Exploring the Potential of Benzene-1,3,5-tricarboxamide Supramolecular Polymers as Biomaterials

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ABSTRACT: The fast dynamics occurring in natural processes increases the difficulty of creating biomaterials capable of mimicking Nature. Within synthetic biomaterials, water-soluble supramolecular polymers show great potential in mimicking the dynamic behavior of these natural processes. In particular, benzene-1,3,5-tricarboxamide (BTA)-based supramolecular polymers have shown to be highly dynamic through the exchange of monomers within and between fibers, but their suitability as biomaterials has not been yet explored. Herein we systematically study the interactions of BTA supramolecular polymers bearing either tetraethylene glycol or mannose units at the periphery with different biological entities. When BTA fibers were incubated with bovine serum albumin (BSA), the protein conformation was only affected by the fibers containing tetraethylene glycol at the periphery (BTA-OEG₄). Coarse-grained molecular simulations showed that BSA interacted with BTA-OEG₄ fibers rather than with BTA-OEG₂ monomers that are present in solution or that may exchange out of the fibers. Microscopy studies revealed that, in the presence of BSA, BTA-OEG₄ retained their fiber conformation although their length was slightly shortened. When further incubated with fetal bovine serum (FBS), both long and short fibers were visualized in solution. Nevertheless, in the hydrogel state, the rheological properties were remarkably preserved. Further studies on the cellular compatibility of all the BTA assemblies and mixtures thereof were performed in four different cell lines. A low cytotoxic effect at most concentrations was observed, confirming the suitability of utilizing functional BTA supramolecular polymers as dynamic biomaterials.

INTRODUCTION

Nature is the source of inspiration for the fabrication of fascinating materials.1,2 Although the field of biomaterials has made huge progress in recent years, mimicking Nature remains an extremely challenging task. This is mainly due to the highly dynamic nature of natural systems, which is complex to simulate artificially or synthetically. Noncovalent recognition motifs occurring through hydrogen bonding, π−π stacking, metal chelation, van der Waals, and hydrophobic interactions allow natural systems to continuously evolve, reshape and reorganize in time and space to fulfil specific functions. Thus, biomaterials assembled through these noncovalent interactions have emerged as promising candidates to mimic the unique dynamics of Nature. Supramolecular chemistry offers the essential tools to reach this ambitious goal. As a result, water-compatible supramolecular polymers have been extensively studied as building blocks for the fabrication of dynamic biomaterials.3,4

A plethora of supramolecular polymers assemble into one-dimensional (1D) fibers in water and buffers,5 making them extremely powerful to mimic the abundant number of fibrillar components in natural systems.6 In particular, host–guest interactions have been used to assemble 1D supramolecular polymers by exploiting their hydrophobic character.7,8 Nevertheless, achieving complex modular host–guest biomaterials is challenging due to synthetic limitations. Alternatively, π-conjugates have been chosen as modular supramolecular building blocks, keeping the hydrophobic interactions as driving forces for the assembly. In this case, direct coupling to specific biofunctional moieties is feasible, which facilitates the synthetic challenge.9,10

A further example of fiber-like materials is represented by nature-inspired peptide assemblies that fully mimic 1D fibrillar native entities, containing natural amino acids and thus having intrinsic biofunctionality. Among them, peptide amphiphiles (PA), made of peptide sequences covalently bound onto hydrophobic chains, have shown promising properties as
biofunctional materials. These PA fibers have been used in a number of exciting applications, while at the same time, the fundamental understanding of their structure–property relationship has been crucial to progress.

In order to achieve broader modularity and expand the bioapplications of supramolecular polymers, ureido-pyrimidine (UPy)-based 1D materials have been developed. UPy moieties dimerize via quadruple hydrogen bonding, and the addition of urea groups allows for lateral stacking into fibers held together by a combination of hydrogen bonding, hydrophobic interactions and π-π stacking. In dilute form, UPy fibers have been enriched with cationic moieties to allow intracellular delivery of siRNA, while the incorporation of specific peptides enabled growth factor stabilization.

Another example that involves hydrogen bonding and hydrophobic effects is benzene-1,3,5-tricaboxamide (BTA)-based supramolecular polymers. The increased understanding gathered from fundamental studies on their assembly in water renders this class of materials highly promising for bioapplications. BTAs are known to form μm-long 1D fibers via intermolecular 3-fold hydrogen bonding between the amides and hydrophobic effects and have been proven to be highly dynamic. Additionally, in analogy to covalent polymers, copolymerization has been successfully achieved thus, allowing the modular insertion of multiple functionalities. Peptides, charges, carbohydrates, and DNA have been used to decorate BTA fibers, inducing cellular recognition, allowing intracellular delivery, mimicking the glycocalyx, and recruiting proteins. Furthermore, hydrogels were recently obtained by increasing the concentration of fibers and self-healing characters of these hydrogels make them extremely powerful tools to mimic the extracellular matrix. Nevertheless, their application as biomaterials in, for example, tissue engineering is still limited, also due to the lack of detailed knowledge on how these supramolecular materials behave in physiological environments and, especially, on how they interact with biologically relevant species, that is, proteins and cells.

In this study, we assess the potential of water-compatible BTA derivatives as biomaterials both in the fiber and in the hydrogel state. By applying a fully realistic approach, the stability of different BTA homopolymers (BTA-OEG₄, BTA-OEG₄-Man, and BTA-Man; Scheme 1) and copolymers (BTA-OEG₄-Man/BTA-OEG₄ and BTA-Man/BTA-OEG₄) upon incubation with physiological concentrations of bovine serum albumin (BSA) was investigated. Remarkably, the BTA-s contained their fiber-like structure in protein-rich media, as proven by complementary techniques. Furthermore, several cell lines showed high viability, even upon exposure to high BTA concentrations, providing unprecedented insight into the applicability of BTA-based supramolecular polymers as biomaterials.

## EXPERIMENTAL SECTION

### Materials

- Dulbecco’s Modified Eagle Medium (DMEM), Eagle’s Minimum Essential Medium (MEM), penicillin–streptomycin, trypsin, and fetal bovine serum (FBS) were purchased from Gibco (Life Technologies).
- Phosphate buffer saline (PBS), bovine serum albumin (BSA), urea, 10X DMEM, 10X MEM, and human recombinant insulin were purchased from Sigma-Aldrich. (4,5-Dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium (MTS) solution (CellTiter 96Aqueous One Solution Assay) was purchased from Promega. Cy5-BSA was obtained from ProteinMods.
- BTA-OEG₄, BTA-OEG₄-Man, BTA-Man, and BTA-Cy3 were synthesized and assembled into 1D fibers as previously reported.

### Scheme 1. Chemical Structure of the Water-Compatible BTA Monomers Selected for This Study

BTA-OEG₄ and BTA-OEG₄-Man samples were assembled by weighing the solid material into a glass vial, adding Milli-Q (MQ) water to obtain the desired concentration, stirring the sample at 90 °C for 15 min, and vortexing the sample for 15 s immediately afterward. The samples were then left to equilibrate overnight at room temperature before being used for any measurements. The samples in MEM in DMEM were prepared by first preparing a 1 mM sample of BTA-OEG₄ in MQ water following the standard heating–cooling method. After equilibration for 8 h, the samples were diluted to 250 μM with medium (including 10% FBS) and allowed to equilibrate overnight at 4 °C. BTA-Man was dissolved in methanol to obtain a concentrated stock solution, which was subsequently injected into MQ water. After overnight equilibration, dialysis against MQ water was performed to remove the methanol content. Co-assembled samples were prepared by allowing the individual stocks to cool down to room temperature for 10 min after heating the stocks for 10 min at 80 °C. The 1:1 mixture was prepared by pipetting the correct amounts together. The samples were vortexed for 10 s, heated for another 10 min at 80 °C, vortexed for 10 s, and allowed to equilibrate at room temperature overnight before measuring. Hydrogels were prepared using the same procedure as in the assembly but with an additional step of equilibration in an ice bath after vortexing for 15 s.

### Tryptophan Fluorescence Emission

BSA (20 mg/mL in PBS) was incubated overnight at 37 °C in a 96-well-plate with different concentrations (1.6 to 100 μM in PBS) of BTA-OEG₄ and the copolymers of BTA-OEG₄-Man/BTA-OEG₄ and BTA-Man/BTA-OEG₄. Tryptophan fluorescence emission was measured with a TECAN SPARK Multimode microplate-reader in the wavelength range of 320 to 500 nm after excitation at 280 nm. The experiments were performed in triplicate, and the average of the spectra was calculated. BSA alone was measured at negative control and BSA incubated with urea (9 M) was considered as positive control.

### Förster Resonance Energy Transfer (FRET)

Cy5-BSA (1 μM in PBS) was incubated overnight at 37 °C in a 96-well plate with different concentrations (1–20 μM in PBS) of 5% BTA-Cy3, 95% BTA-OEG₄, 5% BTA-Cy3, 95% BTA-OEG₄-Man/BTA-OEG₄, and 5% BTA-Cy3, 95% BTA-Man/BTA-OEG₄. FRET was measured with a TECAN SPARK Multimode microplate reader in the wavelength range of 560 to 800 nm after excitation at 520 nm. Cy5/Cy3 ratio was calculated after subtraction of the Cy5-BSA fluorescence and normalized for the Cy3 amount in each sample.
Cryogenic Transmission Electron Microscopy (CryoTEM). Imaging was performed on samples with a BTA concentration of 250 μM incubated overnight with 20 mg/mL BSA in PBS. Vitrified films were prepared in a “Vitrobot” instrument (PC controlled vitrification robot, patent applied, Frederik et al. 2002, patent licensed to FEI, Vitrobot Mark III) at 22 °C and at a relative humidity of 100%. In the preparation chamber of the “Vitrobot”, 3 μL samples were applied on lacey grids (LC200-Cu, Electron Microscopy Sciences), which were surface plasma treated just prior to use (Cresington 208 carbon coater operating at 5 mA for 40 s). Excess sample was removed by blotting using filter paper for 4 s at −3 mm, and the thin film thus formed was plunged (acceleration about 3 g) into liquid ethane just above its freezing point. Vitrified films were transferred into the vacuum of a CryoTITAN equipped with a field emission gun that was operated at 300 kV, a postcolumn Gatan Energy filter (97335 Nikon). Images were recorded with an EMCCD camera. Virtrified films were observed in the CryoTITAN microscope at temperatures below −170 °C. Micrographs were taken at low dose conditions, starting at a magnification of 6500 with a defocus setting of −40 μm and at a magnification of 24000 with a defocus setting of −10 μm.

Total Internal Reflection Fluorescence Microscopy (TIRF). TIRF images were acquired with a Nikon N-STORM system. Cy3 was excited using a 561 nm laser. Fluorescence was collected by means of a TIRF images were acquired with a Nikon N-STORM system. Cy3 was excited using a 561 nm laser. Fluorescence was collected by means of a TIRF microscopy instrument (PC controlled vitrification method,47 with a coupling time of 2 ps, while the pressure was kept at 1 bar by means of Parrinello—Rahman barostat,36 with a time constant of 6 ps.

CG-MD Simulations—Mechanism of BSA—BTA Interaction. Metadynamics (MetaD)49 was employed as previously described49 to efficiently explore the configurations and exchange of a BTA-OEG4 monomer across the interface between the fiber and the BSA dimer. The applied bias acted along two collective variables A and B, relative to a preselected monomer M, initially interacting with both the fiber and the protein surface. A was defined as the minimum distance between the M core atoms (the benzene ring of BTA-OEG4) and the backbone of the fiber. B was defined as $v_{fiber} - C_{protein}$, where $v_{fiber}$ was the number of contacts between the core atoms of M and the backbone of the fiber (the other BTA cores) and $C_{protein}$ was the number of contacts between the core of M and the BSA atoms. The bias was constructed by depositing every 1000 CG-MD steps Gaussian kernels of height 0.2 kJ/mol and width of 0.05 (in the A direction) and 0.25 (in the B direction). All MD and MetaD simulations and analyses were performed with GROMACS 2018,50,51 equipped with PLUMED 2.5.2.

Rheology. Rheological measurements were performed on a TA Instruments DHR-3 rheometer (TA Instruments). A hydrogel was deposited onto the rheometer stage. A 20 mm stainless-steel cone plate was used, and the gap height was set to 56 μm for measurement. The temperature was controlled strictly at 37 °C using a water trap for all the measurements, and each measurement was repeated multiple times for reliable data. We monitored the storage modulus $G’$ and loss modulus $G''$ under an applied strain of 0.1% to 100% at a frequency of 1 rad s−1 for the strain sweep.

Cell Culture. HEK293, RAW264.7, and MDA-MB-231 cells were cultured in a medium composed as follows: DMEM, 10% FBS and 1% penicillin-streptomycin. MCF7 cells were cultured in a medium composed as follows: MEM, 10% FBS, 0.01 mg/mL human recombinant insulin, and 1% penicillin-streptomycin. The cell culture was performed under an atmosphere with 5% CO2 at 37 °C. Detachment of the cells from the culture flask was performed by trypsinization for HEK293, MDA-MB-231, and MCF7 and by scraping for RAW264.7.

Cytotoxicity. For the determination of the cytotoxicity of BTAs, cells were seeded in a 96-well cell culture plate at a density of 5 × 104 cells per well for HEK293 and MDA-MB-231, 10 × 104 cells per well for MCF7 and 25 × 103 cells per well for RAW264.7 in complete media (100 μL) and allowed to grow for 24 h at 37 °C in a humidified atmosphere with 5% CO2. Subsequently, the medium was aspirated, and the cells were washed twice with PBS before adding different concentrations of BTAs (from 0.8 to 150 μM). The final concentration of BTAs was obtained by diluting the samples with 10× DMEM and 10× MEM. The samples were incubated for 24 h at 37 °C in a humidified atmosphere with 5% CO2. Control cells were treated with equivalent volumes of the corresponding medium. Afterward, 20 μL of MTS solution (CellTiter 96Aqueous One Solution Assay) was added to each well of a 96-well plate containing cells. The cells were then incubated for 3 h at 37 °C in a humidified atmosphere with 5% CO2, and the absorbance was read at 490 nm. The experiments were run in triplicate. The average of the triplicate was normalized based on the controls.
RESULTS AND DISCUSSION

Interaction of BTA Fibers with BSA Probed by Fluorescence Spectroscopy. Since protein adsorption onto materials is depicted as the first event occurring in any biological system, the interaction of BTA fibers with proteins was first investigated by using bovine serum albumin (BSA) as model protein. The BSA quaternary structure shields two tryptophans in hydrophobic pockets, allowing them to emit fluorescence. Upon protein unfolding, these amino acids get exposed to the surrounding aqueous environment, and their fluorescence quenches. By spectrscopically monitoring this process, interactions between BSA and BTA fibers were investigated.

Different amounts of BTA-OEG₄ fibers, assembled through the standard heating–cooling procedure, were incubated overnight at 37 °C with 20 mg/mL (303 μM) BSA in PBS, which corresponds to its physiological concentration, and the changes in tryptophan fluorescence emission were followed. The fluorescence quenching was overall low and only at the highest concentration of BTA-OEG₄, when the ratio BTA-OEG₄/BSA was close to 1:3, the fluorescence emission spectra overlapped with the one containing urea, which is a known protein denaturing agent (Figure 1a). In order to investigate the

![Figure 1](image1.png)

Figure 1. BSA tryptophan fluorescence emission showed interaction between BTA-OEG₄ and physiological concentration of BSA at 37 °C (a). The introduction of mannose moieties at the fiber periphery reduced this interaction (b, c). FRET experiments (d) showed enhanced interaction between Cy3-labeled BTA fibers and Cy5-BSA increasing the fibers concentrations.

![Figure 2](image2.png)

Figure 2. (a) CryoTEM of BTA-OEG₄ incubated with BSA (scale bar: 50 nm) and (b) TIRF imaging of Cy3-BTA-OEG₄ incubated with BSA (scale bar: 10 μm).
mechanism of this interaction, the same experiment was repeated with more hydrophilic BTA-based supramolecular copolymers. In particular, a physiological concentration of BSA was exposed to different amounts of 1:1 copolymers composed of BTA-OEG4-Man and BTA-OEG4 (BTA-OEG4-Man/BTA-OEG4) and BTA-Man and BTA-OEG4 (BTA-Man/BTA-OEG4). The tryptophan fluorescence showed almost no quenching by these two copolymers (Figure 1b,c), indicating a minor interaction between BSA and BTA copolymers bearing mannose moieties at the periphery. Remarkably, the role of mannose in reducing the BTA interaction with proteins was further confirmed by lower Forster Resonance Energy Transfer (FRET). Both Cy3-labeled BTA-OEG4-Man/BTA-OEG4 (Cy3-BTA-OEG4-Man/BTA-OEG4) and Cy3-labeled BTA-Man/BTA-OEG4 (Cy3-BTA-Man/BTA-OEG4) showed lower FRET when incubated overnight with Cy5-BSA than Cy3-labeled BTA-OEG4 (Cy3-BTA-OEG4) incubated with Cy5-BSA (Figure 1d). These experiments also showed that higher BTA concentrations induced an increase in interactions, corroborating the tryptophan fluorescence emission results.

Since experimentally BSA showed a somewhat stronger interaction with BTA-OEG4 fibers compared to the two mannose copolymers, the interaction and stability of BTA-OEG4 fibers was further unraveled in the presence of BSA.

Consequences of the Interactions between BSA and BTAs Probed by Microscopy. In order to assess whether BTA-OEG4 assemblies were retaining a fiber-like morphology upon incubation with BSA, cryogenic transmission electron microscopy (cryoTEM) and total internal reflection fluorescence microscopy (TIRF) were measured. For cryoTEM imaging, BTA-OEG4 fibers were incubated overnight with BSA at physiological protein concentration in PBS. Even though fluorescence spectroscopy on tryptophan exposure and FRET experiments suggested an interaction of BSA with BTA-OEG4...
fibers, long 1D assemblies were visualized (Figure 2a). Nevertheless, some fibers showed chain ends, suggesting that the interaction with BSA partially affects the BTA-OEG4 microstructure by shortening the fiber length, as seen by cryoTEM imaging of BTA-OEG4 with and without BSA at the same magnification (6.5k) and BTA concentration (250 μM; Figure S4). The BTA-OEG4 fiber-like structure preservation upon incubation with BSA was further confirmed by TIRF microscopy, which showed the presence of micrometer-long fibers (Figure 2b). In some cases, those were slightly shorter compared to Cy3-BTA-OEG4 not incubated with BSA (Figure S5). This result showed the correlation with cryoTEM results and suggested that even though the fibers retained their morphology, their length was partially affected due to the interaction with BSA.

Molecular Simulations Elucidating the Interaction of BTAs with BSA. Two mechanisms were initially hypothesized for the BTA-OEG4/BSA interaction: either BTA-OEG4 monomers were sequestered by BSA and trapped into the protein hydrophobic pockets (i.e., monomer exchange out of the fibers into the protein), or BSA was destabilized by BTA-OEG4 fibers (i.e., interaction of BSA with BTA-OEG4 fibers, rather than monomers). In the first case, the BSA/BTA-OEG4 interaction would be governed by the dynamics of the monomer exchange out of the BTA-OEG4 fibers, which determines how many disassembled monomers were present in the solution. In the second case, the destabilization would involve a macro-molecular interaction between BSA and the whole BTA-OEG4 fibers.

In order to shed light onto the most probable BSA/BTA-OEG4 interaction mechanism, we employed coarse-grained molecular dynamics (CG-MD) simulations. In particular, two in silico experiments were designed to assess the effect on the BSA in the two types of interactions, that is, BSA interaction with disassembled BTA-OEG4 monomers in solution vs BSA interaction with assembled BTA-OEG4 fiber. We thus simulated two cases where one BSA dimer interacts with 100 initially disassembled BTA monomers versus one pre-equilibrated BTA fiber composed of 100 assembled monomers in aqueous solution. The effect of the interaction with BTA-OEG4 on the BSA structure in both cases was then compared.

In general, both CG-MD simulations showed that the protein had a strong interaction with BTA-OEG4. In the first case, the monomers tended to distribute across the BSA surface, targeting specific hydrophobic patches (Figure 3a). In the second situation, the BTA-OEG4 fiber strongly attached to the protein surface and bent surrounding it (Figure 3b). Analysis of the CG-MD trajectories indicated that the number of contacts between BSA and BTA-OEG4 was higher in the case of free/disassembled BTA-OEG4 monomers in solution (Figure 3c). Nevertheless, the BSA 3D structure deviated more from its native conformation when interacting with the BTA-OEG4 fiber (Figure 3d).

Since CG models can be intrinsically limited by the approximations in the CG scheme and by the use of the elastic network applied on the protein model, we cross-checked these comparative simulations in presence and in absence of an elastic network potential (preserving the BSA structure). These additional simulations confirmed the trends observed in the data shown in Figure 3d for both cases. This demonstrated that the BSA/fiber interaction entailed a strong effect on the protein secondary and tertiary structure. This phenomenon, which can
be compatible with protein denaturation, is experimentally corroborated by the similar effect of urea and BTA-OEG₄ seen in the fluorescence spectroscopy. On the other hand, although the BSA interaction with free monomers was observed in the CG-MD (Figure 3c: number of contacts with monomers was even higher than those with the BTA-OEG₄ fiber), the effect on the BSA’s 3D structure was considerably smaller. Altogether, this indicated that the interaction with a full fiber had a stronger destabilization effect on the BSA structure than the interaction with the same number of individual BTA-OEG₄ monomers. Since the concentration of free monomers when the fibers are present is very low, BSA proteins interacted more likely with BTA-OEG₄ fibers than with disassembled BTA-OEG₄ monomers. Thus, these CG-MD simulations suggested that the tryptophan fluorescence quenching and the shortening of the fibers observed experimentally were caused by the interaction of BSA with the BTA-OEG₄ fiber rather than with the monomers. This was further confirmed by thermodynamic data extracted from the CG-MD simulations, which showed a more favorable interaction energy for BSA with BTA-OEG₄ fibers compared with the same number of disassembled BTA-OEG₄ monomers (with a difference in interaction energy of ∼2.2 kcal/mol per monomer, see Table S1).

Additional evidence for stronger interaction of BSA with BTA-OEG₄ fibers rather than with BTA-OEG₄ monomers was obtained by CG metadynamics (CG-MetaD) simulations. Starting from a BTA-OEG₄/BSA bound state (i.e., BTA-OEG₄ fiber bent and bound to the BSA surface), a CG-MetaD simulation was performed where one BTA-OEG₄ monomer at the interface was biased to exchange/diffuse from the BTA-OEG₄ fiber within the protein and back. During the CG-MetaD run, the monomer crossed multiple times back and forth from the fiber into the protein. From this CG-MetaD simulation, we computed the free energy surface (FES) of the monomer interaction between fiber and protein. The FES showed that a BTA-OEG₄ monomer could exist in different states with favorable free energy (see Figure 4, dark free energy minima). In the first case, the monomer could be stacked into the BTA-OEG₄ fiber backbone having negligible interaction with BSA (Figure 4i). Another possibility could see the monomer stacked onto a defect of the BTA-OEG₄ fiber backbone at the interface between BSA and the BTA-OEG₄ fiber and interacting with both (Figure 4ii). The monomer could also be placed at the interface between BSA and the BTA-OEG₄ fiber but adsorbed on the fiber surface (Figure 4iii). In this case, the monomer interaction was stronger with BSA than with the fiber. Finally, the monomer could be sequestered by the protein, having no interaction with the fiber (Figure 4iv).

Stability of BTA-OEG₄ Fibers and Hydrogels in FBS and Cell Media. Since the interaction between BTA-OEG₄ and BSA did not disrupt the fiber morphology, the stability of the BTA fibers was further assessed in the presence of fetal bovine serum (FBS) as a more complex biological medium. For this purpose, Cy3-BTA-OEG₄ fibers were incubated overnight with FBS, and TIRF microscopy was measured (Figure 5a). Compared to Cy3-BTA-OEG₄ alone (Figure S5) and Cy3-BTA-OEG₄ incubated with BSA (Figure 2b), the presence of FBS induced heterogeneity between fibers, as micrometer long assemblies could be visualized, together with smaller aggregates. This indicated that FBS was capable of disrupting some of the fibers into smaller assemblies, but also that long micrometer fibers remained present. The stability of BTA-OEG₄ fibers was further assessed by overnight incubation in DMEM and MEM cell media containing 10% FBS. CryoTEM imaging of these samples showed the presence of 1D supramolecular polymers with visible chain ends in both media that were shorter in length when dispersed in MEM than in DMEM (Figure S6). These results show that even in complex cell culture media, the fiber morphology was retained and only shortened due to interactions with proteins.
All the above measurements were done at relatively low concentrations. As a next step, we investigated how the presence of FBS affects the material properties of BTA-OEG₄. At a concentration of 2 wt %, BTA-OEG₄ forms a hydrogel. This hydrogel was exposed to FBS by adding 100 μL of pure FBS on top of the gel and incubating the system overnight. To assess how far FBS has a negative effect on the hydrogel, the mechanical properties of the hydrogels were investigated by rheological measurements (Figure 5b). Remarkably, both hydrogels showed the same linear behavior and a similar gel-to-sol transition upon 300% strain, indicating that, upon FBS incubation, the hydrogel retained its viscoelastic properties. Furthermore, no significant difference of G’ was observed upon addition of the FBS, indicating that the strength of the hydrogel was hardly affected by the presence of FBS. Even though TIRF imaging showed that FBS affected the length of the BTA fibers, hydrogels exposed to FBS maintained their mechanical properties. This suggests that the interaction of FBS with the fibers is not strong enough to inhibit fiber entanglement as the driving force for the hydrogel formation. These results showed that, although interactions exist at the microscopic scale between serum proteins and BTA-based fibers, they did not have a large impact on the material properties; thus, showing that BTA-based supramolecular biomaterials can be used for bioapplications.

Cytotoxicity Studies of BTA Fibers Using Different Cell Lines. Once the stability of the BTA fibers in the presence of serum proteins was assessed and cell culture media, the cytotoxic effect of different BTA assemblies was evaluated in different cell lines. First, the cytotoxicity of pure BTA-OEG₄, BTA-OEG₄-Man, and BTA-Man was assessed. Whereas BTA-OEG₄ and BTA-Man form micrometer long 1D fibers, BTA-OEG₄-Man forms small micelles instead of 1D fibers. The human embryonic kidney cell line HEK293 was chosen as a common human cell line that does not express mannose receptor on the cell membrane and the breast cancer cell line MDA-MB-231 was chosen as human cell line expressing mannose receptor. The three homoassemblies were incubated for 24 h with the two different cell lines at increasing concentrations of BTAs and the cytotoxicity was assessed using the MTS viability assay. The homoassemblies showed no cytotoxic effect when incubated with HEK293 cells at any of the concentrations tested (Figure 6a). Interestingly, the mannose receptor expressing cell line MDA-MB-231 showed a decrease in cell viability to 60% and 70% when incubated with the highest concentrations of BTA-OEG₄ (100 and 50 μM, respectively), which did not contain mannose (Figure 6b). The two mannose-containing homoassemblies of BTA-OEG₄-Man and BTA-Man did not show any cytotoxic effect at any of the concentrations tested.

To further assess whether the coassemblies of each of the monomers could affect the cell viability, all the combinations of the three monomers were coassembled (BTA-OEG₄-Man/BTA-OEG₄, BTA-Man/BTA-OEG₄, BTA-OEG₄-Man/BTA-Man) and the cytotoxicity was tested. Since only the mannose receptor expressing cell line showed a slight decrease in cell viability upon exposure to a high concentration of BTA-OEG₄, two additional cell lines containing mannose receptor were chosen for the cytotoxicity assay, that is, the human breast cancer cell line MCF-7 and the murine macrophages RAW264.7. All the materials were incubated at different
concentrations with the cells for 24 h and the cell viability was assessed using the MTS assay. In contrast to the viability trend shown with MDA-MB-231 breast cancer cells, none of the BTAs showed a cytotoxic effect against MCF-7 breast cancer cells at any of the concentrations tested (Figure 6c). The three BTA homopolymers incubated with macrophages RAW264.7 showed a slight decrease in cell viability at the highest concentration of materials. This phenomenon was particularly pronounced in the case of BTA-OEG4-Man homopolymers that showed only 40% of cell viability (Figure 6d). Since it is known that spherical nanoparticles are internalized by macrophages to a higher extent compared to other more elongated morphologies, this could be attributed to the mannose-displaying micelle-like morphology of BTA-OEG4-Man. Nevertheless, this cytotoxic effect was improved by coassembly of the BTA monomers with each other, leading to negligible cytotoxicity even at the highest concentrations. Hence, in summary, the BTA assemblies did not show a general cytotoxic trend at any of the concentrations studied.

## CONCLUSION

In order to assess the suitability of water-soluble BTA supramolecular polymers as biomaterials, three functional BTA materials bearing tetraethylene glycol, mannose, or both tetraethylene glycol and mannose at the periphery (BTA-OEG4, BTA-Man, and BTA-OEG4-Man, respectively) were studied in the presence of different biological entities (BSA, FBS, and cells). By following the tryptophan fluorescence of BSA as well as FRET between labeled BTAs and BSA, it could be concluded that only BTA-OEG4 interacted with BSA. CG-MD simulations corroborated this interaction and showed that the BSA conformation was affected by the interaction with BTA-OEG4 fibers rather than with BTA-OEG4 monomers. Additional CG-MetaD simulations showed that the exchange of monomers from the BTA-OEG4 fibers to BSA occurred most likely via a direct contact between the fiber and the BSA surface without the need of a monomer exchange to water as an intermediate step. Gratifyingly, microscopy imaging of BTA-OEG4 in the presence of BSA and FBS showed that the fiber conformation was mostly preserved. Only the length was affected and in the case of FBS, the samples were more polydisperse, but still containing long fibers. Moreover, upon incubation with FBS, BTA-OEG4 hydrogels retained the viscoelastic properties of the materials showing their stability in protein-rich media. However, when a specific binding of fibers to certain proteins should be limited, the oligoethyleneglycol groups should be avoided and carbohydrate units are preferred. BTA fiber’s biocompatibility was further assessed in four different cell lines. Some BTA materials showed a slight cytotoxic effect in specific cell lines (BTA-OEG4 with MDA-MD-231 and BTA-OEG4-Man with RAW264.7), but this phenomenon only happened when incubated at the highest concentrations (100 and 150 μM). No cytotoxicity was observed in any of the other cases. The stability of the functional BTA fibers, as well as their compatibility with different cell lines, make these polymers excellent candidates for biomaterials development. Furthermore, their direct accessibility to modularity paves the way for the fabrication of tailor-made materials for selected bio-applications.

### ASSOCIATED CONTENT

#### Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.biomac.0c00904.

CD spectra of BTA fiber incubated with BSA, cryoTEM images of BTA-OEG4 fibers incubated with and without BSA, TIRF imaging of Cy3-BTA-OEG4 in PBS, and cryoTEM images of BTA-OEG4 fibers in cell culture media (PDF)

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