/kg) and fentanyl (0.3 mg/kg) for subsequent intracardiac infusion with 0.9% solution Saline followed by 4% buffered paraformaldehyde (0.1 M phosphate buffer). After perfusion, the brain was removed and washed with 0.1 M PB and cryoprotected by immersion in 30% sucrose. The cuts were performed in a cryostat (model Micron 1800) at 50 microns thick containing the infarct area and the biomaterial implant area.

Histochemistry and immunofluorescence analysis
The sections were washed with PBS, permeabilized with 0.1% Triton X-100 and blocked with 10% normal goat serum. The following primary antibodies were then applied overnight at 4°C: anti-HuNu (Millipore, human cells); GFAP (Dako Cytomation, astrocytes); Anti-CD31 (Abcam, newly formed endothelium); Anti-NeuN (Millipore, neurons); Anti- DCX (Santacruz, neuroblasts); Anti-BrdU (Millipore, thymidine analogue); Anti-NF 160 and anti-NF 200 (Millipore, neurofilaments). Subsequently the samples were washed in PBS three times for subsequent incubation in Alexa 488, 555, or 647 (1:500, Invitrogen) conjugated secondary
antibodies, DAPI-contracted and mounted with the Fluorsave reagent (Calbiochem). Immunofluorescence images were obtained with the Olympus AF2000 confocal microscope.

The quantitative study both at the SVZ and at the areas around the biomaterial consisted in the analysis of 10 different fields for each of the antibodies studied (BrdU, DCX, and GFAP) and the result was the average of 10 determinations. Within the BM, a volumetric analysis was performed using confocal microscopy of 12 optical segments 800 nm thick.

The NF were counted in four optical planes by confocal microscopy in z projection. In cases where the unit of measurement was the observed mark quantity per field (Optical Density, DO), the software program Image J, v1.46r of the National Institutes of Health was used for its calculation.

Volume analyzed 0.69 mm³.

RESULTS

After 72 h of incubation, cells were easily inserted into the scaffolds, and they were adhered to the walls of the same in a homogeneous way (Figure 1). Karyotype and phenotype did not show any alteration.

The application of FeCl₃ induced a limited and constant brain infarction located at the left parietotemporal cortex (Supporting Information Figure 1). Neurological deficits appeared 4 h after the surgery, as forelimb flexion, resistance to lateral push and circling behavior; but all animals returned to a normal neurological examination 12 h later.

Animals did not show any further neurological alteration after the implant of the biomaterials, without inducing or inciting fever or implant rejection, suggesting the absence of pyrogenic elements. Magnetic resonance imaging showed that the implanted biomaterials were fully integrated into the host tissue and reduced the cavitation provoked by stroke in control animals. The reduction was more evident when ASC were added to the biomaterials (Figure 2).

After stroke, there was an astrocytic reaction around the infarcted area. The astrocytic reaction to the biomaterial was not intense (similar to the one produced with stroke alone), but it was much less to the biomaterials precharged with ASC, as we show in Figure 3 (Sham: 0.341 0.029 T Stroke: 1.780 0.106; Stroke+BM: 1.681 0.066; Stroke+BM+ASC: 0.935 0.086).

Neurofilaments were scarcely seen at the infarcted area in control animals, but they were found within the biomaterials. After 30 days of scaffold implantation (with or without ASC), more neurofilaments, both immature (160 kD) and mature (200 kD), were encountered inside the biomaterial

**TABLE I. Counting of Immature (NF 160 kD) and Mature (NF 200 kD) Neurofilaments Found in the Ischemic Zone in Stroke Controls, Stroke Animals After Placement of Biomaterales, and After Placement of Biomaterials with ASC (Units: Neurofilaments Per mm³)**

<table>
<thead>
<tr>
<th>Group</th>
<th>NF 160 kD</th>
<th>NF 200 kD</th>
<th>NF 160 kD</th>
<th>NF 200 kD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stroke</td>
<td>9.3 ± 6.7</td>
<td>2.1 ± 1.9</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Stroke+ASCs</td>
<td>–</td>
<td>–</td>
<td>36.2 ± 17.9</td>
<td>12.6 ± 9.4</td>
</tr>
<tr>
<td>Stroke+BM+ASC</td>
<td>–</td>
<td>–</td>
<td>89.7 ± 13.2</td>
<td>20.4 ± 3.7</td>
</tr>
</tbody>
</table>

Volume analyzed 0.69 mm³.

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**FIGURE 4.** Immunohistochemical image showing the neurofilaments found inside the biomaterial+ASC 30 days after implantation. (HA: hyaluronic acid, DAPI: nucleus, NF160: 160 kD neurofilaments, NF200: 200 kD neurofilaments. Scale Bar: 100 μm B: Detail from image A. Scale Bar: 10 μm.
than in the periphery of the lesion or in the controls (Table I).

More neurofilaments were seen in the biomaterial plus ASC than in the one without them. Most of the neurofilaments were immature (160 kD) (Stroke: 9.3 \pm 6.7; Stroke+BM: 36.2 \pm 17.9; Stroke+BM+ASC: 89.7 \pm 13.2; Table I and Figure 3), but mature neurofilaments (200 kD) were also seen (Stroke: 2.1 \pm 1.9; Stroke+BM: 12.6 \pm 9.4; Stroke+BM+ASC: 20.4 \pm 7.7, data expressed in NF into biomaterial; Table I and Figure 4).

CD31 positive cells were seen inside the implants (Figure 5), but not in the infarcted area in control animals. Some of them were organized in tubule-like structures, resembling vessels. Also, more CD31 positive cells were seen inside the biomaterials pre-charged with ASC.

There was a proliferative reaction at the SVZ after the stroke, in the number of BrdU (Sham: 122 \pm 8; Stroke: 174 \pm 16; Stroke+BM: 241 \pm 11; Stroke+BM+ASC: 305 \pm 26) DCX (Sham: 0.51 \pm 0.07; Stroke: 0.94 \pm 0.08; Stroke+BM: 1.52 \pm 0.09; Stroke+BM+ASC: 1.83 \pm 0.12) positive cells. The implantation of biomaterials increased this reaction. The preloaded cells (ASC) increased even further this effect (Figure 6).

Neuroblast cells showing DCX were not seen inside the infarcted areas in control animals, although some of them were seen in the vicinity of the core zone. More cells were seen in the vicinity of the scaffold alone and even more around the biomaterials plus ASC. Inside the biomaterials we found some newly formed neuroblasts, and this number increased when ASC were co-grafted (Stroke: 0.032 \pm 0.0052; Stroke+BM: 0.061 \pm 0.0096; Stroke+BM+ASC: 0.096 \pm 0.0169; Figure 7).

Neither HuNu positive cells, neither teratomas or any other neoplasic formation were found inside the biomaterials or in the surrounding brain tissue after 30 days of implantation of human ASC.

DISCUSSION

Here, we show that the implant of biomaterials made of HA at the core of a brain stroke lesion permits the migration of endogenous new neurons, glia, and endothelial cells, and the proliferation of neurofilaments and vessel-like structures.
into the biomaterials. Furthermore, the implants increase the proliferation of neuroblasts at the SVZ. The addition of ASCs to the biomaterials increase the proliferative response and the homing effect of biomaterials, plus decreasing the astrocytic reaction to stroke, implantation surgery and biomat-erials. It also increases the number of neurofilaments found inside the scaffolds and the number of newly formed vessels inside the biomaterials. The biomaterials are easily inte-grated into the core zone, and this integration was even better when ASC were associated (Supporting Information Figure 2).

ASC has been used to diverse inflammatory pathologies by many authors; being recovery objectives not only in the central nervous system,52,56 ASC have shown to increase SVZ neurogenesis,27–31 and have antiinflammatory and cytotoxic properties when administered in stroke models,22,33 and to improve function in animal models, both when administered locally or even through the systemic route.21,23,34,35 In addition, it has been shown their neuroprotector effect that intervenes directly in the apoptosis mechanism as describe Wei et al.22,36; and their secretor effect as show Cunningham et al.36 However, these approaches do not seem to replace the totally brain tissue lost.37,38

Conversely, the use of scaffolds leads to true regeneration or re-formation of new cerebral tissue, not only reducing the bulk cavitation provoked by stroke, but with the presence of new neurons, neurites and supporting cells and vessels, which might functionally integrate with the sur-rounding brain, as shown Fuhrmann et al., who propose the strategy of favoring the microenvironment.9,39–41

Scaffolds made of HA showed compatibility with the brain, as previously noted.13,14 This confirms the innocuous nature of this biomaterial, which can be designed to fit the particularities of the host tissue, therefore favoring the migration of healthy cells adjoining the lesion zone in several models of injury.20,42,43 Recently, biomaterials is a growing sector, with very different compositions, such as plasma, PCL, or hydrogels with adhered molecules. All of them share a structural function to restructure the damage tissue.44–48

The combination of HA BM+ASC has a synergistic effect, promotes the proliferative activity in the SVZ (increase BrdU and DCX positive cells), enhancing the attraction of endogenous cells, therefore, promoting an appropriate niche to recovery.49,51 The increase in proliferation of the SVZ analyzed by BrdU agrees with the increase of DCX cells in the group with HA plus ASC. In the group where ASC was not administered, a lower amount of DCX and a greater amount of GFAP is observed. So the administration of ASC and its immunomodulatory or anti-inflammatory effects possibly influence the proliferation and differentiation of GFAP and DCX cells.23,49–51

Furthermore, ASC increases the biocompatibility and homing ability of HA scaffolds.52–54 This is done probably by the antiinflammatory and cytotoxic properties of ASC, but not by becoming part of the final implanted structure, since no antigens marking human nuclei were found 30 days after implantation. Also, ASC showed a good integration with HA when the scaffolds were preloaded with these cells, as shown earlier. The combination of the two is safe, as no abnormal cell proliferation of ASC was seen neither in culture nor after implantation. Simultaneously to enhance an appropriate niche and the damage tissue is reduced; the plasticity mechanism is favored.55–57

We used immunocompromised animals to explore safety in terms of possible neoplastic proliferation of ASC after implantation.23,48 However, these cells have not shown rejection issues when used in immunocompetent rodent stroke models.36,37,58

In control animals receiving stroke, newly formed neurons (DCX) coming from the ipsilateral SVZ were seen in the vicinity of the core zone, but were not seen inside the infarcted areas. This confirms the observations by others showing that stroke induces per se a repair reaction, but these cells are unable to stay and organize within the core zone.5,59

The stroke model we used induced an infarction pattern which is more constant than other models.60 However, as this model produces few neurological side effects in the animals, we could not observe if this treatment strategy may produce functional improvement.61–63 Further studies in different models should be done in rodents, or higher mammals should be tested to determine this.

In conclusion, in stroke, the use of HA and ASC is novel. The present results show how scaffolds of HA implanted at the core zone of a stroke lesion permit the proliferation and migration of new neurons, neurites, glia, and vessels, and the addition of ASC increases this proliferation and homing effect (Supporting Information Figure 2). This may permit formation of new brain tissue after stroke.

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CONFLICT OF INTEREST
The authors declare no other conflict of interest.

REFERENCES


