Differentiation of Human Neural Progenitor Cells in Functionalized Hydrogel Matrices

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Abstract

Hydrogel-based three-dimensional (3D) scaffolds are widely used in the field of regenerative medicine, translational medicine, and tissue engineering. Recently, we reported the effect of scaffold formation on the differentiation and survival of human neural progenitor cells (hNPCs) using PuraMatrix™/C212(RADA16-I) scaffolds. Here, we were interested in the impact of PuraMatrix modified by the addition of short peptide sequences, based on a bone marrow homing factor and laminin. The culture and differentiation of the hNPCs in the modified matrices resulted in an approximately fivefold increase in neuronal cells. The examination of apoptotic and necrotic cells, as well as the level of the anti-apoptotic protein Bcl-2, indicates benefits for cells hosted in the modified formulations. In addition, we found a trend to lower proportions of apoptotic or necrotic neuronal cells in the modified matrices. Interestingly, the neural progenitor cell pool was increased in all the tested matrices in comparison to the standard 2D culture system, while no difference was found between the modified matrices. We conclude that a combination of elevated neuronal differentiation and a protective effect of the modified matrices underlies the increased proportion of neuronal cells.

Key words: neuronal differentiation; PuraMatrix; ReNVM cells; tissue engineering

Introduction

Hydrogel-based scaffolds are widely used in the field of stem cell research, providing in vitro model systems that study different aspects as tissue regeneration, self-renewal, and differentiation of stem cells. Hydrogels consisting of synthetic nanofibres are excellent tools that generate such model systems, as they contain defined components, such as amino acids, resulting in a high reproducibility, and more over, they can be fabricated free of any animal products. Such scaffolds comprise three-dimensional (3D) networks of overlapping nanofibres, and have proved to be an effective environment for neural cells. Their mechanical properties, for example stiffness and flexibility, can be adjusted to the needs of the hosted cells. Pore sizes ranging between 50 and 100 nm allow the diffusion of gases, metabolites, and macromolecules for nutrition. In addition, 3D scaffolds of a consistent composition with predictable properties are ideal for combination with biometric cues. Such functionalized scaffolds have been used to study critical cell functions such as proliferation, differentiation, and migration of cells within the 3D-culture systems. A well-described hydrogel-based self-assembling scaffold, complying with the properties just mentioned, is PuraMatrix™ (RADA16-I). PuraMatrix (PM) was used in a series of studies to investigate proliferation and neuronal differentiation with stem and progenitor cells of different origins.

The next generation of self-assembling scaffolds is featured by the functionalization of the self-assembling backbone sequences with specific biological motifs. Laminin-derived sequences have been shown to be supportive of neuronal differentiation and survival when inserted into scaffolds. Functionalization of PuraMatrix with different short peptides, including among others a laminin-derived sequence (GGSDPGYIGSR) and a sequence found in the bone marrow homing factor (GGPFSSTKT), was shown to affect the mechanical properties of the matrix and alter neuronal differentiation. In a recent study, we supplemented PuraMatrix with laminin and described the fate of human neural progenitor cells (hNPCs) encapsulated in these 3D scaffolds, regarding differentiation and survival. Based on those findings, here we have tested the influence of modified
formulations of PuraMatrix (kindly provided by BD) on the neuronal differentiation of human NPCs. We used the immortalized hNPC line ReNcell VM (Millipore). This cell line has been described as an in vitro model system in a variety of studies dealing with neuronal differentiation in standard culture systems, as well as in 3D scaffolds, though the immortalization precludes any application in human studies. The cell line is characterized by a fast proliferation and a rapid onset of differentiation on the withdrawal of growth factors. The hydrogel was modified by the addition of short peptide sequences to the backbone, modifications that have been shown to influence the adhesion and differentiation of mouse neural stem cells in vitro. In this study, we were interested in the influence of the modifications on the differentiation of the hNPCs. Therefore, we proliferated and differentiated the cells in standard and two modified formulations of PuraMatrix. To overcome obstacles regarding the analysis of the cells hosted in the matrices, we adapted a protocol that releases the cells from the 3D scaffolds. The hydrogel was modified by the addition of growth factors. The hydrogel was modified by the addition of growth factors. The hydrogel was modified by the addition of growth factors.

**Methods**

**Culture of hNPCs in 3D scaffolds**

The hNPCs were cultured as previously described. In short, the cells were cultivated in Dulbecco's modified eagle medium (F12), supplemented with Glutamax, B27 media supplement, heparin sodium salt, and gentamycin (all Invitrogen). During proliferation, epidermal fibroblast growth factor (eFGF, 20 ng/mL) and basic fibroblast growth factor (bFGF, 10 ng/mL, both Roche) were added to the media. hNPCs were proliferated for 4 days and, subsequently, embedded in the 3D scaffolds as previously described. The hNPCs were proliferated for 7 days in all scaffolds. Differentiation was induced by the withdrawal of growth factors. The cells were differentiated for approximately 10 days with the media changed every 2–3 days. Based on the results of a previous study, we used a PuraMarix concentration of 0.25% to host the cells. In this study, PuraMarix and formulations (kindly provided by BD Bioscience) modified with two different peptide sequences, namely (1) -GGSDPGYIGSR- (denoted as PM-SDP) and (2) -GGFPSSTKT- (denoted as PM-PFS) were used. Nonmodified PuraMatrix is denoted as PM.

**Immunocytochemistry**

For immunocytochemistry, 3D cultures were fixed with paraformaldehyde (4% in 0.1 M phosphate buffered saline [PBS]) for 30 min and stored at 4°C in PBS with 0.02% Na-azide. For staining, nuclei labeling was performed with 4',6-diamidino-2'-phenylindoldihydrochloride (DAPI, 100 ng/mL in PBS, Molecular Probes). Subsequently, matrices were mounted with Mowiol/Dabco. For the staining (2 h at RT), the cells were centrifuged and resuspended in saponin buffer (PBS+0.5% bovine serum albumin [BSA]+0.02% Na-azide). For the staining, H$$\text{C}_{176}$$/D (Invitrogen, 1:100, mouse monoclonal), HuC/D (Invitrogen, 1:100, mouse monoclonal), PSA-NCAM (Millipore, 1:100, mouse monoclonal, IgM), Bcl-2 (Santa Cruz, 1:500, mouse monoclonal), or Alexa Fluor 488 (Molecular Probes, 1:1000, goat anti-mouse) or Alexa Fluor 488 (Molecular Probes, 1:1000, goat anti-mouse) for 1h in saponin buffer. The cells were washed again twice with saponin buffer and resuspended in wash buffer for analysis. A total amount of 50,000 cells of each probe was measured. Measurements were done using an FACScalibur instrument (Beckton-Dickinson) in combination with Cell Quest Pro software.

**TUNEL assay**

For TUNEL assay, we used an In Situ Cell Death Detection Kit (Roche). The cells were prepared as just described. Fixed cells (1% PFA for 15 min) were resuspended in HBSS buffer with 0.2% HSA. Next, the cells were permeabilized (0.1% Triton X-100 + 0.1% sodium citrate in PBS) for 2 min on ice and subsequently incubated in a TUNEL reaction mix for 1h at 37°C in a humidified atmosphere, in the dark. Samples were subsequently washed and stored in PBS until analysis. For positive controls, cells were incubated with DNase I (3000 IE/mL) to induce DNA strand breaks. Measurements were done using FACScalibur (Becton-Dickinson) in combination with Cell Quest Pro software.

For flow cytometry, 3D scaffolds were mechanically disrupted, and cells were released by adding trypsin. After centrifugation at 3000 g at RT for 5 min, the cells were washed twice with HBSS buffer. Subsequently, large cell/matrix aggregates were removed with a cell strainer (70 μm). After fixing the cells with 1% PFA for 15 min, the cells were resuspended in buffer (PBS+0.5% bovine serum albumin [BSA]+0.02% Na-azide). For the staining (2 h at RT), the cells were centrifuged and resuspended in saponin buffer (PBS+0.5% saponin+0.5% BSA+0.02% Na-azide) containing the first antibody against the jIL6-tubulin antibody (Santa Cruz, 1:100, mouse monoclonal), HuC/D (Invitrogen, 1:100, mouse monoclonal), PSA-NCAM (Millipore, 1:100, mouse monoclonal, IgM), Bcl-2 (Santa Cruz, 1:500, mouse monoclonal), or without the primary antibody providing a negative control. Afterward, the cells were washed twice with saponin buffer and incubated with the secondary antibody Alexa Fluor 647 (Molecular Probes, 1:1000, goat anti-mouse) or Alexa Fluor 488 (Molecular Probes, 1:1000, goat anti-mouse) for 1h in saponin buffer. The cells were washed again twice with saponin buffer and resuspended in wash buffer for analysis. A total amount of 50,000 cells of each probe was measured. Measurements were done using an FACScalibur instrument (Becton-Dickinson) in combination with Cell Quest Pro software.

**AnnexinV/propidium iodide apoptosis detection**

To determine the amount of apoptotic and necrotic cells, an FITC Annexin V Apoptosis Detection Kit I (BD Bioscience) was used. Cells were released from the scaffolds as just described and washed twice with HBSS. The cells were resuspended in binding buffer (0.1 M HEPES/NaOH [pH 7.4], 1.4 M NaCl, 25 mM CaCl$$\text{2}\text{)}$$, and Annexin V and propidium iodide (PI) were added according to the manufacturer’s manual. After 15 min incubation at RT on a shaker in the dark, the measurement was done by using FACScalibur (Becton-Dickinson) in combination with Cell Quest Pro software.
Statistics

All statistical analysis were performed with Prism 5 (GraphPad Prism, Inc.) using one-way analysis of variance with the Bonferroni Post test. p-value ≤ 0.05 was considered as indicating significant statistical differences. Values represent mean ± s.e.m.

Results

The aim of this study was to evaluate the effects of the two modified formulations of PuraMatrix on the hNPCs. Therefore, the proportion of neuronal cells and several parameters regarding neuronal differentiation and survival of the cells were analyzed by means of flow cytometry.

Expression of neuronal markers of hNPCs in modified PuraMatrix

Since the modifications are described as influencing the differentiation of adult mouse stem cells, we were interested in the effect of the modifications on the differentiation of human NPCs. Therefore, we cultured and differentiated hNPCs in PM, PM-SDP, and PM-PFS. In all three conditions, the cells grew in spheroid-like structures wherein slight differences in the growth pattern were observed by transmission light micrographs (Fig. 1A, C, E). The cells cultured in PM-PFS showed processes tending to leave the spheroid-like structures (Fig. 1C). To a certain extent, this could be observed in PM-SDP (Fig. 1E) but was hardly seen in PM (Fig. 1A).

On the induction of differentiation by the withdrawal of bFGF and eFGF, the cells started expressing the neuronal marker βIII-tub. It was striking that in both modified matrices, a higher amount of positive staining was observed in comparison to the normal PM (Fig. 1B, D, F), indicating a beneficial effect of the modified matrices on the differentiation of the hNPCs. The cells differentiated in PFS displayed the highest amount of βIII-tub⁺ cells and developed a very dense network of processes (Fig. 1D). Based on these results, the amount of neuronal cells in the scaffolds consisting of PM, PM-SDP, or PM-PFS was investigated.

Amount of neuronal cells in scaffolds consisting of different PM formulations

Recently, we demonstrated that the formation and structure of 3D scaffolds can impact the neuronal differentiation, where we performed immunocytochemical staining and determined the percentage of neuronal cells by manual counting. Since this method is extremely time consuming and only a small proportion of the cells can be analyzed, we adapted a method that extracts cells from 3D scaffolds and subsequently examines them by means of flow cytometry. We quantified the cells cultured in scaffolds consisting of PM, PM-SDP, and PM-PFS. An unstained sample (Fig. 1G) was used to determine the proper gate for the analysis of positive cells (Fig. 1H).

As evident from the fluorescence pictures (Fig. 1B, D, F), the quantification of βIII-tub⁺ cells revealed significantly higher proportions of positive cells in the PM-SDP and PM-PFS scaffolds in comparison to the PM scaffolds (Fig. 2A). After 4 days of differentiation (dd), we detected 17.7% ± 3.3% of βIII-tub⁺ cells in PM-SDP and 21.9% ± 4.2% in PM-PFS in comparison to 3.7% ± 0.9% of βIII-tub⁺ cells in PM. The difference was significant between PM and the modified formulations but not between the PM-SDP and PM-PFS; however, a slightly higher number was observed in PM-PFS. Interestingly, the amount of βIII-tub⁺ cells was stable over time in the modified matrices, where the number of cells in the PM scaffolds was increasing until the latest analyzed time point of 10 days. At this stage, the amount of βIII-tub⁺ cells was twice to thrice higher in PM-SDP and PM-PFS compared with PM. Noticeably, we observed βIII-tub⁺ cells in proliferating cells (day 0) in PM-SDP (2.8% ± 0.6%) and PM-PFS (3.3% ± 0.6%), most likely indicating the spontaneous differentiation of the cells in the modified matrices.

To consolidate these findings, we used an antibody against the human neuronal protein (HuC/D) as another marker for neuronal cells (Fig. 2B). We found a higher amount of positive cells at certain time points. After 4 dd, we found 7.4% ± 2.2% positive cells in PM, 9.9% ± 2.5% in PM-SDP, and 11.7% ± 1.6% positive cells in PM-PFS, where the amount was not significantly different. We found a significantly higher proportion of positive cells in PM-PFS after 7 dd in comparison to PM. In contrast to the results of the βIII-tub staining, the number of HuC/D⁺ cells was declining in PM scaffolds after 4 dd, where the number of positive cells in PM-SDP and PM-PFS changed at certain time points; however, the number of neuronal cells was always lower in PM in comparison to the modified scaffolds. Since HuC/D is described as marking cells shortly after their leaving the cell cycle, these results might indicate different developmental states in the different scaffolds. As a next step, we wanted to elucidate the mechanism contributing to the higher proportion of neuronal cells in PM-SDP and PM-PFS matrices.

Expression of Bcl-2 in scaffolds consisting of different PM formulations

Anticipating a protective effect of the modified matrices on the cells, we analyzed the amount of the anti-apoptotic protein Bcl-2. The cells that were released from each type of scaffold were analyzed by means of flow cytometry (Fig. 3). The analysis revealed a significant increase between day 0 (proliferating cells) and the cells differentiated for 4 days in all conditions. After 4 dd, the level of Bcl-2 changed only slightly in all conditions, though there was a prominent, but not significant, decline at days 7 and 10 in PM-SDP (Fig. 3, gray bars). With this exception, the lowest amount at the different time points was always detected in the scaffolds consisting of PM-PFS.

Amount of apoptotic and necrotic cells in the different matrix formulations

To further investigate a possible protective effect of the PM formulations on the hNPCs, we next analyzed the proportion of apoptotic and necrotic cells. We used an AnnexinV and PI staining to determine the amount of cells in an early apoptotic state (AnnexinV positive), a late apoptotic state (AnnexinV/PI positive), and the amount of necrotic cells (PI positive). Comparable to the level of Bcl-2, we found an increase of AnnexinV⁺ cells between day 0 (proliferating cells) and day 4 in all conditions (Fig. 4A). Although we had observed an increase in the different scaffolds, neither the difference between the formulations nor the difference between later time points was significant, most likely
due to the variation of the sampled data. For the amount of cells in a later stage of apoptosis, indicated by a double-positive staining for AnnexinV and PI (Fig. 4B), we observed a slight increase in double-positive cells between day 0 and day 4, followed by a slight decrease in positive cells over time, for all scaffolds. We found no significant differences between the scaffolds or over the time. Next, we analyzed the proportion of necrotic cells indicated by PI-positive cells (Fig. 4C). In all three types of scaffolds, we found an increase in necrotic cells between day 0 and day 4. In PM-SDP and PM-PFS scaffolds, the proportion of PI-positive cells declined between day 7 and day 10, where the number of necrotic cells in PM-PFS never reached the level of the necrotic cells in PM or PM-SDP. Moreover, the level of PI-positive cells did not decline but rather increased in the scaffolds consisting of PM. This result suggests that at least a certain proportion of the increased number of neuronal cells in the modified matrix formulations is based on a protective influence of the modified scaffolds.

**Proportion of TUNEL-positive neuronal cells in the different matrix formulations**

Since we had obtained the amount of Bcl-2+ cells and the proportion apoptotic/necrotic cells from the whole population of cells hosted in the matrices, we next analyzed the...
amount of neuronal cells undergoing apoptosis or necrosis by means of a combination of a TUNEL assay and βIII-tub staining. The quantification was done by means of flow cytometry measurements, and the percentage of TUNEL+ cells within the population of βIII-tub+ cells was determined.

The analysis of cells released from PM revealed an increase of TUNEL+ and βIII-tub+ cells from 41.1%–18.7% at day 0 (proliferating cells) up to 89.4%–5.6% after 10 dd (Fig. 5, white bars). Notably, the peak of neuronal cells in PM-SDP and PM-PFS was reached after 4 dd and was stable until the latest time point of observation (10 dd), whereas the amount of βIII-tub+ cells tends to increase over time in PM. (B) As a second neuronal marker, we used HuC/D. Comparable to the analysis of βIII-tub+ cells, we found a higher number of positive cells in the PM-SDP and PM-PFS. *p ≤ 0.05; **p ≤ 0.01. n.s., not significant; dd, days of differentiation.

FIG. 2. Quantification of neuronal cells. Comparison of the amount of neuronal cells hosted in the different PM formulations. To determine the size of the neuronal population, we used the neuronal marker βIII-tub and HuC/D. (A) The flow cytometry analysis revealed a significantly higher amount of βIII-tub+ cells in the modified formulations PM-SDP (gray bars) and PM-PFS (black bars) in comparison to PM (white bars). Notably, the peak of neuronal cells in PM-SDP and PM-PFS was reached after 4 dd and was stable until the latest time point of observation (10 dd), whereas the amount of βIII-tub+ cells tends to increase over time in PM. (B) As a second neuronal marker, we used HuC/D. Comparable to the analysis of βIII-tub+ cells, we found a higher number of positive cells in the PM-SDP and PM-PFS. *p ≤ 0.05; **p ≤ 0.01. n.s., not significant; dd, days of differentiation.

FIG. 3. Quantification of Bcl-2. Determination of the anti-apoptotic protein Bcl-2. To test the hypothesis of a protective effect on the modified PM formulations, we analyzed the amount of the anti-apoptotic protein Bcl-2 by flow cytometry. In all three conditions (PM [white bars], PM-SDP [gray bars], PM-PFS [black bars]), we observed a significant increase of the Bcl-2 level after 4 dd. After 10 dd, the lowest amount was found in scaffolds consisting of PM-PFS (19.5±1.9), and the highest amount was found in PM scaffolds (29.4±6.2), indicating a benefit of the PM-PFS scaffolds over PM scaffolds, with regard to a protective effect. *p ≤ 0.05; **p ≤ 0.01; ***p ≤ 0.001.
Discussion

The influence of 3D scaffolds on growth, proliferation, and neuronal differentiation is of great interest in order to find new methods for cell-based therapies. In the context of regenerative medicine, the combination of biomaterial scaffolds with neural stem and progenitor cells holds great promise as a therapeutic tool. Culture systems emulating a 3D environment have been shown to influence proliferation and differentiation in different types of stem and progenitor cells. Here, the formation and functionalization of the 3D microenvironment is important for the survival and fate of the embedded cells. PuraMatrix (RADA16), a peptide-based hydrogel scaffold, has been well described and used to study the influence of a 3D environment on different cell types. Recently, we have demonstrated the impact of the formation of PuraMatrix on the survival and differentiation of hNPCs. The combination of biodegradable scaffolds as PuraMatrix and hNPCs is of special interest, as scaffolds can be fabricated xenofree and in a highly standardized manner in comparison to animal-derived products with a high batch variability.

In this study, we used two modified formulations of PuraMatrix, wherein both formulations of short peptide sequences were linked to PuraMatrix. In PM-SDP, the peptide sequence—SDPGYIGSR—was added, which is a motif found in laminin and described as promoting cell adhesion and the
extensibility of neural cells. The second modification consists of the addition of the peptide sequence -GGPFSSTKT- a motif of the bone marrow homing factor that is shown to stimulate neuronal stem cell adhesion and differentiation in vitro. Both modified matrices have been used to promote the proliferation and differentiation of adult mouse neural stem cells. In accordance with the results of Cunha et al., we found a significantly increased amount of neuronal cells in both modified matrices, where in PM-PFS an approximately sevenfold increase was observed (Fig. 2A). The modifications are discussed to change the 3D structures of the matrices, resulting in changed mechanical properties as stiffness or flexibility. It was recently reported that the mechanical properties of a matrix can determine the fate of cells toward a neuronal phenotype (soft gels) or an osteogenic differentiation (hard gels). The stiffness of the matrices comparable to the matrices used here showed no differences, most likely ruling out a contribution to the observed increase in neuronal cells.

Besides changes in the mechanical properties of matrices due to modifications, changes in the permeability of a matrix might result in a more effective delivery of nutrients and oxygen to the cells and, thus, an improvement in the survival rate of cells. Based on this assumption, we determined parameters indicating the level of apoptotic or necrotic cells cultured in the different matrices by means of flow cytometry.

We observed the tendency of a higher survival rate of the cells in the modified matrices, although we did not find significant differences between the modified matrices and the nonmodified PuraMatrix. Consistently, we found a trend toward higher amounts of apoptotic cells and a higher proportion of apoptotic/necrotic neuronal cells in PM in comparison to the modified matrices PM-SDP and PM-PFS.

As recently described, the used hNPCs can be protected by the Na⁺ channel agonist Veratridine, indicating a protective effect depending on the activity of neuronal cells in the culture system. In addition, we described a changed functionality of the cells co-cultured with hippocampal brain slices, where we found the cells that receive synaptic input from the host tissue proofing the expression of neurotransmitter receptors.

The importance of the expression of neurotransmitter receptors in very early developmental stages of neurons or even neuroblasts is shown to be important for the survival of the cells in an increasing number of studies. For human mesencephalic NPCs, it was shown that a persistent application of the glutamate receptor agonist NMDA during the proliferation results in an increased number of TH immunopositive cells, indicating activity-dependent mechanisms. Similar results were obtained in a study using cortical NPCs. Taken together, one can deduce that NPCs need neuronal activity or a certain level of neurotransmitters to survive and mature. To support the idea that an increased neuronal activity might contribute to the higher amount of neuronal cells we observed in the modified matrices, we performed patch-clamp studies. Therefore, the cells were differentiated within the matrices, subsequently released, and plated on cover slips, as the cells situated in the scaffolds cannot be accessed for such studies. First, results indicated and evaluated the number of cells expressing functional Na⁺ channels (data not shown). However, further studies are necessary to elucidate differences in synaptogenesis and, subsequently, neuronal network activity determined, for example, by recording the synaptic currents underlying neuronal network activity.

Another aspect is that a higher density of cells leads to an improvement of cell-cell interactions. Cell-cell interactions play a crucial role in the proliferation, development, and survival of cells and might contribute to the higher amount of neuronal differentiation observed in the modified matrices.

Besides investigating the protective effects of the matrices, we asked whether the pool of neuronal-determined progenitor cells contributes to the higher amount of neuronal-differentiated cells. We used PSA-NCAM as a marker for cells that were determined to undergo neuronal differentiation. Although PSA-NCAM is not solely a marker for neuronal progenitor cells but also for glial progenitors, it was used for neuronal lineage selection in the past. Recently, it was shown for human ES cells that a preselection of PSA-NCAM-positive cells results in highly purified neuronal lineages, where the size of the PSA-NCAM⁺ pool increased over the time of differentiation, indicating an increasing pool of neuronal cells. In contrast, we found in our study that the level of PSA-NCAM⁺ cells did not change on the induction of differentiation (data not shown). Furthermore, we found no significant differences in the pool of PSA-NCAM-positive cells in proliferating cells (Fig. 6) within the different matrices. However, interestingly, we found a five to eight times higher amount of PSA-NCAM⁺ cells in the 3D scaffolds compared with cells cultured in a 2D monolayer system (Fig. 6). This leads us to the conclusion that the 3D environment, regardless of the composition used, has a massive impact on the neuronal fate-determined progenitor pool, although the underlying mechanism remains unknown, and that the modified matrices PM-SDP and PM-PFS possess benefits over the PM regarding the survival of differentiated cells over time.

In summary, our results demonstrate the importance of a lineage-specific proliferation, as shown by the impact of a 3D culture system on the pool of neuronal fate-determined cells. Furthermore, we have shown the beneficial influence of the environment on the survival of hosted cells. Taken together, these results demonstrate that the 3D scaffolds might provide a powerful tool for optimizing the yield of neural progenitors, and subsequent neuronal differentiation, for research and therapeutic applications.

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Authors’ Contribution

A.L. designed the study, performed experiments, analyzed data, and wrote the article. P.J.M. designed the study, performed experiments, and analyzed data. S.F., A.R., and M.J.F. designed the study, analyzed data, and wrote the article.

Author Disclosure Statement

The authors have nothing to disclose.

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