Cell therapy in experimental models of Parkinson’s disease replaces the lost dopamine neurons (DAN), but we still need improved methods to guide dopaminergic axons (DAx) of grafted neurons to make proper connections. The protein Semaphorin 3C (Sema3C) attracts DAN axons and enhances their growth. In this work, we show that the hydrogel PuraMatrix, a self-assembling peptide-based matrix, incorporates Sema3C and releases it steadily during 4 weeks. We also tested if hydrogel-delivered Sema3C attracts DAx using a system of rat midbrain explants embedded in collagen gels. We show that Sema3C released by this hydrogel attracts DAx, in a similar way to pretectum, which is known to attract growing DAN axons. We assessed the effect of Sema3C on the growth of DAx using microfluidic devices. DAN from rat midbrain or those differentiated from human embryonic stem cells showed enhanced axonal extension when exposed to hydrogel-released Sema3C, similar to soluble Sema3C. Notably, DAN of human origin express the cognate Sema3C receptors, Neuropilin1 and Neuropilin2. These results show that PuraMatrix is able to incorporate and release Sema3C, and such delivery guides and promotes the axonal growth of DAN. This biocompatible hydrogel might be useful as a Sema3C carrier for in vivo studies in parkinsonian animal models.

Introduction

Axons grow with spatiotemporal specificity through the response to different cues during embryonic stages. Such growth is directed by the expression of ligands and their cognate receptors. The large family of Semaphorins (Sema) includes both soluble and membrane-bound proteins that mediate axonal guidance through Rho-related G proteins. Class 3 Sema (Sema 3) are guide molecules well conserved in several species. Sema3A was initially reported as Collapsin, as exposure of sensory neuron explants of the chick dorsal root ganglia to this protein produced growth cone collapse. Although initially characterized as inhibitors of axonal growth, it was later discovered that Sema3 can induce either repulsive or attractive effects depending on the cellular context. Sema3A on cortical axons or Sema3E on retinal ganglion cells produced repulsive effects. On the contrary, Semaphorin 3C (Sema3C) secreted by a transient neuronal population exerted an attractive influence on crossing callosal axons during development. Exposure of developing mouse hippocampal subicular neurons to Sema3E promotes and attracts the axonal growth, an effect mediated by Neuropilin (NRP) 1.

Several Sema proteins influence the growth of mesencephalic dopaminergic axons (DAx) during development: Sema3A, Sema3C, Sema3F, Slit-2, and Netrin-1 are all implicated in the guidance of DAx to reach the striatal area. Growing DAx respond differentially to Sema3 signaling: Sema3F is repulsive for DAx emanating from mesencephalic explants embedded in collagen gels. In contrast, Sema3C and Sema3A increase the axonal growth of DAx, whereas Sema3C also attracts DAx of mesencephalic
neurons and those derived in vitro from mouse embryonic stem (ES) cells. Neutralizing antibodies for the Sema3C receptors NRP1 and NRP2 block Sema3C effects on rodent DAx.

The low ability of axon regeneration in the adult central nervous system (CNS) has been widely documented and is mainly due to the lack of signals present during development and to the expression of inhibitory molecules by reactive glial cells. These impediments become relevant when axonal regeneration in the CNS is needed to repair the damaged or diseased brain. In Parkinson’s disease (PD), DAx that form the nigrostriatal pathway are lost, secondary to the death of dopamine neurons (DAN) in the substantia nigra (SN). Grafting of DAN in the striatum, the region originally innervated by DAx, causes behavioral recovery in adult parkinsonian rats. However, this strategy applied to PD patients did not cause consistent recovery.

In contrast, when DAN are grafted in the SN, they are unable to project their axons to the striatum probably due to the adverse environment for axonal growth. Enhancing and directing DAx growth is an appealing idea to be tested when grafting neurons in PD models. In a rat model of PD, we previously demonstrated that transfected HEK-293 cells could release Sema3C along an artificial trajectory between the SN and the striatum, promoting the DAx growth of neurons grafted in the SN, resulting in behavioral improvement.

The use of biomaterial scaffolds might constitute an ideal strategy to replace transplanted cells for delivery of Sema3C to the brain. Hydrogels are polymers that incorporate large quantities of water in their structure and gels in response to several physiological conditions, such as temperature or osmolarity. Recent studies have shown the usefulness of hydrogels as drug carries due to their ability for in vivo or in vitro release (reviewed in Carballo-Molina and Velasco37). For example, neurotrophic factors such as glial cell-derived neurotrophic factor (GDNF), brain-derived neurotrophic factor (BDNF), or ciliary neurotrophic factor can be incorporated to the hydrogel and then released.

Hydrogels are also a good substrate for neural cells, allowing their survival, growth and proliferation (reviewed in Carballo-Molina and Velasco). Neural stem cells can proliferate and differentiate when embedded in hydrogels in vitro and in vivo. Some researchers have investigated the use of hydrogels as cell carriers of rat neurons and human neural progenitor cells. Several groups have reported that the implantation of hydrogels to rodent and primate brains does not cause tissue rejection and promotes axonal growth (reviewed in Carballo-Molina and Velasco). The hydrogel PuraMatrix is a synthetic molecule constituted by self-assembling peptides with the sequence arginine-alanine-aspartate-alanine repeated 16 times (RADA16). This hydrogel promoted recovery in a hamster model of blindness when implanted in the brain and helped recover from cortical lesions, mainly by allowing axonal growth and the formation of synapses in vitro. Furthermore, PuraMatrix can incorporate different proteins into its structure and release them in vitro. In this study, we evaluated if PuraMatrix can release Sema3C and guide/promote DAx growth. We show that PuraMatrix-embedded Sema3C is stably released and that such delivery causes attraction of DAx from midbrain DAN.

Quantification of the growth of DAx of rat and human origin showed that Sema3C released by this hydrogel caused significant larger axons, compared to controls.

Materials and Methods

Conjugation of proteins to hydrogel and release assays

We used the RADA16 BD PuraMatrix hydrogel (1% w/v solution) based on self-assembling peptides. The hydrogel was diluted with sterile water to obtain a 0.15% w/v final concentration as recommended by the manufacturer. The hydrogel was combined with recombinant human Sema3C peptide (Abcam) and incubated at 37°C in a sterile saline solution for 28 days; an aliquot of the solution was recovered and the same volume was added every day. The experiment was repeated five times for each concentration. Sema3C was mixed with this hydrogel solution previous to the gelling process, which was triggered by adding saline solution (0.9% NaCl). To measure the release of Sema3C conjugated at 5 and 10 μM/mL with hydrogel, an ELISA was performed, according to the manufacturer’s protocol (Cloud-Clone Corp).

Animals

Adult Wistar rats and embryos at 14 days of gestational age (E14) were used. The animals were maintained and handled in accordance with the local animal care and use committee and complied local (NOM-062-ZOO-1999) and international guidelines (Animal Welfare Assurance A5281-01). Rats were housed with a 12-h light–12-h dark cycle at room temperature (RT, 22°C ± 2°C). Water and food were freely available. Animals were euthanized. All efforts were made to minimize the number of animals used and to avoid pain or discomfort.

Collagen extraction from rat tails

The collagen extraction procedure was made as reported previously. Four adult frozen rat tails were rinsed with 70% ethanol. Tendons were extracted using sterile tweezers and rinsed with 70% ethanol. We obtained 2 g of tendons that were cut into 0.5 cm fragments. The dry tendons were transferred to 60 mL of ice-cold 0.1% acetic acid solution. Tendons were stirred for 20 h at 4°C. The solution was diluted with 80 mL of sterile water and centrifuged at 9000 rpm for 30 min at 4°C. This solution was used to prepare collagen gels.

Assays of explants in collagen gels to measure axonal attraction

Ventral midbrain (Mb), isthmic region (Ist), and pretectum (PT) explants were obtained from E14 embryos as described and were maintained in Krebs solution. Ventral Mb explants were embedded in a collagen matrix by mixing 300 μL of N2 medium (DMEM/F12 1:1, supplemented with 25 μL/L of human insulin, 30 nM sodium selenite, 100 μM putrescine, 20 nM progesterone, and 100 mg/L of apotransferrin) with 100 μL of collagen solution, 10 μL of 1.537 M NaCl solution (final concentration of 36.6 mM), and 10 μL of 0.89 M of NaHCO3 (final concentration of 21.2 mM); this mixture was maintained on ice before the gelling process. The Mb explants were embedded together with PT (n = 23) or with Ist (n = 17) explants, on four-well dishes (Nunc), and let to polymerize at 37°C for 1 h. Special care was taken to keep a distance of 500–
1000 μm between explants before adding another layer of collagen and let to polymerize for an additional hour. Finally, 700 μL of N2 medium was added.

For hydrogel experiments, a drop of PuraMatrix coupled to either Sema3C (n = 21) or gelatin from pig skin coupled to Oregon Green 488 (Molecular Probes) as a control protein (Cp), since this protein has a similar weight and charge to Sema3C (n = 11), both at 200 μg/mL, was jellified for 1 h by adding medium at RT. A layer of collagen was added and immediately the Mb explants were introduced. After 1 h at 37°C, an additional layer of collagen was added and let to polymerize for 1 h, and finally, 700 μL of N2 medium was added. Explants were cultured for 5 days without medium change.

To determine the attraction/repulsion effects on Mb explants, the TH+ axon length was measured with ImageJ software (National Institutes of Health) using NeuronJ plugin.51 The Mb explants were divided into two areas, proximal (P) or distal (D) (as indicated in the scheme of Fig. 2A). In cases where explants extended many axons, only the 10 largest were considered; when axons were fewer than 10, all axons were measured. The average axonal length for D and P sides was obtained to calculate the P/D ratio. When the value was greater than 1.0, it was considered an attractive effect; values smaller than 1.0 were considered repulsive.52 When the Mb explant does not present axonal growth, a zero value was registered. One methodological problem we encountered in the hydrogel groups was that the gel often fragmented when collagen was placed on top. We discarded experiments where many pieces of hydrogel were present.

**Dopaminergic differentiation of H9 human ES cells**

The ES cell line H9 was used after lentiviral transduction for GFP expression as reported.53 Dopaminergic neuron differentiation was made with the dual-SMAD inhibition and floor plate induction.54 Briefly, H9 cells were plated on Matrigel (BD), grown for 11 days in a knockout serum replacement medium (KSR) containing knockout DMEM, 15% serum replacement, 2 mM GlutaMAX, 1X nonessential amino acids, and 10 μM β-mercaptoethanol. Cultures were exposed to LDN193189 (100 nM, Stemgent; days 0–11), SB431542 (10 μM, Tocris; days 0–5), the sonic agonists SAG (1 μM, Sigma; days 1–9) and Purmorphamine (2 μM, Stemgent; days 1–9), FGF8 (100 ng/mL, Preptech; days 1–7) and CHIR99021 (3 μM, Stemgent; days 3–13). The KSR medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal medium was gradually shifted to N2 medium starting on day 5. On day 11, the medium was changed to the Neurobasal

**Calcein staining, immunocytochemistry, and microscopy**

To visualize live rat cells by fluorescence, cultures were incubated with 1 μM calcein-AM (R&D Systems) for 15 min. This assay allowed us to clearly identify the growing axons inside the microgrooves of microfluidic devices, which in some cases were difficult to observe in phase-contrast microscopy. For immunodetection, cells in microfluidic devices were fixed with 4% paraformaldehyde for 20 min at RT or with methanol for 5 min at −20°C. Cells were permeabilized and blocked for 1 h at RT with 0.3% Triton X-100 and 10% normal goat serum in PBS. Cells were incubated overnight with the following primary antibodies in PBS plus 10% normal serum: mouse anti-β III Tubulin 1:1000 (Covance); rabbit polyclonal anti-TH 1:1000 (Pel-freeze). Secondary antibodies were incubated for 2 h at RT: Alexa-Fluor 488 goat anti-rabbit 1:1000 (Molecular Probes); Alexa 568 goat anti-mouse 1:1000 (Molecular Probes). For human cells expressing GFP, Alexa 488 was substituted by Alexa Fluor 647 goat anti-rabbit 1:500 (Jackson ImmunoResearch). In all experiments, negative controls that included the addition of only secondary antibodies did not show unspecific staining.

**Axonal growth chambers**

Polydimethylsiloxane microfluidic devices were purchased from Xona Microfluidics (Standard Neuron Device Cat.No: SND900) and were used according to the manufacturer’s protocol. Briefly, they were attached by pressure to a coverslip precoated with poly-L-ornithine (15 μg/mL)/laminin (1 μg/mL)/fibronectin (2 μg/mL). The ventral Mb region was dissected from E14 rat embryos in Krebs solution and dissociated with TrypLE Express (Gibco) to obtain DAN. Cells were plated (250,000 cells/5 μL) in the cell compartment (Fig. 3A). After 10 min, 100 μL of N2 medium was added. The other side of the chamber was filled with 200 μL of N2 medium alone (control) or with 20 μg/mL Sema3C. For hydrogel experiments, 100 μL of the medium was first added in one supply well in the opposite compartment where cells were plated (Fig. 3A), then 3.5 μL of the hydrogel containing 200 μg/mL of Sema3C or Cp and 100 μL of medium were added. The cultures were maintained at 37°C in humidified 5% CO2/95% air for 5 days. Each chamber contains a variable number of axons, although the cells were plated at the same initial density; the total numbers of axons measured were as follows: medium alone (n = 48), medium plus soluble Sema3C (n = 59), hydrogel with Cp (n = 90), and hydrogel coupled to Sema3C (n = 92). Axonal growth in the microgrooves of the chambers was measured by ImageJ software using NeuronJ plugin.51 We decided to use this software because it allows the measurement of axons inside the microgrooves, the axons entering the microgrooves, and also axons emerging from the channels. Human DAN were introduced to microfluidic devices at day 21 of differentiation and kept for 14 days.
Contrast, Ist and Hyd with Cp mostly induced repulsion. PT and Hyd with Sema3C mainly promote dopaminergic attraction; in contrast, Ist explant exposed to Hyd with Cp, which showed 64% of DAx exposed to hydrogel with Cp, which showed 64% of DAx toward Ist explant. In all cases, the top pictures are reconstructions of low magnification and the images below are enlargements of the boxed areas. (C) Mb explant cultured with PT. DAX grew toward the PT. (D) Mb explant exposed to Hyd with Sema3C. DAX are preferentially present toward the hydrogel. (E) Mb explant cocultured with Hyd Cp. The axons grow randomly. (F) DAX were classified as proximal, distal, or absence of growth and the proportion of attraction, repulsion, or no growth was calculated. PT and Hyd with Sema3C mainly promote dopaminergic attraction; in contrast, Ist and Hyd with Cp mostly induced repulsion. (G) P/D ratios were plotted for the different conditions. PT and Hyd Sema3C promoted attraction, whereas Ist and Hyd Cp conditions induced repulsion. No significant differences were found between PT and Hyd Sema3C. (H) The length of DAX regardless of the direction of growth did not change between all the analyzed conditions. Mean ± standard error of mean *p < 0.05; **p < 0.01; ***p < 0.001 (one-way ANOVA and Tukey’s multiple comparisons test). Scale bars: 1000 μm (reconstruction images) and 100 μm (magnification images). Color images available online at www.liebertpub.com/tea

Explants were processed following reported methods. Briefly, collagen gels were fixed with 4% paraformaldehyde for 2 h at RT. Explants were permeabilized and blocked for 3 h at RT with 0.3% Triton X-100 and 10% normal goat serum in PBS. Collagen gels were exposed to the same primary and secondary antibodies used in microfluidic devices. The times of incubation were extended: explants were exposed to primary antibodies for 2 days and secondary antibodies for 1 day.

Statistical analysis

Results

Hydrogel releases proteins to the medium

We conjugated recombinant human Sema3C to PuraMatrix hydrogel at two different concentrations (5 or 10 μg/mL) and measured the release of the protein. We collected the solution in contact with the hydrogel every week and quantified the amount of Sema3C released by ELISA. After 28 days, we detected constant levels of the protein for both concentrations (Fig. 1).

Sema3C released by hydrogel induces attraction of DAx

Collagen gels form a permissive three-dimensional environment for axonal growth and also allow the diffusion of molecules. To test if hydrogel-released Sema3C could influence the direction of DAX growth, we used a collagen gel culture system that included ventral Mb explants, which contain DAN, together with pretectum (PT) or isthmic region (Ist) explants. We also combined the Mb explants growing in collagen with the polymerized PuraMatrix hydrogel containing either Sema3C or a Cp. We used Mb-PT explants as a positive control because this embryonic region exerts axonal dopaminergic attraction, as a negative control, we combined Mb with Ist, which does not attract DAX. In parallel, we cocultured Mb explants with hydrogel coupled to either Cp or Sema3C (Fig. 2A). After 5 days, we immunostained for TH and β III Tubulin to visualize the direction of dopaminergic axonal growth. A proximal/distal attraction ratio was calculated as described in Methods (Fig. 2A).

We did not observe growing axons toward Ist in cocultured Mb-Ist explants (Fig. 2B). In contrast, we observed an attractive effect on DAX when Mb was cocultured with PT (Fig. 2C). We observed a similar attractive effect when Mb was exposed to the Sema3C-embedded hydrogel, where DAX reached mainly the P area, and the larger axons grew toward the hydrogel (Fig. 2D). DAX grew in random directions when exposed to hydrogels conjugated to Cp (Fig. 2E).

Seventy percent of axons in the Mb-PT cocultures showed axonal attraction and 30% repulsion. In contrast, for the Mb-Ist group, Mb DAX showed repulsion in 65% of cases, in 12% attraction, and in 23% no growth (Fig. 2F). Mb explants exposed to Sema3C-embedded hydrogel presented 90% of attraction and 10% of repulsion, in contrast to DAX exposed to hydrogel with Cp, which showed 64% of repulsion, 27% attraction, and 9% without growth (Fig. 2F).
We calculated the proximal/distal ratio to measure dopaminergic attraction in each condition (Fig. 2G). Mb-PT and Mb-hydrogel Sema3C presented proximal/distal ratio values above 1 (Fig. 2G), indicating that Sema3C released by hydrogel attracts DAx similarly to PT. Mb-Ist and Mb-hydrogel Cp cocultures had values below 1 (Fig. 2G). We also measured the length of TH+ axons regardless of their proximal or distal position and found that it was similar in all groups (Fig. 2H).

**Sema3C released by hydrogel promotes axonal growth of rat DAN**

To evaluate if Sema3C released by hydrogel could increase DAx length, we used commercial microfluidic devices designed to assess the axonal response to soluble molecules. These chambers contain two sides that are separated by microgrooves of 900 μm of length (Fig. 3A), since it has been established that only axons are able to grow this distance.61,62

One side of the chamber contains more culture medium than the other, creating a hydrostatic force that allows the one-direction movement of soluble molecules from the axonal to the soma compartment (Fig. 3A). As indicated by the scheme, we plated DAN in the soma side of the chamber and Sema3C was present in the axonal compartment. One key advantage of these devices is that axons follow a linear trajectory, in contrast to three-dimensional approaches such as collagen gels. We measured DAx of dissociated rat Mb DAN after 5 days of exposure to different treatments: (1) medium alone; (2) medium plus soluble Sema3C (20 μg/mL); (3) hydrogel

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**FIG. 3.** Axonal growth of embryonic rat dopamine neurons (DAN) is promoted by soluble Sema3C and Sema3C released by hydrogel in microfluidic devices. (A) Scheme of neurons cultured into the microfluidic device. (B) Immunostaining with β III Tubulin and TH antibodies shows that in the axonal compartment, there are more and larger DAx in Medium + Sema3C and Hydrogel + Sema3C conditions, compared to control medium and Hydrogel + Cp, respectively. In the soma compartment, the density of TH/β III Tubulin double-positive cells close to the zone of microgrooves is similar in all conditions. In the right panel, a low-magnification picture after live calcein staining is presented. In all cases, axons that entered the microgrooves were similar in number. The orange and blue boxes correspond to the areas in the soma and axonal compartments, respectively. (C) Quantification of total dopaminergic axonal length. Medium + Sema3C and Hydrogel + Sema3C conditions exert a similar effect on axonal growth, in contrast to control medium and Hydrogel + Cp, which had significantly lower levels. (D) In all conditions, the number of DAx that entered to the microgrooves was not statistically different. Mean ± standard error of mean **p<0.01; ****p<0.001 (one-way ANOVA and Tukey’s multiple comparisons test). Scale bars: 100 and 300 μm (low-magnification image of calcein staining). Color images available online at www.liebertpub.com/tea
coupled to Cp (200 μg/mL); and (4) hydrogel coupled to Sema3C (200 μg/mL). Calcein staining of live cells revealed that the cell density was similar in all conditions and that neurons group in clusters. A few growing axons of unknown identity can be observed entering and crossing to the axonal compartment (Fig. 3B).

β III Tubulin and TH staining after fixation allowed the identification of DAx. In medium alone, only a few TH+ axons reached the axonal compartment (Fig. 3B). In the case of soluble Sema3C, we observed an increase in the crossing DAx; remarkably, the length and number of branches also increased compared to controls (Fig. 3B). The hydrogel combined with Cp presented a similar growth as the medium alone. For the Sema3C-hydrogel condition, we found more axons with a higher structural complexity (Fig. 3B). We measured the length of TH+ axons that enter and those that crossed the microgrooves with the ImageJ software. Considering all DAx that enter the microgrooves, soluble Sema3C and Sema3C-hydrogel had significantly longer axons than the corresponding control groups (Fig. 3C). Hydrogel embedded with Cp increased the length of TH+ processes when compared to control medium. Notwithstanding, the hydrogel + Sema3C condition significantly extended 1.6-fold DAx relative to Cp. The average number of DAx that entered the microgrooves was not different between groups (Fig. 3D).

Dopamine neurons differentiated from human ES cells express NRP1 and NRP2 receptors

We differentiated GFP-expressing H9 human ES cells, according to a published protocol to produce DAN.54 Using this protocol, after 28 days, 36.6% of neurons were TH+, and this proportion increased to 46.6% at day 35 of differentiation. To determine if DAN obtained from human ES cells express NRP1 or NRP2, receptors required to respond to Sema3C, we immunostained cells derived from the same cell suspension used for the assays with microfluidic devices presented below. At day 28, we quantified TH+ neurons and observed that NRP1 was present in 86% of DAN and 87.6% of cells were double-positive for NRP2 and TH. This high proportion was preserved in differentiated cultures of 35 days (Fig. 4), where we observed 88.8% and 90% of colocalization of TH with NRP1 or NRP2, respectively.

Sema3C released by hydrogel promotes dopaminergic axonal growth in neurons differentiated from human ES cells

At 21 days of differentiation, human ES cells were dissociated and plated on microfluidic devices. Immunostaining for β III Tubulin and TH showed that treatments with soluble Sema3C or Sema3C-hydrogel promoted the growth of more and larger DAx, in sharp contrast to treatments with medium only and hydrogel-Cp (Fig. 5A). In the soma compartment, the density of TH/β III Tubulin double-positive neurons was similar in all conditions. The right column of Fig. 5A shows a low-magnification GFP image, where it can be appreciated that the proportion of axons that entered the microgrooves is similar in all conditions. The orange and blue boxes correspond to the areas in the soma and axonal compartments, respectively. Similar to the observations in the rat mesencephalic neurons, the average length of axons was significantly higher in soluble Sema3C and Sema3C-hydrogel than in their corresponding controls (Fig. 5B). The axons located in the microgrooves were similar in all studied conditions (Fig. 5C).

Discussion

In this study, we show that hydrogel-embedded Sema3C is released to the medium maintaining constant levels of the protein for at least 28 days. PuraMatrix was reported to incorporate and release different molecules such as basic fibroblast growth factor, vascular endothelial growth factor, BDNF, bovine serum albumin, immunoglobulin G, and insulin.48,63,64 Our results confirm that proteins released by this hydrogel are functional as they exert the expected function in vitro.48,63 All these proteins have molecular weights between 7 and 38 kDa; Sema3C has a higher mass (82 kDa), but still can be released by PuraMatrix.

Using a three-dimensional collagen assay, we demonstrated in vitro that DAx of neurons differentiated from
FIG. 5. The axonal growth of human ES cells differentiated to DAN is promoted by Sema3C. Dopamine cells were studied in microfluidic devices in control (medium alone or hydrogel-Cp) or Sema3C (either soluble or coupled to hydrogel) conditions. (A) Immunostaining for β III Tubulin and TH shows that in the axonal compartment, there are more DAx in Medium + Sema3C and in Hydrogel + Sema3C conditions, compared to control medium and Hydrogel + Cp, respectively. In the soma compartment, a similar density of neurons positive for both TH and β III Tubulin is found close to the zone of microgrooves. The H9 ES cells express GFP that was used to visualize cell density in the soma compartment. In the right-most panel, a low magnification picture of the GFP+ axons in the microgrooves is shown. Note that in Medium + Sema3C and Hydrogel + Sema3C conditions, many axons were larger than the microgroove length (900 μm). (B) Quantification of total dopaminergic axonal length. Medium + Sema3C and Hydrogel + Sema3C conditions exert a positive effect in length, in contrast to control medium and Hydrogel + Cp, which had significantly shorter axons. (C) In all conditions, a similar number of DAx entered the microgrooves. Mean ± standard error of mean **p<0.01; ***p<0.001 (one-way ANOVA following Tukey’s multiple comparisons test). Scale bars: 100 and 300 μm. Color images available online at www.liebertpub.com/tea
mouse ES cells are attracted by Sema3C released by aggregated HEK cells. The current study shows that the PT embryonic region exerts axonal attraction, in contrast to the 1st region, which does not. Exposure of Mb explants to hydrogels coupled to Sema3C generates a similar attractive effect to that observed in PT. Conversely, hydrogel with Cp has analogous effects to that of 1st explants. These results indicate that the use of hydrogel coupled to Sema3C guides DAX growth, with a comparable efficiency to PT, which attracts these axons during development. Using a similar methodological approach, other groups have demonstrated that Netrin-1 exerts dopaminergic axonal attraction. To our knowledge, the current report is the first in which an hydrogel alone is able to deliver a molecule that guides dopaminergic axonal growth. Some articles showed that hydrogels can be used as scaffolds that require the combination with other particles that release molecules in vitro: a scaffold of synthetic poly(ethylene glycol) gel conjugated with biodegradable poly-lactic-co-glycolic acid particles with different concentrations of insulin-like growth factor directed the axonal growth of mouse ES cell-derived neurons. An increase of the axonal length of peripheral nervous system neurons with a scaffold of agarose hydrogel, combined with collagen and poly-lactic-co-glycolic acid particles that released nerve growth factor (NGF), has been also demonstrated. On the contrary, the guidance of axons during development is influenced by several molecules that interplay, sometimes with opposite effects, to direct axonal growth. The use of a choice point system allows the assessment of various cues simultaneously. A device based on immobilized and soluble proteins with different gradients was recently developed to test the effects of Sema6A and NGF on the axonal growth of dorsal root ganglia. Sema6A induces repulsion of neurite growth of sensory neurons of the lumbosacral region, but not other spinal regions, and NGF can attenuate this effect.

Collagen gels have been used previously to evaluate axonal growth. In such a matrix, axons can develop in all directions, making difficult to follow a trajectory in one plane. Thus, the measurement even with specialized software can be challenging. Therefore, we decided to use microfluidic devices, in which axons follow a linear trajectory for 900 μm in one plane. This system has been used to evaluate the effect of diffusible molecules on growing axons. In this work, we specifically looked at the effects of Sema3C using these devices, which unequivocally identify growing axons. Using DAN of rodent and human origin, we found that exposure of such neurons to hydrogel embedded with Sema3C results in an increase of DAX length, similar to soluble Sema3C. Therefore, Sema3C released by PuraMatrix to Sema3C seems to be equally functional as the soluble form, with regard to promotion of DAX growth, although with a 10-fold higher initial concentration. It is noteworthy that the times required to observe axonal growth in rodent and human neurons were different. PuraMatrix-released Sema3C was effective for rat cells (5 days) and also for human neurons (15 days), strongly suggesting that the delivery time of Sema3C by the hydrogel was optimal for these assays.

Microfluidic devices have been used to isolate axons of neurons differentiated from human ES cells, to observe mitochondrial movement across axons, to evaluate axonal injury, regeneration, and transport, to assess differentiation and neuronal entrance to microgrooves, and to identify bidirectional communication between motor neurons and muscle cells. Microfluidic systems with different characteristics in design have also been successfully used for generating chemical gradients and evaluating axonal growth cone responses. Previous data from our group have indicated that Sema3C increases DAX growth in vitro of DAN obtained from mouse ES. This is the first report on the positive effects of Sema3C on DAN derived from human ES cells, suggesting similar responses between human and rodent cells. We show that the dopaminergic differentiation protocol for human cells used in this study, as well as the protocol for rodent cells used previously, generates DAN neurons that respond to Sema3C. In fact, this response of human DAN was correlated with a high proportion of TH+ neurons that express the Sema3C receptors NRPI and NR2B.

Previously, the presence of NRPI and NR2B has been demonstrated in DAN obtained from the developing rat ventral Mb and DAN differentiated from mouse ES. These rodent DAN presented similar high proportions of TH+ neurons that express NRPI (77% for rat and 82% for mouse) or NR2B (48 for rat and 59% for mouse). Blocking antibodies for these receptors inhibit the action of Sema3C on attraction and axonal extension. Therefore, it was of interest to analyze if human DAN express NRPI and NR2B since no previous reports were available. Our results show that approximately 90% of DAN derived from human ES presented immunoreactivity to anti-NRPI or anti-NR2B antibodies, suggesting that the effects of Sema3C to promote axonal growth are mediated by these receptors.

Grafting DAN into the striatum of parkinsonian rats provides recovery in apomorphine- and amphetamine-induced rotational test. However, this ectopic graft induces behavioral complications such as increasing levodopa-induced dyskinesias. Furthermore, grafts in the putamen do not cause a consistent and steady recovery in PD patients. An alternative to such ectopic grafts in experimental animals is placing DAN in the SN and guiding DAX to reach the dorsal striatum. Previous data from our group have demonstrated that the intranigral graft of DAN derived from mouse ES cells, combined with HEK cells transfected with Sema3C, along an artificial trajectory from SN to striatum, promotes behavioral recovery of parkinsonian rats lesioned with 6-OHDA after 3 weeks. We report that Sema3C release was maintained during 4 weeks in vitro, so we can speculate that the use of this biomaterial scaffold could replace HEK cells to promote and guide axonal growth in vivo.

We used a biocompatible and biodegradable hydrogel that has been implanted into rodent brain with negligible glial reaction. This is a key issue, because hydrogels should incorporate and release proteins but without adverse tissue reactions. For example, a Poly-(ethylene glycol)-silica hydrogel has been used to release Sema3A, which in vitro promotes DAX growth; however, when it was implanted into the rat brain, the hydrogel produced an inflammatory response. Our results suggest that it might be possible to use PuraMatrix combined with Sema3C to promote and guide axonal growth in animal models of PD, using of DAN differentiated from human ES cells.

In addition to the effect of Sema3C on axonal growth, this protein can influence the development of new vessels, which has implications on vision. Sema3C inhibits the formation
of pathological endothelial tufts in an animal model of retinopathy of prematurity. This protein, acting through NRPs1 and Plexin D1, prevented AKT- and P38 MAPK-induced phosphorylations, resulting in apoptosis of developing microvasculature, decreased branching, loss of sprouting tips, and abnormal cell junctions. 5 It might be of interest to test if the controlled release of Sema3C by a hydrogel could be applied in this model.

Conclusion

In this study, we demonstrate that Sema3C can be embedded in the PuraMatrix hydrogel and this protein is released for at least 28 days in vitro. Hydrogel-released Sema3C can attract the growth of rodent embryonic DAs in a similar way to that of PT in collagen gels and also increases the axonal growth of human and rodent DAN. Remarkably, the response of these human neurons to Sema3C is very similar to that of rodent cells. This is the first time that a hydrogel is used as a protein carrier to increase dopaminergic axonal length.

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Disclosure Statement

No competing financial interests exist.

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