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Self-Assembled Arginine-Capped Peptide Bolaamphiphile Nanosheets for Cell Culture and Controlled Wettability Surfaces

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¶Supporting Information

ABSTRACT: The spontaneous assembly of a peptide bolaamphiphile in water, namely, RFL4FR (R, arginine; F, phenylalanine; L, leucine) is investigated, along with its novel properties in surface modification and usage as substrates for cell culture. RFL4FR self-assembles into nanosheets through lateral association of the peptide backbone. The L4 sequence is located within the core of the nanosheets, whereas the R moieties are exposed to the water at the surface of the nanosheets. Kinetic assays indicate that the self-assembly is driven by a remarkable two-step process, where a nucleation phase is followed by fast growth of nanosheets with an autocatalysis process. The internal structure of the nanosheets is formed from ultrathin bolaamphiphile monolayers with a crystalline orthorhombic symmetry with cross-β organization. We show that human corneal stromal fibroblast (hCSF) cells can grow on polystyrene films coated with films dried from RFL4FR solutions. For the first time, this type of amphiphilic peptide is used as a substrate to modulate the wettability of solid surfaces for cell culture applications.

INTRODUCTION

The self-assembly of short peptides into ordered nanostructures is widely used as a tool for the development of new materials in biotechnology and related fields.1,2 Peptide-based nanomaterials present both an intrinsic biocompatibility and rich structural diversity because their building blocks are amino acids.3 In particular, peptide amphiphiles (PAs) have been a popular choice to formulate peptide biomaterials because of their ability to self-assemble in aqueous medium and their versatile design, where biofunctionality can be easily introduced through the appropriate linkage of specific amino acid sequences.2

During the last 3 decades, the self-assembly of synthetic PAs has been extensively characterized, and their interaction with plasma membranes has been investigated.4,5 More recently, there has been increasing interest in the development of new peptide materials with bioactive units that are suitable to produce coatings for cell culture6 and antimicrobial surfaces.7,8 Bioinspired moieties, derived from functional domains available in naturally occurring proteins, have often been used to design sequences with enhanced cell attachment and proliferation properties.9,10 The lipopeptide C16-KTTKS, incorporating a pentapeptide segment from human collagen type I, has been found to stimulate collagen production in a concentration-dependent manner in both dermal and corneal fibroblast cultures.11 Also, peptide films containing the fibronectin epitope RGDS (R, arginine; G, glycine; D, aspartic acid; S, serine) show good adhesion properties and can significantly enhance the proliferation of human cornea stromal fibroblasts (hCSF).10,12–14 Arginine-rich sequences are often used in the synthesis of PAs with antimicrobial7 and cell membrane-permeation capabilities.15 In addition, the extensive presence of arginine residues in cell-penetrating oligopeptides such as the transactivator of transcription of HIV-I and the penetratin sequence derived from Drosophila Antennapedia protein also inspires its usage in cell-targetting substrates.16 The exact mechanism by which arginine-based peptides exhibit higher membrane permeation across the cytoplasmic barrier remains elusive. However, it has been proposed that the positive charge might favor association with anionic species in the cell membrane, whereas the planar Y-shaped geometry and the strong H-bond-donating capabilities of the guanidinium groups

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in the R residue could induce negative curvature, resulting in caveolae and membrane disruption.15

The vast majority of reports on self-assembling PAs describe surfactant-like species, with a polar head consisting of a short hydrophilic sequence linked to a longer hydrophobic tail of nonpolar amino acids or lipidated hydrocarbon chains.17,18 An interesting alternative to this common layout is that within so-called bolaamphiphilic peptides. They consist of hydrophilic peptide groups at both ends of a relatively long hydrophobic hydrocarbon chain or peptide sequence. The double-headed design of bolaamphiphilic peptides increases solubility and makes possible the formation of nanostructures with well-defined external hydrophilic surfaces.19 In particular, the higher propensity of bolaamphiphilic peptides to self-assemble into flat layers through lateral association of hydrophobic cores19 makes them good candidates for the formulation of coating substrates.

Although these characteristics make bolaamphiphiles attractive for developing peptide coatings, research on these compounds has been rarely reported in the literature. To the best of our knowledge, there has been no previous work on the self-assembly and cytotoxicity properties of double-headed arginine-based peptides. Here, we present a detailed investigation of the self-assembly and cytotoxicity of the linear octamer RFL4FR (R, arginine; F, phenylalanine; L, leucine). The core of RFL4FR consists of the strongly hydrophobic tetraleucine segment containing aliphatic side chains. 16 The central spacer is flanked on either side by F residues, which favors self-assembly through π−π interactions between neighboring chains, as recently observed.1,2,22 The polar ends of the peptide sequence are made from cationic arginine groups with cell attachment properties, exploited in the creation of substrates for cell culture using RFL4FR. In the following sections, we probe the structure of RFL4FR using X-ray diffraction (XRD), small-angle neutron and X-ray scattering (SANS and SAXS), circular dichroism (CD) spectroscopy, and crioegenic transmission electron microscopy (cryo-TEM). Kinetic data following the time evolution of secondary structure through CD provides unique insights on the self-assembly pathway. The remarkable ability of this peptide to modulate wettability properties on solid substrates is investigated by a number of techniques, including cryogenic transmission electron microscopy (cryo-TEM). The instrument was a cryo-electron microscope, JEOLEMS200FSC, with an acceleration voltage of 300 kV. The device was used in bright-field mode with a final resolution of 20 eV, and images were recorded with a Gatan Ultrascan 4000 CCD camera. Aliquots of 3 μL of solution were cast on copper grids, blotted once for 1 s, and then vitrified (in a FEI Vitrobot device) in a 1:1 mixture of liquid ethane and propane at −180 °C. Grids with vitrified samples were maintained in liquid nitrogen and then cryo-transfered into the microscope.

**X-ray diffraction (XRD).** Patterns from oriented samples were obtained from dried stalks. Droplets of a 5 wt % peptide solution were suspended between the ends of wax-coated capillaries and allowed to dry at room temperature. Stalks were vertically positioned onto a RAXIS IV++-X-ray diffractometer (Rigaku) using a rotating anode generator as the X-ray source. Data were recorded using a Saturn 992 CCD camera at a sample-to-detector distance of 40 mm. Data reduction was performed using Fit2D software (ESRF), and unit cell optimization and simulations were carried out with the program CLEARMON.23

**Synchrotron Radiation Circular Dichroism (SRS).** Data were obtained on module B of the B23 beamline at Diamond Light Source (Didcot, UK). Hellma cuvettes with path lengths of 0.01, 0.2, or 1 mm, depending on concentration, were used to optimize the signal-to-noise ratio, with the absorbance values A ≤ 2. The ring current was 230 mA, and the beamline parameters were as follows: 1 mm slits, 1 mm per step, 1 s per step, and 1 mm bandwidth. Data were averaged over 4 accumulations, and background was subtracted. Raw data were converted into mean residue ellipticity units to allow comparison between different concentrations and path lengths. Kinetic data were obtained using a sealed cuvette to avoid evaporation. The temperature was kept at 20 °C, spectra were obtained every 10 min, and the data collection started only after ultrasonication and thermal equilibration.

**Cryo-Transmission Electron Microscopy (Cryo-TEM).** Imaging was performed as described elsewhere.10 The instrument was a cryo-electron microscope, JEOLEMS200FSC, with an acceleration voltage of 300 kV. The device was used in bright-field mode with a final resolution of 20 eV, and images were recorded with a Gatan Ultrascan 4000 CCD camera. Aliquots of 3 μL of solution were cast on copper grids, blotted once for 1 s, and then vitrified (in a FEI Vitrobot device) in a 1:1 mixture of liquid ethane and propane at −180 °C. Grids with vitrified samples were maintained in liquid nitrogen and then cryo-transfered into the microscope.

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**Fluorescence Assays.** Samples containing between ~10−4 and ~1 wt % peptide were dissolved into solutions containing 1 × 10−5 wt % pyrene. Samples were incubated overnight at room temperature (RT, 20 °C) prior to fluorescence assays. Data were recorded using a Cary Varian Ellipse spectrometer, with excitation wavelength λex = 338 nm and slits at 5 × 5 nm. Fluorescence intensities at 375 nm were plotted as a function of log[peptide concentration], and the critical aggregation concentration (cac) was estimated from the crossover between linear fits corresponding to different emission regimes.

**Fourier Transform Infrared (FTIR).** Spectra were obtained on a Nicolet Nexus spectrometer. Droplets of peptide dissolved in D2O were sandwiched between CaF2 windows with mica spacers of 12 μm. One-hundred and twenty eight accumulations were obtained using a resolution of 4 cm−1. The data were corrected by subtraction of the D2O spectra.

**Peptide Synthesis and Samples Preparation.** Peptides were custom-synthesized by CS Bio (Menlo Park, CA) using TFA as a counterion (catalog no. CS12075, batch N013). Purity was assessed by Fourier Transform Infrared (FTIR). Spectra were obtained on a Nicolet Nexus spectrometer. Droplets of peptide dissolved in D2O were sandwiched between CaF2 windows with mica spacers of 12 μm. One-hundred and twenty eight accumulations were obtained using a resolution of 4 cm−1. The data were corrected by subtraction of the D2O spectra.

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was recorded in the interval 0.01 Å⁻¹ ≤ q ≤ 0.25 Å⁻¹. SANS data have been fitted using the step-model presented in the Supporting Information, which has been implemented in the curve fitting toolbox available in MATLAB. The intermediate angle region was probed on the NIMROD beamline at ISIS. A 1 wt % RFL4FR solution prepared into D₂O was loaded into a silica Hellma cell of 30 × 30 × 1 mm³ and illuminated with a white beam (0.5 Å ≤ λ ≤ 12 Å), with a square cross-section of side 30 mm and incident flight path of 20 m.

**Contact Angle.** Measurements were performed using a GBX-Digidrop instrument. Five microliter water droplets were deposited onto peptide-coated surfaces. Photographs shown in Figures 6A and S2 were taken immediately after casting, and images shown in Figure 7G were obtained ~10 min after water deposition. Room temperature was controlled at 22.5 ± 1 °C, and relative humidity was 60%. Angle measurements were obtained using ImageJ software.

**AFM Imaging.** Atomic force microscopy (AFM) assays were carried out at the Brazilian Nanotechnology National Laboratory (Campinas, SP). A Digital Instruments Nanoscope III instrument was operated in tapping mode. Samples were prepared to reproduce the same conditions used in contact angle measurements. Images were obtained by scanning 512 × 512 pixels covering surfaces between 3 and 10 μm². Image treatment and roughness measurements were performed with the Gwydion package.

**Cell Assays.** Viability and proliferation assays were performed using human corneal stromal fibroblasts (hCSFs). These cells were isolated from postmortem human corneal rings, following epithelium cell depletion. The tissue was finely minced prior to digestion in 2 mg/ml collagenase type-I (Invitrogen, USA) in Dulbecco’s modified Eagle’s medium (DMEM) supplemented 5% fetal bovine serum (FBS) for 5 h under gentle agitation at 37 °C/5% CO₂. Isolated cells were plated onto standard culture plates (Corning, USA) and maintained in DMEM/F12 media (Invitrogen) supplemented with 5% FBS, 1 mM ascorbic acid, and 1% penicillin/streptomycin. Aqueous RFL4FR solutions ranging from 0.1 to 1 wt % peptide were prepared and incubated overnight to ensure the formation of self-assemblies. Peptide films were produced by dropping 50 μL from solutions onto 96-well polystyrene culture plates (Corning, USA) and left to dry overnight at room temperature. Cells were seeded onto the peptide films at approximately 1 × 10⁴ cells per cm² and incubated in DMEM/F12 under sterile conditions at 37 °C/5% CO₂. Fluorescence imaging was performed after 24 h and on day 5 after seeding. Prior to fluorescence assays, cell culture media was removed, wells were gently rinsed three times with sterile phosphate buffered saline (PBS), and a live/dead fluorescent dyes: the membrane-permeable Cyto-Dye (fluorescence in green) and membrane-impermeable propidium iodide (fluorescence in red). Cells were incubated for 15 min with both dyes before again being rinsed three times with PBS. Fluorescence images were registered using an AxioVert 200 (Zeiss) microscope. Green and red fluorescence images were obtained using separate filters and superposed with ImageJ. In the superposed image, live cells display green cytoplasmic staining only, whereas dead cells exhibit a red nuclear stain. Live and dead cells were manually counted. Averages and standard deviations were obtained over triplicates.

**RESULTS**

The critical aggregation concentration (cac) of the RFL4FR bolaamphiphile was assessed using a pyrene fluorescence assay. Peptide solutions containing from 3 × 10⁻⁵ to 1 wt % RFL4FR were prepared using a 1 × 10⁻⁵ wt % pyrene solution as the solvent. The fluorescence emission of the solutions (λₑx = 338 nm) was monitored in the interval 350–450 nm. In Figure 1A, the emission intensity at 373 nm is shown as a function of peptide concentration, revealing a sharp association around a cac ~ 0.085 wt %. The discontinuity in the concentration dependence of the fluorescence intensity indicates the trapping of pyrene molecules within the hydrophobic core of peptide aggregates formed above the critical concentration. The cac found for RFL4FR is relatively low compared to the values obtained for single-headed arginine-rich amphiphiles investigated recently. For instance, peptides A6RGD and A6R, whose hydrophobic tails are composed of alanine residues, show aggregation concentrations around one order of magnitude higher than that reported in Figure 1. The finding is especially interesting considering that the presence of a second polar head is expected to increase solubility of the sequence, and it points to the unique composition of the bolaamphiphile investigated here. Previous studies have noted the lowering of critical concentrations in bolaamphiphile self-assemblies due to the formation of H-bond networks induced by amide linkages along peptide backbones. In the case of RFL4FR, not only amide groups but also guanidinium moieties within arginine residues are likely a source of H-bond donors that contribute toward stabilizing self-assemblies at lower concentrations. A6RGD peptides also exhibit backbones rich in amide linkages, but the lack of a second arginine group decreases its abilities to form H-bond networks and could play a role in its higher cac. In addition to H-bonding between adjacent strands, the entropy gain arising from the hydrophobic effect is also an indispensable driving force of self-assembly at low concentrations. In fact, H-bonding alone is not enough to stabilize well-ordered nanosized structures in aqueous medium due to compensation of enthalpy gain by entropy loss upon H-
bond formation.36,37 Regarding RFL, we presume that the tetrapeptide segment flanked by phenylalanine moieties boosts the amphiphatic nature of the bolaamphiphile. Indeed, the RFL FR bolaamphiphile exhibits an average Kyte–Doolittle33 hydrophathy ∼1.5, whereas, for A4RGD and A4R, these averages are ∼0.3 and ∼0.9, respectively.

The secondary structure of the peptide in solution has been analyzed using FTIR and CD. The amide region of the FTIR spectrum obtained for a 1 wt % solution shows a very sharp peak at 1620 cm⁻¹ (Figure 1B), assigned to β-sheet conformation.33 Figure 1B also shows two bands at 1674 and 1684 cm⁻¹ attributed to TFA counterions bound to R residues34 and to the antiparallel β-sheet conformation, respectively.33 In Figure 1C, CD data from three mixtures below and above the cac show a close relationship between peptide conformation and concentration. The spectrum for the sample prepared at 0.005 wt % RFL 4FR exhibits a minimum below and above the cac show a close relationship between peptide conformation and concentration. The spectrum for the sample prepared at 0.005 wt % RFL 4FR exhibits a minimum around 192 nm and a shallow positive maximum at ∼218 nm, associated with mixed transitions arising from peptide backbone and phenylalanine residues in an unordered conformation.35 At a higher concentration, 0.05 wt % RFL 4FR, the spectrum reveals some features typically found in β-sheet structures. In this case, a small negative peak is noticed at ∼190 nm and a strong maximum is found at ∼200 nm, along with a broad minimum near 220 nm. These features suggest the formation of β-sheet structures in the vicinity of the cac. The β-sheet characteristics are even more evident at 1 wt % peptide, with an intense 200 nm maximum appearing accompanied by a shallow (red-shifted) minimum at ∼220 nm.

CD is a valuable tool to investigate the time evolution of peptide aggregation and protein folding in solution.36,37 Here, we followed the kinetics of evolution of the secondary structure of RFL 4FR to provide insight into the self-assembly pathway of this bolaamphiphile. Figure 2 shows CD data measured for a presumably results from the strong amphiphilicity of the RFL FR sequence. In the following, we assume that the ellipticity at ∼200 nm is proportional to the number of β-sheets in solution. It should be mentioned that other conformational changes not necessarily related to the self-assembly could play a role in the CD spectra. However, using methods described previously,36 useful kinetic information can be obtained from the procedure described below.

Figure 2 indicates that the time evolution of ellipticity at 200 nm follows a sigmoidal profile similar to typical signatures found in a large variety of protein aggregation processes.38 To describe this behavior, we have used the two-step model proposed by Fink and Watzky (F–W).38,40,41 The F–W model describes two self-assembly reactions, A → B and A+ B → 2B, where A represents the monomeric form of the protein and B is the monomer incorporated into an aggregate.40,42 It should be noted that these reactions are composed of a multitude of elementary steps (monomers → dimers → trimers → ...) and that their corresponding constant rates, k₁ and k₂, embody average values for all underlying steps.38 The model comprises a slow nucleation phase, where monomers are converted into aggregated nucleation centers, and a fast growth with primary species absorbing onto the nucleation centers and forming well-developed aggregates. Although the two-step model does not account for underlying molecular mechanisms present in aggregation, it has been successful at providing average rates, which allow comparison between data sets from different proteins.38,41 Also, it should be noted that nanosheets are both a catalyst and a product; thus, the second part of the reaction represents an autocatalysis process.41

Mathematically, the model is expressed in its integrated form by the simple three-parameter equation38,41

\[
[B]_t = [A]_0 - \frac{k_1}{k_2 + [A]_0} \exp(k_1 + k_2[A]_0)t
\]

[1]

[B]ₜ and [A]₀ are proportional to the concentration of aggregates at time t and the initial concentration of protein, respectively.41 Parameters k₁ and k₂ are average rate constants related to characteristic times of nucleation and autocatalytic growth. Equation 1 has been extensively used to fit kinetic data sets from a number of proteins undergoing folding and peptides aggregating into amyloid fibrils, as investigated through a wide variety of physical methods, including CD.43,44,45 In addition, it is one of the simplest nonempirical functions providing average kinetic constants to describe the macroscopic behavior of aggregation.41 We have normalized our data to an initial minimum of zero and used equation 1 to fit the temporal variation of ellipticity. As indicated by the red line in Figure 2, the fit exhibits very good agreement with experimental points, although only three parameters are left free in the model. Parameter [A]₀ has been found to be at 5.56 × 10⁻²⁹ deg·cm²·dmol⁻¹, and here it is associated with the ellipticity (at 200 nm) induced by the self-assembly. The resulting rate constants k₁ = 1.32 ± 0.04 × 10⁻¹¹ h⁻¹ and k₂ = 1.96 ± 0.35 × 10⁻¹¹ mM⁻¹·h⁻¹ are around one order of magnitude lower than those found in other protein analogs and show that the self-assembly of this short bolaamphiphilic peptide is relatively fast compared, for example, to that of the amyloid fibril formation of Aβ segments investigated under similar conditions.36–38 In addition, this relatively quick self-assembly process at a...
concentration close to the cac is consistent with the strong amphiphilicity of RFL\textsubscript{4FR}.

To investigate the morphology of the structures resulting from the self-assembly, cryo-TEM imaging has been performed on vitrified specimens. Figure 3A shows a representative micrograph from a 1 wt \% RFL\textsubscript{4FR} solution, revealing the formation of extensive peptide nanosheets. Additional images can be found in Supporting Information, Figure S1. The formed membranes are apparently highly homogeneous in thickness and exhibit defect-free interfaces with large surface areas. Also, the micrographs show that the sheets are highly flexible, since partially folded edges are clearly identified. These characteristics suggest an interesting potential for using these peptide sheets as arginine-rich flexible, since partially folded edges are clearly identified. These characteristics suggest an interesting potential for using these peptide sheets as arginine-rich layers for coating substrates and subsequent exploration in cell-related applications, as discussed below.

The molecular packing within nanosheets has been probed through XRD experiments performed on dried stalks. In Figure 3B, a partially oriented pattern shows intense diffraction rings, indicating a high degree of ordering within the fibrils. Diffraction data from oriented samples systematically reveal sharp meridional reflections at \(d = 4.8\) Å. This repeat distance has been widely ascribed to separation between hydrogen-bonded \(\beta\)-strands organized into \(\beta\)-sheets either in peptides or protein aggregates.\textsuperscript{2}\textsuperscript{5}\textsuperscript{46} In the case of the RFL\textsubscript{4FR} assemblies, we find ordering of anti-parallel-aligned strands with periodicity \(d = 4.8\) Å and held together by H-bonds between NH and CO groups along the backbones.\textsuperscript{2} This information also agrees with FTIR and CD data shown in Figure 1. In addition to the strong meridional reflection, the XRD pattern also contains peaks corresponding to \(d = 11.7\) Å, which are also found by neutron scattering for samples in solution, as discussed below (Figure 5, inset). This periodicity arises from the intersheet spacing between laterally stacked \(\beta\)-sheets, likely stabilized through H-bonds between guanidinium groups at polar termini or through \(\pi\)-stacking of aromatic rings in phenylalanine's side chain. From the results derived above, we conclude that self-assembly are formed by galleries of strands separated by steps of 4.8 Å along the parallel direction and associated into indefinitely long \(\beta\)-sheets laterally spaced by distances of about 11.7 Å. In the third direction, i.e., perpendicular to the plane of nanosheets, the size is determined by the molecular length of RFL\textsubscript{4FR}. Closer to the beam stop, two very strong peaks are found that correspond to spacings \(d = 33.0\) Å, consistent with the length of a octapeptide chain (8 \(\times\) 3.5 Å = 28 Å) plus TFA counterions adsorbed onto arginine groups.\textsuperscript{26} These reflections appear along the equator of the figure, oriented perpendicularly to the diffraction from the \(\beta\)-strand spacing. This configuration confirms that the sheets are organized into a cross-\(\beta\) structure\textsuperscript{26} self-assembled through lateral association between RFL\textsubscript{4FR} chains.\textsuperscript{46} In addition to the strong diffraction peaks mentioned above, other weaker spots are observed at \(d = 16.9\) and 3.9 Å. To index this set of Bragg reflections, we have used the unit cell optimization module available in the program CLEARER.\textsuperscript{23} Our data have been indexed according to an orthorhombic unit cell with lattice parameters \(a = 32.9\) Å, \(b = 11.4\) Å, and \(c = 9.6\) Å. To provide a comparison between calculated and experimental data, we have simulated the theoretical diffraction patterns using the parameters from the indexed unit cell. The result is exhibited in the inset of Figure 3B along with the corresponding Miller indices. The simulated pattern accurately describes the major features of the experimental diffractogram, in particular the positions of the Bragg rings are predicted with good precision (see table in Figure 3B, inset). The simulated pattern also exhibits other reflections that are not observed experimentally.

The reason for this is related to limitations of the simulation model, which reproduces neither the exact arrangement of side chains nor the crystallite packing.\textsuperscript{46} In situ investigations have been performed using synchrotron small-angle X-ray scattering (SAXS) and small-angle neutron scattering (SANS). In Figure 4A, SAXS curves obtained for solutions between 0.1 and 1 wt \% RFL\textsubscript{4FR} are exhibited. Across the concentration range, the SAXS profiles are characterized by an intensity decay proportional to \(q^{-2}\) at very low scattering angles, consistent with flat structures\textsuperscript{49} and thus in agreement with the morphology revealed by cryo-TEM. Therefore, peptide sheets similar to those observed in Figure 3A are likely present also at lower concentrations. To extract quantitative information, the data was fitted using the bilayer form factor detailed in the Supporting Information, eq S2, and included in the SASFit program. The resulting fits (red lines in Figure 4) show good agreement with the experimental data. The obtained parameters reveal effective membrane thicknesses ranging from 25 to 29.4 Å (see Table S1, Supporting Information), indicating
that the peptide sheets are made up of bolaamphiphilic monolayers since this thickness is close to the estimated octamer length. It should be noted that, here, the longitudinal size of the peptides appears to be lower than that one revealed by XRD assays because TFA counterions are released from peptide interfaces in solution.

The fitting procedure provides information in addition to layer thickness. Direct-space reconstructions of electron density profiles also have been retrieved by substituting the fitted parameters into eq S4, Supporting Information. In Figure 4B, it can be seen that the electron density profiles are dominated by two positive Gaussians, associated with the arginine groups at both termini of the bolaamphiphilic sequence. Interestingly, the central hydrophobic portion is also represented by positive (or neutral) electron density. The electron density for the nonpolar segments is different from profiles observed for other surfactant bilayers, which typically exhibit lower electron densities compared to those of solvent.30,51 The reason for this unusual behavior is presumably related to the chemical structure of the hydrophobic core in RFL4FR, which is made up of amino acid moieties rather than just alkyl chains, as is the case for conventional surfactants and lipids. The extensive presence of side chains and O and N atoms along the backbone leads to higher electron densities in comparison with those of hydrocarbon tails found in lipidated surfactants. In addition, as revealed by XRD experiments, RFL4FR chains are organized into a partially crystalline structure; thus, the molecules are closely packed into organized structures.

Neutron scattering provides independent information on the structure of peptide nanosheets since the contrast is related to scattering length densities (SLD) of nuclei in the sample. In Figure 5, SANS data from a 1 wt % RFL4FR sample show the same q−2 descent found in the SAXS curves (Figure 4). Due to q-range constraints and the lower resolution of the SANS measurements, we have implemented the step-model proposed by Nallet et al.52 and expressed in eq S5 in MatLab to describe the data with a minimal number of parameters. The fit exhibits excellent agreement with the experimental data, highlighting the robustness of the procedure. The fitted value \( \delta = 21.7 \pm 3.0 \) Å for the thickness of the hydrophobic core is compatible with the membrane thickness revealed by SAXS, within uncertainties. The cross-section SLD profile has been obtained from eq S6 and is reproduced in Figure 5 (inset). Features arising from the structure factor of peptide chains organized within nanosheets appear in the wide-angle region. In Figure 5, inset, we observe a remarkable Bragg peak at \( q = 0.53 \) Å, arising from a repeat distance \( d = 11.9 \) Å. This periodicity is consistent with the separation between \( \beta \)-sheets found in XRD assays, thus showing that the \( \beta \)-sheet stacking within the supramolecular structure is also present in solution and is not merely a consequence of drying during the production of stalks for XRD assays.

The solutions studied above have been used to modify solid surfaces and modulate their wettability properties. Samples have been prepared just by casting drops from RFL4FR solutions onto the substrates and left to dry overnight. These modified surfaces have been examined through optical microscopy, and the resulting films have been found to be very stable, remaining adsorbed onto the substrates after several weeks or upon mechanical stressing induced by gentle twisting of the surfaces. Contact angle measurements from films obtained from a 0.3 wt % solution are shown in Figure 6A, corresponding to substrates with different degrees of hydrophobicity, a highly wettable (freshly cleaved mica) and quasi-hydrophobic surface (polystyrene, PS). Although the technique is not able to provide a detailed picture of peptide organization on the surfaces, it is very useful for obtaining insights into the average properties.53 On mica, the contact angle increases from 0° to 72°, showing a dramatic growth of hydrophobicity upon peptide modification. On the other hand, a significant decrease is noticed on PS, from 86° to 58°. On glass surfaces, which exhibit intermediate hydrophilicity, we observed the lowest change from 38° to 45° (Supporting Information, Figure S2).

To get deeper insights into the nature of surface modification by peptides, we have performed detailed AFM measurements. Figure 6B shows topology images of samples on mica and PS substrates. AFM data reveal that peptide deposition produces remarkable morphological changes and that coatings with thicknesses of a few hundred nanometers appear to be physically adsorbed on both substrates. Moreover, coatings
are homogeneously distributed across the surfaces, and corrugations are restricted to the nanometer scale, indicating the smoothness of peptide coverage. To quantify surface roughness and provide an index for comparison between different films, we calculated the roughness parameter $R_a$, which is defined as the average of absolute vertical deviations from the mean plane (see Supporting Information for further details). These measurements reveal that roughness is systematically

Figure 6. (A) Water contact angles from mica and polystyrene substrates modified with RFL4FR films. (B) AFM images showing finer topological details and revealing nanoscale roughness of the surfaces ($3 \times 3 \, \mu m^2$).

Figure 7. (A–E) AFM images showing the topology of RFL4FR coatings deposited onto polystyrene surfaces ($10 \times 10 \, \mu m^2$). Films were dried from solutions at concentrations indicated. (F) Surface roughness, $R_a$, plotted as a function of peptide concentration. Red line: a guide for the eye showing the exponential increase of $R_a$ with increasing concentration. (G) Water droplets onto coated PS surfaces photographed 10 min after casting.
higher on peptide coatings when compared to that of bare substrates, in agreement with the formation of thicker coatings. Accordingly, freshly cleaved mica exhibits a low roughness parameter $R_a = 0.16$ nm, whereas peptide films on this same substrate show $R_a = 14$ nm. On bare and peptide-coated PS, respectively, these values are found to be $R_a = 12$ and 24 nm. In addition to modulation of roughness induced by peptides, analysis of the roughness parameter reveals interesting insights into the influence of substrate composition on wettability. When we compare roughness and contact angles exhibited in Figure 6, we observe that uncoated PS and peptide-coated mica possess very similar morphologies and exhibit almost the same $R_a$ values; however, the water hydrophobicity of these materials is rather different, and they possess distinct wettability behaviors. Therefore, in this case, we can infer that the chemical composition of substrates also plays an important role in macroscopic wettability. We hypothesize that the highly hydrophilic and charged nature of the mica surface likely favors adhesion of arginine groups and should assist exposure of nonpolar sites on the top, resulting in stronger hydrophobic behavior. On the other hand, when PS is used as a substrate, the extensive presence of nonpolar clusters could favor polar arginine groups facing upward and increasing polarity at the interface with water.

Previous studies have suggested that both chemical composition and surface morphology contribute toward wettability properties on polymeric substrates used for cell attachment. To further characterize the relationship between morphology and wettability, we performed AFM assays on coatings on PS surfaces prepared from solutions in the range 0.1–1 wt %. Topology images associated with these assays are shown in Figure 7A–E, and they show that morphology is strongly influenced by concentration. Also, although all samples investigated present homogeneous surfaces, data reveal that corrugations become more pronounced when higher amounts of peptide are present. Roughness parameters $R_a$ have been determined from representative $10 \times 10 \mu m^2$ surfaces and plotted as a function of peptide concentration in Figure 7F. Interestingly, the roughness increases monotonically upon concentration and exhibits exponential behavior until reaching $R_a \sim 85$ nm at 1 wt %.

Roughness on these surfaces has been correlated to wettability through water contact angle experiments. On films containing higher peptide amounts, droplets appear to be completely spread over the surface after an elapsed time of $\sim 10$ min (see Figure 7G), indicating greater wettability possibly due to roughness on these surfaces.

Cell assays have been performed to assess the biocompatibility of RFL4FR. In Figure 8A, digital images obtained by phase-contrast microscopy show hCSFs in culture with good adhesion to peptide coatings obtained from solutions containing 0.1 and 0.3 wt % RFL4FR. At these concentrations, hCSF cells maintained a typically fibroblastic morphology similar to those observed under standard culture conditions and comparable confluence to controls after 5 days in culture, indicating cell functionality. Coatings obtained from more concentrated peptide solutions show reduced cellular adherence and observably rounded, phase-bright cell bodies, indicating nonviable cells. These results were supported by the fluorescent live/dead cell staining (Figure 8B), where only red propidium iodide-stained nuclei (dead cells) were seen for the 0.5 wt % RFL4FR solution. The biocompatibility of RFL4FR...
was quantified by plotting the average cell number density and percentage of viable of attached hCSFs as a function of peptide concentration at 24 h and 5 days postseeding (Figure 8C,D). ANOVA and subsequent post hoc testing shows that at day 1 there is a significant difference between films at 0.1 wt % and the control, whereas at day 5, there is no significant difference between these two conditions and the number of cells per cm² is even higher on films at 0.1 wt % (see data for day 5 in Figure 8C). It is clearly shown that both cell number density and percentage viability decline with increasing concentration of peptide in the films. This is most evident during the first day of incubation, indicating poor initial cell attachment at higher concentrations. For instance, on coatings from solutions containing 0.3 wt % RFL4FR and above, the number of viable cells is seen to be below 50% of the total number of seeded cells. This increase in cell death is consistent with the higher availability of arginine groups on the substrates, which likely leads to disruption of cell membranes, as reported elsewhere.15 However, once left to grow for 5 days, cell density increases on coatings up to 0.5 wt %, with no significant differences between the control and 0.1 wt %. The proportion of viable cells also increases accordingly. This suggests that the RFL4FR has, indeed, formed a coating even at low concentrations and that cells do not immediately bind to this substrate (as they do to optimized tissue culture plastic; control), but they do require a few days to produce their own supporting extracellular matrix; thus, the coating at low concentrations is not toxic to the fibroblastic cells. However, at higher concentrations, i.e., 0.5% and above, the coating could be considered to be toxic, as a significant difference in cell density was observed even after 5 days. The low attachment of cells on day 1 could be associated with the wettability of peptide-modified surfaces. These findings agree with contact-angle images of RFL4FR layers dried from solutions with concentrations above 0.5 wt % (Figure 7G), which show water droplets spread over large areas, suggesting growth of wettability with increasing peptide concentration in the films. In this case, we propose that permeation of water molecules across the coatings is enhanced at higher concentrations and presumably contributes to the detachment of lipid-rich plasma membranes. On the other hand, peptide films prepared from lower concentration solutions show moderate hydrophobicity and seem to be more suitable for cell attachment. This, associated with the lower toxicity of peptides at low concentrations, could enable the formation of the supporting extracellular matrix by the cells.

■ CONCLUSIONS
We have examined, for the first time, the self-assembly, secondary structure, wetting properties, and cytocompatibility of an arginine-rich bolaamphiphilic peptide. This peptide has been found to form nanosheets in solution, likely through lateral association between peptide chains, with arginine moieties exposed on both interfaces. The estimated value for critical concentration, cac ∼ 0.085 wt %, is lower than that for many other PAs that have been explored for use in coating materials, and this represents a potential advantage in terms of cytocompatibility. The temporal evolution of self-assembly has been investigated through CD assays, and data have been successfully fitted according to the Finke–Watzky model. This reveals that the self-assembly of RFL4FR peptide is driven by a two-step process, where a nucleation phase is followed by fast autocatalytic growth. This is the first time, to our knowledge, that this type of mechanism has been observed for an arginine-rich oligo-peptide or a bolaamphiphilic peptide. The characteristic rates revealed from these kinetic assays show that time scales for aggregation are on the order of hours, which ensures relatively fast assembly of nanosheets. Secondary structure is dominated by β-sheet conformations, where peptide backbones form paired β-strings with antiparallel orientation. The crystalline structure is characterized by an orthorhombic symmetry with crossed antiparallel strands, at repeat distances of 4.8 Å, forming galleries separated by 11.7 Å running perpendicular to the plane of nanosheets.

Small-angle scattering data shows that the nanosheets are built from bolaamphiphile monolayers. Very good agreement has been found between SAXS and SANS data fitted with a minimum number of free parameters. The electron density profiles of these structures were described by fitting SAXS data to a three Gaussian layer form factor, and they revealed characteristics that are not usual for amphiphilic membranes. Unlike traditional electron density profiles for lipid bilayers or lipilated peptides, the hydrophobic core of RFL4FR nanosheets is characterized by a positive contrast, with electron density values above that for the aqueous solvent. This behavior is a direct consequence of the unique design and composition of the peptide bolaamphiphile, where side chains are organized into highly packed structures in the core of the membranes, which produce higher electron densities.

A unique ability to use the bolaamphiphilic peptide to modulate the wettability of different surfaces has been demonstrated by contact angle measurements and deeply characterized through high-resolution AFM analyses. Quantitative determinations on the surface roughness have shown a close relationship between wettability and surface morphology; in addition, chemical interaction between peptide films and the supporting substrate also has been identified. Thus, the macroscopic wettability found in these peptide coatings arises from a combination of physical and chemical characteristics of the assemblies. Our data suggest that adhesion and peptide concentration in the coatings are correlated and that dried films obtained from more concentrated solutions, i.e., above 0.5 wt %, possess poor attachment properties regarding cells. Also, if the peptide content in the film is too high, then cell viability is compromised. These findings are most likely related to the presence of arginine groups in the substrates, since the abundance of cationic groups in the surfaces should lead to higher polarity of the surfaces and thus greater potential to disrupt lipid membranes.15 No significant differences in cell morphology or viability were observed between the control and films at 0.1 wt % after 5 days in culture, indicating that coatings at this concentration did not introduce detectable cytotoxicity and that cells remain healthy. These data suggest that coatings obtained from 0.1 wt % RFL4FR are safe and suitable to interface with living matter.

■ ASSOCIATED CONTENT

Supporting Information
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SAXS and SANS modeling, fitting parameters, and additional contact angle measurements and cryo-TEM images (PDF).
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Notes
The authors declare no competing financial interest.

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