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Encapsulation and Delivery of Neutrophic Proteins and Hydrophobic Agents Using PMOXA−PDMS−PMOXA Triblock Polymersomes

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ABSTRACT: Polymersomes are attractive nanocarriers for hydrophilic and lipophilic drugs; they are more stable than liposomes, tunable, and relatively easy to prepare. The copolymer composition and molar mass are critical features that determine the physicochemical properties of the polymersomes including the rate of drug release. We used the triblock-copolymer, poly(2-methyl-2-oxazoline)-block-poly-(dimethysiloxane)-block-poly(2-methyl-2-oxazoline) (PMOXA−PDMS−PMOXA), to form amphipathic polymersomes capable of loading proteins and small hydrophobic agents. The selected agents were unstable neurotrophins (nerve growth factor and brain-derived neurotrophic factor), a large protein CD109, and the fluorescent drug curcumin. We prepared, characterized, and tested polymersomes loaded with selected agents in 2D and 3D biological models. Curcumin-loaded and rhodamine-bound PMOXA−PDMS−PMOXA polymersomes were used to visualize them inside cells. N-Methyl-D-aspartate receptor (NMDAR) agonists and antagonists were also covalently attached to the surface of polymersomes for targeting neurons. Labeled and unlabeled polymersomes with or without loaded agents were characterized using dynamic light scattering (DLS), UV−vis fluorescence spectroscopy, and asymmetrical flow field-flow fractionation (AF4). Polymersomes were imaged and tested for biological activity in human and murine fibroblasts, murine macrophages, primary murine dorsal root ganglia, and murine hippocampal cultures. Polymersomes were rapidly internalized and there was a clear intracellular co-localization of the fluorescent drug (curcumin) with the fluorescent rhodamine-labeled polymersomes. Polymersomes containing CD109, a glycosylphosphatidylinositol-anchored protein, promoted cell migration in the model of wound healing. Nerve growth factor-loaded polymersomes effectively enhanced neurite outgrowth in dissociated and explanted dorsal root ganglia. Brain-derived neurotrophic factor increased dendritic spine density in serum-deprived hippocampal slice cultures. NMDAR agonist- and antagonist-functionalized polymersomes targeted selectively neurons over glial cells in mixed cultures. Collectively, the study reveals the successful incorporation into polymersomes of biologically active trophic factors and small hydrophilic agents that retain their biological activity in vitro, as demonstrated in selected central and peripheral tissue models.

INTRODUCTION

In living cells, many cellular functions are performed by nanometer-scale vesicles.1,2 Biological vesicles store substances, such as neurotransmitters, and control the extent of their release.3−5 There is a current interest to use cellular vesicles, most notably exosomes,6−8 as carriers to deliver drugs, in view of their compatibility with cells and organs. Still, in the context of therapeutic carriers, biological vesicles have few limitations such as complex composition, limited stability, and modest loading of either lipophilic drugs or proteins.9 Synthetic...
vesicles constructed with amphiphilic block copolymers, or polymersomes, are designed to overcome these limitations.\textsuperscript{10,11} They accommodate lipophilic compounds in their membrane and hydrophilic agents in their aqueous core.\textsuperscript{14–15} Like most nanocarriers, polymersomes, were primarily developed as carriers for anticancer agents.\textsuperscript{16–20} Stimulus-controlled drug delivery from polymersomes has been demonstrated as well, using triggers such as changes in pH or in temperature.\textsuperscript{16,20–24}

There is such a diversity in polymer composition, structure, and functionality, that it is possible to design polymersomes adjusted to complement the properties of their cargo, from small synthetic molecules and peptides to large biological compounds such as proteins.\textsuperscript{25} We opted for self-assembling polymersomes made of the triblock copolymer, poly(2-methyl-2-oxazoline)-block-poly-(dimethylsiloxane)-block-poly(2-methyl-2-oxazoline) (PMOXA–PDMS–PMOXA), as their suitability for protein protection and transport was reported earlier.\textsuperscript{26} Moreover, PMOXA, which is known for its nonfouling properties, provides stealth properties to the polymersomes.\textsuperscript{27–29}

We assessed polymersomes in biological models in 2D ("wound in the dish") and 3D explanted ganglia\textsuperscript{30} with more complex structures consisting of different cell populations. Under normal conditions, wound healing occurs through a multistage pathway that includes coagulation, inflammation, formation of granulation tissue, and remodeling.\textsuperscript{31–34} In severe or chronic pathological conditions, the healing process is locked in a state of chronic inflammation due to the presence of cytokines, chemokines, and reactive oxygen species in the milieu, that cause further damage related to oxidative stress. Therapeutic agents currently used for wound healing include agents promoting epithelization and antibiotics to eliminate infection at the site of injury.\textsuperscript{35,36} In pathologies such as diabetes, wound healing is impaired because of inadequate trophic support due to peripheral nerve injury. This repair requires trophic support in the form of nerve growth factor (NGF), brain-derived neurotrophic factor (BDNF), and NT3 normally supplied and retrogradely transported from the target tissues (skin and muscles) to the soma [dorsal root ganglia (DRG)]. NGF promotes axonal growth in both the peripheral and central nervous system.\textsuperscript{37–40} NGF binds to the nerve terminal and is retrogradely transported to the cell body, where it signals the transcription factor CREB to upregulate prosurvival genes.\textsuperscript{41} BDNF is a 27 kDa protein, which when excreted by microglia has been reported to increase neuronal survival.\textsuperscript{41–43} NGF and BDNF exhibit wound healing properties and are excellent candidates for nerve repair.

Other agents have been proposed as therapeutics for wound healing. Anti-oxidants, such as curcumin (diferuoylmethane), a natural small hydrophobic and unstable molecule, have been shown in phase 1 clinical trials to have high safety and wound healing properties.\textsuperscript{44–47} CD109 is a membrane protein, approximately 180 kDa, which binds to TGF-β and represses TGF-β signaling in human keratinocytes.\textsuperscript{48–51} Because curcumin and proteins such as CD109, NGF, and BDNF are unstable in biological environments,\textsuperscript{52,53} it is advantageous to incorporate them into nanocarriers to protect them from hydrolytic and enzymatic cleavage to reach the site of injury.

We investigated the safety and effectiveness of PMOXA–PDMS–PMOXA triblock copolymer in delivering small hydrophobic molecules such as curcumin and larger hydrophobic proteins such as NGF and BDNF to promote wound healing and neural regeneration. We characterized the polymersome structure and drug loading using transmission-electron microscopy (TEM), asymmetrical flow field-flow fractionation, UV spectroscopy, and dynamic light scattering (DLS), while drug release was monitored using a dialysis bag method. In addition, drug-loaded polymersomes were tested in both 2D scratch assays as well as 3D organotypic cultures. This study demonstrates that PMOXA–PDMS–PMOXA is non-toxic and effective in delivering proteins and small molecules to macrophages, fibroblasts, and neurons of the DRG and in the hippocampus.

\section*{RESULTS AND DISCUSSION}

To protect the photolabile curcumin and the labile proteins (CD109, NGF, and BDNF) used in this study against harsh environments, we incorporated them into PMOXA–PDMS–PMOXA polymersomes.\textsuperscript{54} The triblock PMOXA–PDMS–PMOXA copolymer was selected in view of its known chemical stability and low toxicity.\textsuperscript{26,55} It was synthesized following a known procedure,\textsuperscript{26} shown schematically in Figure S1. The chemical structure of the copolymer and the synthetic intermediates was ascertained by \textsuperscript{1}H NMR (Figure S2) and \textsuperscript{13}P NMR spectroscopy (Figure S3). The molar mass of the copolymer (M\textsubscript{n} = 9.84 kDa) was determined by \textsuperscript{1}H NMR spectroscopy and gel permeation chromatography (GPC, M\textsubscript{n} = 8.8 kDa, M\textsubscript{w}/M\textsubscript{n} = 1.4). The chemical structure of PMOXA–PDMS–PMOXA is shown in Figure 1A, where the subscripts 25 and 75 represent, respectively, the numbers of MOXA and DMS units in the blocks, as determined from the \textsuperscript{1}H NMR spectrum of the copolymer.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig1.png}
\caption{(A) Chemical structure of PMOXA–PDMS–PMOXA copolymer. (B) Pictorial representation of polymersomes incorporating proteins (NGF, CD109, or BDNF) or hydrophobic molecules (curcumin) and covalently labeled with rhodamine and the NMDAR antagonist or agonist [memantine or R,S-tetrazolyglycine (TZG)].}
\end{figure}

Curcumin and various proteins were incorporated into PMOXA–PDMS–PMOXA polymersomes (Figure 1B) using the standard film hydration method. The hydrophobic nature of curcumin allowed for its incorporation into the polymersome PDMS membrane with 82.3% encapsulation efficiency (EE, Table 1). The curcumin content of PMOXA–PDMS–PMOXA polymersomes was 2.4% (w/w), which corresponds to an equivalent curcumin concentration of 167.57 μM for a polymersome concentration of 2.5 mg/mL. Higher EE was observed with NGF over bovine serum albumin (BSA); however, because the mixing ration of protein/triblock was much higher for BSA compared to NGF, these numbers...
Table 1. Characteristics of PMOXA–PDMS–PMOXA-Based Polymersomes

<table>
<thead>
<tr>
<th>complex</th>
<th>feed ratio&lt;sup&gt;b&lt;/sup&gt; (wt drug/wt PMOXA–PDMS–PMOXA)</th>
<th>$R_h$ (nm)</th>
<th>LC&lt;sup&gt;c&lt;/sup&gt; (%)</th>
<th>EE&lt;sup&gt;d&lt;/sup&gt; (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>empty polymersomes</td>
<td>N/A</td>
<td>66.1 ± 1.0</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>polymersomes (BSA)</td>
<td>0.322</td>
<td>68.4 ± 1.3</td>
<td>13.55</td>
<td>48.6</td>
</tr>
<tr>
<td>polymersomes (curcumin)</td>
<td>0.03</td>
<td>62.3 ± 0.6</td>
<td>2.41</td>
<td>82.3</td>
</tr>
<tr>
<td>polymersomes (NGF)</td>
<td>0.046</td>
<td>65.2 ± 0.5</td>
<td>4.4</td>
<td>100</td>
</tr>
<tr>
<td>polymersomes (curcumin + NGF)</td>
<td>0.03</td>
<td>67.1 ± 2.1</td>
<td>2.32</td>
<td>80.5</td>
</tr>
<tr>
<td></td>
<td>0.019</td>
<td>65.9 ± 4.0</td>
<td>20.72</td>
<td>4.3</td>
</tr>
<tr>
<td>polymersomes (CD109)</td>
<td>6.1</td>
<td>N/A</td>
<td>20.72</td>
<td>4.3</td>
</tr>
<tr>
<td>polymersomes—memantine</td>
<td>0.01</td>
<td>77.0 ± 3.0</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>polymersomes—TZG</td>
<td>0.01</td>
<td>75.5 ± 3.8</td>
<td>N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>

<sup>a</sup>DLS = dynamic light scattering; LC = loading content; EE = encapsulation efficiency; BSA = bovine serum albumin; RhB = rhodamine; NGF = nerve growth factor; TZG = R₅,5-tetrazolylglycine.<sup>b</sup>Ratio of initial drug to copolymer used to prepare the formulation. <sup>c</sup>LC = weight percent of the drug relative to copolymer in the final product. <sup>d</sup>EE = weight percent of drug content in the final formulation relative to amount added at the beginning of the process.

cannot be compared. A remarkable EE was achieved with NGF (loading of 4.4 wt %) with an initial mixing ratio of 4.6% (Table 1). A possible explanation for the high loading of NGF is its net positive charge at neutral pH (pI of 9.3)<sup>35</sup> compared to the net negative charge of BSA (pI of 4.7)<sup>37</sup> and its smaller size (27 kDa vs 66.5 kDa). These results suggest that PMOXA–PDMS–PMOXA is particularly a suitable nanocarrier for trophic factors, such as NGF.

The size distribution of the polymersomes (60 < diameter < 400 nm), determined by DLS, was relatively wide (Figure S4) with $R_h$ values between ~65 and 75 nm, depending on the sample (Table 1). Additional analysis of the polymersomes was performed by asymmetrical flow field-flow fractionation (AF₄) (Figures 2 and S11) to ascertain the incorporation of curcumin and NGF and to measure polymersome sizes using the online-DLS detector. This method is known to be more precise than batch-mode DLS.<sup>38</sup> The morphology of the polymersomes was assessed before and after loading of NGF by TEM (Figure S12). The electron micrographs show spherical structures with diameters ranging from 10 to 140 nm. Size distribution after loading NGF was narrower.

The fractograms presented in Figure 2A, which monitor the absorption at 280 nm of the eluting species (left-hand ordinate), indicate that curcumin- and NGF-loaded polymersomes elute at elution times similar to the elution time of pristine polymersomes. The three eluting polymersomes have similar $R_h$ values, as judged from the fractograms in Figure 2A (right-hand ordinate). The size distributions of pristine polymersome alone and curcumin-loaded polymersomes are narrow, while polymersomes charged with NGF have a slightly broader size distribution and a larger $R_h$ value. Rhodamine-labeled polymersomes loaded with NGF were fractionated by a modified elution protocol (method 2) designed to fractionate free NGF (elution time ~9 min) and NGF-loaded polymersomes (elution time ≈ 22 min). As seen in Figure 2B, the NGF loading in the polymersomes was complete, with no detectable free NGF dissolved in the polymersome suspension. The rhodamine-labeled polymersomes incorporating NGF were larger than empty polymersomes by about 25 nm in $R_h$ value (Figure 2BC). We confirmed that NGF does not interact with the regenerated cellulose membrane used in the AF₄ channel by analyzing NGF solutions of increasing concentration. The elution signal of NGF measured by the UV detector was proportional to the amount of NGF injected (Figure S5).

Release of curcumin by the polymersomes was assessed by a dialysis-based method. The results presented in Figure S6 indicate that the polymersomes sustain the release of curcumin over extended time. After a 48 h-incubation in a PBS saline solution (pH 7.4), about 70% of the initial curcumin is retained in the dialysis tube, hence within polymersomes. Dialysis of free curcumin in the same conditions leaves less than 4% curcumin in the dialysis bag. The NGF release from the polymersomes was measured using a dialysis bag method with a 50 kDa cutoff membrane. In this case, we used a rhodamine-labeled NGF and measured the time-dependent Rho-NGF content of the polymersomes remaining in the

Figure 2. AF₄ fractograms of PMOXA–PDMS–PMOXA polymersomes. (A) AF₄/UV/DLS fractograms monitored by the UV detector (left ordinate) and by DLS (right ordinate), using the AF₄ method 1, showing the UV absorbance at 280 nm as a function of elution time for empty PMOXA–PDMS–PMOXA polymersomes (black curve), curcumin-loaded polymersomes (blue curve), and NGF-loaded polymersomes (red curve) in phosphate-buffered saline (PBS; 10 mM, pH 7.4); PMOXA–PDMS–PMOXA concentration: 0.5 g/L. The scattered symbols represent the hydrodynamic radii of the eluting polymersomes. (B) AF₄ fractograms from AF₄ method 2 of eluting rhodamine-labeled polymersomes with and without NGF. Both samples were injected at similar PMOXA concentration (2.0 g/L). The elution of the polymersomes from the AF₄ channel was also monitored by the fluorescence detector, the fluorescence elution profile matches well with the elution profile from the UV/vis detector (data not shown). (C) Size distribution of empty, NGF-loaded, and BDNF-loaded polymersomes plotted as the differential weight fraction as a function of the hydrodynamic radius ($R_h$).
dialysis bag by spectrofluorometry using the emission of rhodamine \( (\lambda_{ex} = 522 \text{ nm}) \). Nearly 60% of the NGF remained encapsulated after a 24-h incubation and 35% NGF was still retained after 48 h (Figure S6C,D). The AF4 elution time of the polymersomes (Figure S7A) increases as a function of dialysis incubation time, which could be indicative of polymersome swelling for longer dialysis times. In addition, two new eluting species are observed, one at early elution times (left arrow, Figure S7A) and one at late elution times (right arrow for times 137 and 235 h). The peak at early elution times corresponds to the triblock copolymers and free NGF (Figure S5), which provides evidence of the destabilization of the polymersomes. The large late eluting band (right arrow) is due to species larger than the polymersomes (ca. 700–800 nm, Figure S7B) could correspond to large aggregates which would assemble from the swollen polymersomes.

PMOXA–PDMS–PMOXA polymersomes loaded, or not, with curcumin and BSA were tested first in cell cultures of macrophages and human fibroblasts to assess the toxicity of polymer toward these cells. The cells selected are essential components of the skin, a target tissue innervated by DRG that we intend to use in the in vitro polymersomes evaluations in addition to 3D DRG cultures. The first task was to show that polymersomes are indeed not cytotoxic toward these cells, which were not evaluated in previous studies of PMOXA–PDMS–PMOXA polymersomes.

A concentration-dependent evaluation of pristine PMOXA–PDMS–PMOXA polymersomes confirmed that they do not impair the mitochondrial metabolic activity and do not reduce the number of viable cells (Figure 3). J774A.1 macrophage cells were also incubated for 24 h with equimolar concentrations of polymersome nanoparticles (upward triangle), curcumin-loaded polymersomes (square), and curcumin alone (circle). (A) Mitochondrial metabolic activity was determined using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide (MTT) assay. The data are presented as the mean ± standard error of the mean (SEM) obtained from at least three independent experiments. (B) Cell viability was determined by cell counting. Cell nuclei were labeled with Hoechst 33258 (10 μM, 10 min) and 9 fields per sample were counted using a fluorescent microscope. The data are presented as the mean ± SEM obtained from at least three independent experiments. Statistically significant differences from the untreated control were tested by Dunnett’s test and are indicated by *p < 0.05, **p < 0.01, ***p < 0.001.

Next, we investigated whether polymersome-loaded NGF can increase neurite outgrowth in peripheral DRG dissociated and 3D explant cultures. The restoration of nociceptive fields post skin denervation through the collateral sprouting by nearby undamaged nerves is dependent on endogenously produced NGF. Skin and muscle are the target tissues innervated by the nerves from the DRG. The experiments using DRG dissociated and explant 3D cultures described next were carried out to test if NGF retains its biological activity when incorporated into PMOXA–PDMS–PMOXA polymersomes (Figure 6A,B).

Neurite outgrowth of dissociated DRG neurons was enhanced significantly in the presence of NGF, with or without polymersomes as seen in Figure 6A,B. The extent of neurite outgrowth triggered by free NGF (50 ng/mL) and polymersome-incorporated NGF was comparable, although the NGF concentration within polymersomes was several-fold higher. In NGF release kinetics (Figure S6C,D), we observed that the amount of NGF released in the medium within 24 h corresponds to 40% of the initial content. Nerves in the target tissues (skin and muscle) are NGF-responsive in the wound healing processes. Further studies are necessary to assess if and how polymersome-incorporated trophic factors combined with small molecules are advantageous in re-establishing normal collateral sprouting and enhancing the rate of wound closure in animal models. We tested also NGF-containing polymersomes

 Considering that the site of injury is invaded rapidly by macrophages, it would be advantageous to exploit the phagocytic and macroinocytic properties of macrophages for delivery of drugs. This envisioned strategy promoted us to investigate the rate and extent of polymersomes internalization. Macrophages were treated with rhodamine-B (Rhb)-tagged polymersomes loaded with curcumin. Taking advantage of intrinsic fluorescence of curcumin, we detected the location of the polymersomes and their cargo (curcumin). Confocal fluorescence imaging revealed that polymersomes are internalized by J774A.1 macrophage cells within 5 min (Figure 4). Co-localization of Rhb with curcumin was confirmed, proving that polymersomes are capable of delivering their cargo within cells.

Fibroblast migration is an indicator of wound healing, and models, such as the “wound in the dish” model, allow the evaluation of wound healing therapeutic agents. We used scratch assays to determine if the biological activity of compounds loaded into polymersomes is retained or, possibly, enhanced compared to their free form. The experimental conditions were set to measure the parameters of cell viability and migration required for a successful wound healing using both mouse and human fibroblasts (Figure 5). In this assay, an enhanced migration of fibroblasts toward the scratched surface is considered an indicator of promoted wound healing. We tested CD109-loaded polymersomes. The following control treatments were performed: (i) free fibroblast growth factor (FGF, 0.3 μM, a positive control for cell migration), (ii) free mitomycin C (MitC, 15 μM, an inhibitor of cell migration), and (iii) pristine polymersomes for 24 h. Representative fluorescent micrographs are illustrated in Figure 5A with accompanying quantifications in Figure 5B. Treatment with FGF enhanced the fibroblast migration (1.55 fold), whereas treatment with MitC almost completely blocked fibroblast migration. Results from this assay show a significant increase in fibroblast migration, suggesting that FGF and CD109 might be beneficial in wound healing as noted also in earlier findings.

Figure 3. Polymersomes are well tolerated by macrophage cells. J774A.1 macrophage cells were incubated for 24 h with equimolar concentrations of polymersome nanoparticles (upward triangle), curcumin-loaded polymersomes (square), and curcumin alone (circle). (A) Mitochondrial metabolic activity was determined using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide (MTT) assay. The data are presented as the mean ± standard error of the mean (SEM) obtained from at least three independent experiments. (B) Cell viability was determined by cell counting. Cell nuclei were labeled with Hoechst 33258 (10 μM, 10 min) and 9 fields per sample were counted using a fluorescent microscope. The data are presented as the mean ± SEM obtained from at least three independent experiments. Statistically significant differences from the untreated control were tested by Dunnett’s test and are indicated by *p < 0.05, **p < 0.01, ***p < 0.001.
in a 3D model of the peripheral nervous system (DRG explants, Figure 6C,D) and noted that these polymersomes significantly promote the neurite outgrowth. Three-dimensional models both for peripheral and central nervous system have a number of advantages over dissociated cultures including retained connections, the ability to monitor long term time-dependent changes from progenitor to fully differentiated cells, and more reliable testing of therapeutic interventions.61−63

Both peripheral nerves and central nervous system tissues respond to BDNF. BDNF has been established as a crucial trophic factor in brain development64 and neurological
Spine morphology and density contribute to the synaptic plasticity and functions. Several NMDA receptor agonists and antagonists have been evaluated for modulating synaptic functions in different models.\(^{75,76}\) Compounds investigated include TZG and memantine, potent agonist and antagonist, respectively.\(^{77,78}\) Memantine-coated gold nanoparticles have been shown to target exclusively extrasynaptic NMDAR as the large size prevents their entry in the synaptic cleft. This spatial segregation allows blocking of extrasynaptic NMDAR without affecting synaptic NMDAR. Savchenko et al. have shown that the gold nanoparticles exhibit neuroprotective effect by reducing glutamatergic cytotoxicity.\(^{79}\) We functionalized the surface of polymersomes with TZG or memantine and imaged them in mixed cortical cultures. Preliminary results (Figure S13) indicate memantine, and TZG-functionalized rhodamine-labeled polymersomes are predominantly found in neurons, whereas rhodamine-labeled polymersomes that do not present memantine or TZG are present in both glial cells and neurons. Thus, follow-up studies should evaluate electrophysiological parameters in models of central nervous system pathologies.

In summary, the study presented here demonstrates that PMOXA–PDMS–PMOXA self-assembles to form highly colloidal stable polymersomes that are nontoxic to murine macrophages, human fibroblast cells, and 3D models (DRG explants and hippocampal organotypic slice cultures). It suggests that surface-modified PMOXA–PDMS–PMOXA polymersomes could be useful nanocarriers for various trophic factors, such as neurotrophin-3, insulin-like growth factor, and in combination with small biologically active agents or synaptic modulators. Collectively, PMOXA–PDMS–PMOXA polymersomes merit further investigations in disease models in neurodegenerative disorders.\(^{65–67}\) Despite the positive neurotrophic effects of BDNF, clinical use of BDNF has been hampered by its instability and rapid clearance; BDNF has a circulating half-life of \(\sim 10\) min.\(^{68}\) We explored polymersome delivery of neurotrophic protein BDNF in the central nervous system using mouse hippocampal 3D organotypic slice cultures. BDNF is a crucial regulator of memory formation in the hippocampus.\(^{69}\) It is known to increase dendritic spine density\(^{70,71}\) and to facilitate LTP.\(^{72}\) The hippocampus, an area associated with memory formation, is especially sensitive to traumatic brain injury (TBI). Even mild TBI, which does not produce systematic vascular injury or neuronal loss, causes dendritic and synaptic degeneration.\(^{73}\) Repeated mild TBI leads to loss of long-term potentiation.\(^{74}\)

We observed that polymersome-loaded BDNF has an activity comparable to that of free BDNF on modulating dendritic spine density in organotypic hippocampal cultures (Figure 7A,B). While treatment with pristine polymersomes (concentration-matched control, 211 \(\mu g/mL, 72\) h) did not significantly affect either the subtype or the total density of dendritic spine density in serum-free media, application of free BDNF (100 ng/mL, 72 h) or polymersome-incorporated BDNF (100 ng/mL BDNF, 211 \(\mu g/mL\) polymersomes, 72 h) significantly increased the total dendritic spine density compared to both controls: hippocampi not treated with polymersomes or treated with drug-free polymersomes in serum-free media.

## Figure 6. (A) Dissociated DRG cultures treated with NGF-loaded polymersomes increase neurite outgrowth. Phase contrast micrographs. Scale bar = 100 \(\mu m\). (B) Quantification of the neurite outgrowth. Note the significant increase of neurite outgrowth with NGF treatment. (C) Bright-field photomicrographs of DRG explants after 4 days in culture. To enhance the contrast of neural cells against the background, we used MTT (500 \(\mu g/mL, 3\) h) and imaged with a light microscope at 10x (Leica DMI6000B). Images were analyzed using ImageJ. Scale bar = 400 \(\mu m\). (D) Quantiﬁcation of the neurite area expressed as a percentage of control (PBS) \((n = 8)\). The total neurite area is normalized as shown in Figure S9. Error bars represent SEM. \(* p < 0.05.\)

## Figure 7. Polymersome-incorporated BDNF increases postsynaptic dendritic spine density of pyramidal neurons in organotypic hippocampal slice cultures. Cultures were treated with BDNF (100 ng/mL), polymersome-loaded BDNF (100 ng/mL BDNF; 211 \(\mu g/mL\) polymersome), or empty polymersomes (211 \(\mu g/mL\)) for 72 h in serum-free media (S−). Serum-containing media (S+) acted as a positive control. (A) Representative photomicrographs of dendritic spines. Scale bar = 1.5 \(\mu m\). (B) Quantification of spine density and subtype exposed to above conditions. \(n \geq 12\) spine segments from at least four independent samples. Statistical signiﬁcance was assessed by one-way ANOVA followed by Dunn’s test. \(* p < 0.05\) compared to S− Ctrl.
vivo to evaluate if they are superior to the unincorporated agents at the peripheral and central target tissues. Polymeromes-containing proteins and small lipophilic molecules provide new avenues for multidrug therapies to be considered for interventions in neurological disorders.

EXPERIMENTAL SECTION

Chemicals. BSA (98%, electrophoresis), curcumin (from Curcuma longa (Turmeric), FGF, laminin, PKH67, MitC, MTT, RhB-isothiocyanate, Hank’s balanced salt solution (HBSS), sodium chloride (NaCl), α,α,α-trifluorotoluene, poly(dimethylsiloxane), 2-methyl-2-oxazoline, 1,2-dichloroethane, trifluoroacetic anhydride, deuterated chloroform (CDCl3), dimethyl-sulfoxide (DMSO), triethylamine (99%), 4-(dimethylamino)pyridine (DMAP, 99%), succinic anhydride (99%), N-hydroxysuccinimide (NHS, 98%), N,N′,N′′-dicyclohexylcarbodiimide (DCC, 99%), and anhydrous dichloromethane (99.8%, containing 50–150 ppm amylene as stabilizer) were purchased from Sigma-Aldrich (St. Louis, MO, USA). Regenerated cellulose dialysis membranes [molecular weight (MWCO): 3.5 kDa] were obtained from Spectrum Labs (Rancho Dominguez, CA, USA). NGF 2.5S was purified from male mouse salivary glands. Recombinant human CD109 protein was a gift of Dr. Anie Philip. Anhydrous ethanol was purchased from GreenField Specialty Alcohols Inc. (Toronto, ON, Canada). Sephacryl S-300 HR (Mw: 10–1500 kDa) was purchased from GE Healthcare Bio-Sciences (Pittsburg, PA, USA). PBS, Dulbecco’s modified Eagle’s medium (DMEM), bovine calf serum, and penicillin–streptomycin were purchased from Gibco (Thermo Fisher Scientific, Waltham, MA, USA). Rat tail collagen and RPMI-1640 were purchased from Invitrogen (Thermo Fisher Scientific, Waltham, MA, USA). Paraformaldehyde was purchased from Fisher Scientific (Waltham, MA, USA). Hoechst 33258 and Alexa Fluor 488 goat anti-mouse were purchased from Molecular Probes (Eugene, OR, USA). Trypsin was purchased from Worthington Biochemical Corp. (Lakewood, NJ, USA). β-Tubulin III was purchased from EMD Millipore (Billerica, MA, USA).

Triblock Copolymer Synthesis. PMOXA25–b-PDMS75–b-PMOXA25 was prepared following a known procedure (Figure S1). Its synthesis is given in Supporting Information, together with its structure and characterization by NMR spectroscopy and GPC. The PDMS block has 75 units and the PMOXA blocks have each 25 monomer units.

Polymeromes Preparation and Loading. Polymeromes were formed by the film hydration method. PMOXA25-b-PDMS75-b-PMOXA25 (10 mg) was placed in a round-bottom flask and dissolved with 2 mL of ethanol for 1 h. The ethanol evaporated under reduced pressure at a temperature of 40 °C, and the resulting thin polymer film was dried for 4 h under vacuum at room temperature. Subsequently, the film was rehydrated using a solution of PBS (10 mM, pH 7.4, 137 mM NaCl, 4 mL, final PMOXA–PDMS–PMOXA concentration of 2.5 mg/mL). The film was placed on a rotating agitator at room temperature for agitation overnight and then stirred magnetically for 4 h with a small Teflon stirring bar. The polymeromes suspension was extruded (11 times) through polycarbonate membranes (0.4 μm pore size followed by 0.1 μm pore size, Avestin, Ottawa, ON, Canada) to adjust the polymeromes size and sharpen the size distribution.

Curcumin was loaded during the film formation by adding 0.3 mg of curcumin to the polymeromes solution and sonicated for 30 s prior to ethanol evaporation. For protein loading, NGF 2.5S (26 kDa), BDNF (27 kDa), CD109 (180 kDa), or BSA (66.5 kDa) were dissolved in the PBS aqueous solution used to rehydrate the polymer film. Excess free protein was removed by filtration through centrifugal filter units with an MWCO of 100 kDa (Amicon, EMD Millipore, Billerica, MA, USA). Free CD109 was separated from polymeromes-encapsulated CD109 using Sephadex G-200 gel filtration columns (Sigma-Aldrich, St. Louis, MO, USA).

Curcumin and Protein Loading Content Determination. For curcumin content determination, an aliquot of the solution after rehydration was kept and diluted 200 times using a solution of Tween 80 (1%) in PBS. An aliquot of identical volume of the polymeromes suspension after extrusion and removal of excess free curcumin was also diluted 200 times in a Tween 80 solution. Both samples were analyzed by fluorescence spectroscopy (λex = 431 nm, λem = 534 nm). The EE was found by summing the areas under the emission bands for the two aliquots, and the concentration of encapsulated curcumin was calculated by multiplying the EE and the initial concentration of curcumin used. The equations for EE and loading content calculations are found below:

\[
EE \text{ (weight %)} = \frac{\text{weight of drug or protein in the polymeromes}}{\text{weight of the drug or protein used initially}} \times 100%
\]

\[
\text{Loading content (weight %)} = \frac{\text{weight of drug or protein in the polymeromes}}{\text{total weight of the polymeromes tested}} \times 100%
\]

The amount of protein in the polymeromes was determined by UV absorbance spectroscopy. The absorption spectrum of the sample after preparation was compared to that of polymeromes prepared under identical conditions but without proteins. The protein-loaded polymeromes were purified by centrifugal filtration through 100 kDa membranes (EMD Millipore, Billerica, MA, USA) or G-200 Sephadex columns for CD109 separation to remove free protein. Fresh PBS was added to the recovered polymeromes that were subjected to three additional filtrations. A UV-absorption spectrum of the retentate, diluted to its initial volume with fresh PBS, was measured. The EE was determined as the ratio of the absorbance of the suspensions after and before filtration (λ = 280 nm).

RhB Labeling of PMOXA–PDMS–PMOXA. To detect the formation of polymeromes and their internalization in cells (in vitro), the copolymer PMOXA–PDMS–PMOXA was labeled by conjugating RhB isothiocyanate to the –OH end groups. RhB isothiocyanate in dichloromethane (0.5 μmol) was added to PMOXA–PDMS–PMOXA (0.01 mmol) dissolved in dichloromethane (1 mL) and left to react for 10 h at RT. Excess RhB was removed by dialysis first in ethanol and then in deionized water (MWCO 5 kDa) for 48 h followed by at least three exchanges with water. The RhB-tagged polymer was isolated by lyophilization.

RhB Labeling of BSA and NGF. BSA (10 mg), or NGF (0.5 mg), was dissolved in aqueous sodium bicarbonate buffer (0.1 M, 1 mL, pH 8.5). A solution (50 μL) of RhB
isothiocyanate in dimethylformamide (10 mg/mL), briefly sonicated and stirred, was added to a stirred BSA solution (or 6.3 μL for NGF solution) for 1 h at room temperature. RhB-labeled proteins were purified from free dye by two consecutive elutions through Sephadex G25 columns.

**Labeling of PMOXA–PDMS–PMOXA with Drugs.** Dicarboxyl acid-modified poly(2-methyl-2-oxazoline)-block-poly(dimethylsiloxane)-block-poly(2-methyl-2-oxazoline) tri-block copolymers (HOOC–PMOXA25−b–PDMS75−b–PMOXA25−COOH) were synthesized as follows: succinic anhydride (39 mg, 0.39 mmol), triethanolamine (50 μL, 0.36 mmol), and DMAP (6.9 mg, 0.06 mmol) were added to a solution of HO–PMOXA25−b–PDMS75−b–PMOXA25−OH (500 mg, 0.07 mmol) in anhydrous dichloromethane at 0 °C under argon and reacted for 24 h. The reaction mixture was dialyzed against dichloromethane for 2 days while changing the dichloromethane four times. NHS-activated ester-modified PMOXA25−b–PDMS75−b–PMOXA25 block copolymers (NHS–PMOXA25−b–PDMS75−b–PMOXA25–NHS) were synthesized as follows: NHS (30 mg, 0.26 mmol) and DCC (35 mg, 0.17 mmol) were added to an anhydrous dichloromethane solution of HOOC–PMOXA25−b–PDMS75−b–PMOXA25–COOH (300 mg, 0.04 mmol) at room temperature under argon. The mixture was maintained in this condition for 24 h. Finally, it was dialyzed against dichloromethane for 2 days while changing dichloromethane four times, followed by dialysis against methanol to remove dichloromethane. The NHS–PMOXA25−b–PDMS75−b–PMOXA25–NHS block copolymer was characterized by 1H NMR spectroscopy (solvent: CDCl3; Figure S10).

Polymersomes were prepared with 5% rhodamine-labeled PMOXA–PDMS–PMOXA and 95% unlabeled PMOXA–PDMS–PMOXA or 5% rhodamine-labeled PMOXA–PDMS–PMOXA, 5% NHS-terminated PMOXA–PDMS–PMOXA, and 90% unlabeled PMOXA–PDMS–PMOXA in the case of drug-attached polymersomes. The triblocks were labeled proteins were purified from free dye by two consecutive elutions through Sephadex G25 columns.

**Curcumin and NGF Release from Polymersomes.** Curcumin release from polymersomes was measured using the dialysis bag method in a solution of PBS and Tween 80 (1%, v/v). Tween 80 was added to solubilize curcumin, which has a low solubility in PBS. A suspension of curcumin-loaded polymersomes in PBS-Tween 80 (2 mL, C_{Cur} = 0.07 mg/mL) was introduced in a dialysis tube (MWCO = 20 kDa, Slide-A-Lyzer MINI Dialysis, 2 mL) and dialyzed against PBS-T (44 mL) at 37 °C. At predetermined time intervals, the entire release medium was replaced by fresh medium to maintain perfect sink conditions for curcumin. A solution of curcumin (0.1 mg/mL in PEG400–water–dimethylacetamide (45:40:15 v/v/v) was used as a control. Curcumin concentration in the release samples was measured using spectrofluorometry. The concentration of curcumin in the dialysis compartment was determined as well by taking aliquots (50 μL) and diluting them 100-fold in ethanol before spectra analysis.

A solution of PMOXA–PDMS–PMOXA polymersomes loaded with RhB-labeled NGF (5 mL, polymersome concentration: 0.7 mg/mL) was placed in a dialysis tube (MWCO: 50 kDa) and dialyzed against PBS (1 L) at 37 °C for several days. Aliquots of the solution in the dialysis bag were taken over time and analyzed by UV absorbance, fluorescence, and AF6. The PBS solution was replaced with fresh PBS at these times.

**Asymmetrical Flow Field-Flow Fractionation.** Information about the instrument used and the two methods developed for the analysis of the polymersomes can be found in the Supporting Information section.

**Transmission Electron Microscopy.** Data were collected on a FEI Tecnai 12 BioTwin 120 kV TEM. Images were taken with an AMT XR80C CCD Camera System. Carbon-coated grids were first treated in a glow-discharge apparatus set at negative 25 mA for 30 s. Rapidly after, a drop of the polymersome sample (stock concentration at 2.5 mg/mL, 5 μL) was deposited on the grid and left for 1 min. A Whatman filter paper was used to wick away excess sample by slowly bringing it into contact with the side of the grid. A uranyl acetate solution (2%) was used as a negative staining agent, a 5 μL drop was deposited and left for 1 min before being wicked away with a clean Whatman filter paper. The grids were placed on a Whatman filter paper and left to dry in a Petri dish at room temperature until imaging (>15 min).

**Macrophage Cultures and Live Cell Imaging.** Mouse macrophage (J774A.1) cells (ATCC, TIB-67) were cultured in DMEM containing 10% fetal bovine serum and 1% penicillin–streptomycin. Cells were used between 5 and 25 passages.

In preparation for live cell imaging, macrophages were seeded at a density of 20 000 cells/coverslip onto rat tail collagen-coated cover slips (Fisher Scientific). Following cell treatment, plasma membranes were labeled with 2 μM PKH67 for 15 min and subsequently washed with PBS containing 1% BSA for 5 min. Prior to imaging, cells were washed with PBS containing 1% BSA for 5 min. Fluorescence micrographs were acquired with a Zeiss LSM 710 confocal microscope with Zeiss Zen imaging software at 63× magnification.

**MTT Cell Viability Assay.** We assessed mitochondrial activity, an indirect measure of cell viability using the colorimetric MTT assay. Macrophage cells were seeded at 100 000 cells/well in 24-well plates (Sarstedt, Nümbrecht, Germany). Following treatment, cell media was removed and replaced with serum-free media containing MTT (0.5 mg/mL), and cells were then incubated for 1 h at 37 °C. Following the incubation, media were removed, cells were lysed, and formazan was dissolved with DMSO. The dye absorbance was measured at 595 nm using a Benchmark microplate reader (Bio-Rad Life Science Research, Hercules, CA, USA). All measurement points were done in triplicates in three or more independent experiments.

**Human and Mouse Fibroblast Cultures and Scratch Assay.** Human fibroblast cells and mouse 3T3 fibroblast cells were seeded in DMEM containing 10% of fetal bovine serum
and 1% penicillin—streptomycin. Cells were used between 5 and 25 passages.

Both mouse 3T3 fibroblasts and human fibroblasts were seeded at 100,000 cells per well in 24 well plates, grown to confluency and scratched with a 100 μL pipette and washed with DPBS to remove detached cells. Fresh serum-containing media was added, and cells were treated with (i) FGF (0.3 μM); (ii) MitC (15 μM); and (iii) curcumin (24.4 μM) for 24 h. Cells were then washed once with PBS and fixed with paraformaldehyde (4%) for 15 min at room temperature. Cell nuclei were labeled with Hoechst 33258 (10 μM, 10 min), and fluorescent images of the scratch area were acquired. Fluorescent images of cells were acquired at 20× with a Leica DFC350FX monochrome digital camera connected to a Leica DMI4000B inverted fluorescence microscope. Cell migration was quantified using ImageJ and expressed as fold change in the number of cells in the scratch area over the control.

**DRG Cultures Preparation and Imaging.** *Dissociated DRG Preparation, Live Imaging, Neurite Outgrowth Assay.* L1–L5 DRG neurons were cultured from P5–P7 C57BL/6J mice. The ganglia were dissociated in 0.1% trypsin in 1× HBSS for 45 min at 37 °C followed by gentle mechanical trituration using a fire-polished pipette. Dissociated cells were grown on laminin-coated 96-well plates (Corning Inc., Corning, NY, USA) and in growth media consisting of L-15 (Wisent Bio Products, St. Bruno, QC, Canada) supplemented with vitamins, cofactors, and penicillin—streptomycin.

In preparation for live cell imaging, dissociated DRG were cultured in 50 ng/mL free NGF for 24 h. After 24 h, cells were incubated for 24 h with a 50,000 ng/mL suspension of RhB-labeled polymersomes loaded with NGF or a polymersome-NGF suspension. Fifteen minutes before the end of treatment, the cell nucleus was stained with Hoechst 33342 at 10× with a Leica DFC350FX monochrome digital camera connected to a Leica DMI4000B inverted fluorescence microscope. Neurite outgrowth was quantified as the total area of bright areas (MTT-labeled regions), excluding the explant body, normalized against the length of the explant body in the image, and expressed as percentage control.

**Primary Dissociated Hippocampal Neural Cultures Preparation and Live Imaging.** Primary dissociated hippocampal neural cultures were prepared from P0–P2 postnatal C57/BL6 mice. Hippocampi were dissected and placed in ice-cold 1× HBSS. The tissues were softened in 0.25% trypsin (2.5% Trypsin (Thermo Fisher) diluted 10-fold into 1× HBSS) for 20 min at 37 °C and 0.1% DNase I (Sigma DN25) for 5 min at 37 °C. Tissues were washed twice in plating media (Neurobasal medium (Thermo Fisher) + 2 mM l-glutamine (Thermo Fisher) + 10% fetal bovine serum + 1% penicillin—streptomycin) and mechanically dissociated by triturating with a fire-polished pipette. Cells were plated onto PDL (Sigma)-coated chamber slides in plating media for 24 h and switched to serum-free maintenance media (Neurobasal medium + 2 mM l-glutamine + 2% B27 + 1% penicillin—streptomycin). Neural cultures were feed every 3 days by changing 50% of the maintenance media and were used after 12 days in culture. Cells were treated as described in the corresponding figure and live-imaged using fluorescence microscopy (Leica DMI6000B). All animal uses followed the McGill University animal use guidelines and were approved by the McGill Animal Care Committee.

**Organotypic Hippocampal Culture Preparation, Imaging, and Spine Classification.** Preparation of organotypic hippocampal cultures were extensively described in Gähwiler et al. 82 Cultures were prepared from P6-8 transgenic mice that express Thy-1 driven eGFP in a subpopulation of CA1 neurons. Hippocampi were transversely sliced 400 μm thick and mounted onto glass coverslips using chicken plasma clot (Cocalcol Biologicals; Reamstown, PA, USA). The culture medium consisted of 25% heat-inactivated horse serum, 25% HBSS, and 50% Basal Medium Eagle. Serum-free medium consisted of 67% Basal Medium Eagle and 33% HBSS supplemented with 12.5 mM glucose and 1.25 mM l-glutamine. Cultures were incubated for 3 weeks in a roller drum incubator before experimentation. Cultures were treated with vehicle (PBS, control), empty polymersomes, free BDNF (100 ng/mL), or BDNF-loaded polymersomes. After treatment, cultures were fixed in 4% PFA overnight at 4 °C, washed with PBS, and mounted onto microscope slides. Image stacks of CA1 pyramidal neurons expressing eGFP were taken at Z = 0.2 μm. Images were acquired using an upright LEICA TCS SP2 confocal microscope (Leica Microsystems, Heidelberg, Germany) with an HVX PL APO 63× NA 1.4 oil immersion objective. Image processing and spine classification were performed as previously described. 83

**Statistical Analysis.** Statistical significance was determined by one-way ANOVA followed by Dunnett’s test, independent t-test, or by one sample t-test. Data were analyzed using SYSTAT 10 (SPSS).

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.8b02311.
Methods section for the synthesis of triblock copolymer and the AF₄ methodology; schematic representation for the synthesis of PMOXA–PDMS–PMOXA; H¹ NMR and ¹⁹F NMR charts of PMOXA–PDMS–PMOXA; calibration curve for BSA and curcumin-loaded polymersomes; hydrodynamic radius distribution of polymersomes prepared by the rehydration method determined by DLS; kinetics of the release of curcumin and NGF from PMOXA–PDMS–PMOXA triblock-based polymersomes; AF₄ fractograms of PMOXA–PDMS–PMOXA-based polymersomes loaded with NGF-rhodamine and functionalized with memantine and TZG; AF₄ calibration of NGF alone; electron micrographs of polymersomes; fluorescent micrographs of primary dissociated DRG cultures exposed to RhB-labeled polymersomes containing NGF; and image transformation for measurement of the neurite surface area using ImageJ software (PDF).

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