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Native collagen: electrospinning of pure, cross-linker free self-supported membrane

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ABSTRACT. Rebuilding biological environments is crucial when facing the challenges of fundamental and biomedical research. Thus, preserving the native state of biomolecules is essential. We use electrospinning (ES), which is an extremely promising method for the preparation of fibrillar membranes to mimic the ECM of native tissues. Here, we report for the
first time (1) the ES of pure and native collagen into a self-supported membrane in absence of
crosslinker and polymer support, (2) the preservation of the membrane integrity in hydrated media
in absence of crosslinker and (3) the preservation of the native molecular structure and recovery
of the hierarchical assembly of collagen. We use a multiscale approach to characterize collagen
native structure at the molecular level using circular dichroism, and to investigate collagen
hierarchical organization within the self-supported membrane using a combination of multiphoton
and electron microscopies. Finally, we show that the membranes are perfectly suited for cell
adhesion and spreading, making them very promising candidates for the development of
biomaterials and finding applications in biomedical research.

KEYWORDS. Collagen; electrospinning; hierarchical organization; extra-cellular matrix; tissue
engineering; second harmonic generation microscopy

INTRODUCTION. Increasing the biological relevance of *in vitro* models is an ever-increasing
goal in biological and medical research. This implies preserving the native state of living systems,
i.e. their native conformation, multi-scale organization, molecular recognition properties.
Mimicking the structure of the extracellular matrix (ECM) is of primary importance to provide the
adequate tridimensional (3D) environment and biological cues for cell attachment, proliferation
and function. This key 3D organization mainly arises from the hierarchical structure of type I
collagen, the major component of ECM. Collagen triple helices self-assemble into fibrils of 20 to
500 nm in diameter. It is therefore crucial to preserve its native state when processing biologically
relevant scaffolds. Electrospinning (ES) has become increasingly popular as a cost effective
technology to produce nanofiber scaffolds for tissue engineering, providing self-supported
membranes with controlled porosity.\textsuperscript{1,2} ES allows to produce fibers with tunable diameter (smaller
than 1 μm) and orientation. Biopolymers are the most relevant candidates to be electrospun. However, ES of collagen remains challenging. Collagen solutions do not have the viscoelastic response necessary for jet stabilization partly due to the stability of collagen tertiary structure.\textsuperscript{3,4} As a result, denaturing fluoroalcohols are often used, such as 1,1,1,3,3,3-hexafluoro-2-propanol\textsuperscript{5-10} and 2,2,2-trifluoroethanol,\textsuperscript{8} allowing stretching and rearrangement of collagen. Since then, the preservation of the native state of collagen has continuously been a subject of debate.\textsuperscript{11,12} Several works evidence that collagen completely unfolds in fluorinated solvents. Efforts have been made to find alternative solvents, such as phosphate buffer saline (PBS)/ethanol\textsuperscript{13} or acetic acid/DMSO\textsuperscript{14} mixtures. All of them involve the use of chemical crosslinkers (glutaraldehyde\textsuperscript{5,15} or carbodiimide\textsuperscript{13,14,16,17,18}) to prevent the membrane from falling apart in hydrated environment. However, the use of crosslinkers alters the fibrillogenesis of collagen and the biocompatibility of the materials.\textsuperscript{19} Original approaches have been developed using a polymer support, where collagen is electrospun together with a synthetic polymer (polycaprolactone,\textsuperscript{7,8,20,21} polyethylene oxide,\textsuperscript{18,22,23} polyvinyl-pyrrolidone\textsuperscript{24}) eventually washed out after crosslinking, or with a biopolymer\textsuperscript{25,26} including elastin,\textsuperscript{18} chitosan,\textsuperscript{9} or gelatin.\textsuperscript{27}

In this work, we report for the first time the ES of pure collagen into a self-supported membrane in absence of polymer support (Fig. 1A). The key is to use an aqueous acidic/alcoholic solvent mixture. We also stabilize the membrane structure for further applications in biological hydrated environment in absence of crosslinkers by inducing collagen pre-fibrillation in basic conditions. Since the hierarchical structure of collagen is crucial, we present a multiscale characterization approach to probe the material from its molecular- up to its micro-scale. We claim this approach to be a key step for collagen-based material characterization. It combines highly specific and biologically adapted methods including multiphoton microscopy based on second
harmonic generation (SHG), field emission scanning electron microscopy in a cryo mode (Cryo-FE-SEM) and synchrotron radiation-circular dichroism (SRCD).

Mastering native collagen electrospun membranes is the base for building more advanced biomaterials and use them as a support to study even further the environment of living cells. As an example we show the processing of hybrid biomaterials and the use of pure collagen membrane into microfabricated chips for biological microfluidics.

**Figure 1.** (A) Scheme of collagen ES and membrane processing. (B) Photo, (C,D) SEM images of a self-supported membrane ([collagen] = 1.7 mg.mL$^{-1}$ in 30 mM HCl:EtOH (25:75 v/v)).

**EXPERIMENTAL SECTION.**

**Collagen extraction and purification.** Type I collagen was extracted and purified from rat tail tendons as previously described by substituting 500 mM acetic acid with 3 mM hydrochloric acid.$^{28,29}$ Collagen purity was assessed by electrophoresis and its concentration estimated by
hydroxyproline titration. All other chemicals were purchased and used as received. Water was purified with a Direct-Q system (Millipore Co.).

**Collagen electrospinning.** 1.2 mL of the solution was electrospun for 4 hours at ca. 12 kV with a flow rate of 0.005 mL.min\(^{-1}\) in a home-made chamber flushed with dry air (ca. 10% humidity) at room temperature (\(\sim 20^\circ\)C). Humidity is a key parameter to be controlled to ensure the good electrospinning of collagen and the reproducibility of the experiment. For this reason, we used synthetic dry air (Air Liquid). The counter electrode was a copper tape (2.5 * 1.5 cm) fixed on a Teflon support. The distance between the metal needle and the counter electrode was 9 cm. After processing, the electrospun membrane was detached from the copper support, and a homogeneous self-standing membrane was obtained. Casted collagen membranes were prepared as control and obtained by pouring 1.2 mL of a collagen solution at a concentration of 1.7 mg.ml\(^{-1}\) in 30 mM HCl and 75% vol. ethanol into a Teflon mold. The solvent was allowed to evaporate overnight under ambient conditions.

**Pre-fibrillation of electrospun membranes by ammonia exposure.** 4 mL of ammonium hydroxide solution (25%, Carlo Erba) in a glass beaker were placed in a desiccator (153*191 mm). After 20 min exposure to ammonia vapors, collagen membranes were rinsed three times a day for 10 days in culture medium to remove all ammonia residues before cell proliferation testing.

**Synchrotron-radiation circular dichroism.** 25 \(\mu\)L of collagen solutions in 3 mM HCl were loaded in quartz cells (Hellma) with a path length of 0.01 cm. The measurements were performed on the DISCO Beamline at SOLEIL synchrotron (Saint Aubin, France).\(^{31,32}\) Melting temperatures were obtained by increasing temperature from 20 to 74°C with steps of 3°C and an equilibration time of 5 min. Raw spectra were acquired from 320 to 170 nm with a 1 nm spectral resolution.
Spectra were treated with the CDTool software. They are the average of three spectra. A background (HCl 3 mM acquired in the same conditions) was subtracted from them. Intensity calibration was obtained with a CSA sample.

**Second harmonic generation / 2-photon excited fluorescence.** We used a custom-built laser-scanning multiphoton microscope and recorded Second Harmonic Generation (SHG) and 2-photon excited fluorescence (2PEF) images in parallel as previously described. Excitation was provided by a femtosecond titanium–sapphire laser (Mai-Tai, Spectra-Physics) tuned to 860 nm, scanned in the XY directions using galvanometric mirrors and focused using a 25× objective lens (XLPLN25XWMP2, Olympus), with a resolution of typically 0.4 μm (lateral) × 1.7 μm (axial). We used either circular polarization in order to image all structures independently of their orientation in the image plane, or a set of linear polarizations with different orientations in order to perform polarization-resolved measurements. All polarization-resolved images were acquired at 36 excitation angles θ regularly spaced between 0° and 360°, using 200 kHz acquisition rate and 300×300 nm² pixel size. Power at the sample was typically 12 mW for dry and stabilized membranes, and 2 mW for the hydrated samples. Epi-detection was used for dry and stabilized membranes and 2x2 binning was applied before pSHG processing. Transdetection was used for hydrated sample, with no binning. pSHG images were processed as already described to provide two images: (i) the average of all images acquired with the set of linear polarizations, that is similar to an SHG image acquired with circular polarization; (ii) a map of the main orientation of collagen in the image plane. The latter pSHG image is displayed using the HSV Look-up-table, where H is the orientation displayed in the insert, and V is the brightness, which is set to 1 if the pSHG processing is satisfactory (R2>0.5). Three areas were observed on each sample to check for the sample structure homogeneity. Each sample was investigated in triplicate.
SEM observations. SEM imaging was performed using a Variable Pressure Hitachi S-3400N working at an accelerating voltage of 10 kV, using an in lens secondary electron (SE) detector, with a working distance at / around 4.2 mm. Self-supported membranes were deposited on carbon-tape coated aluminum pads. Samples were coated with a 10 nm gold layer before observations.

Cryo-FE-SEM observations. Sample preparation and observations were performed at the Electron Microscopy Facility (EMF) of the Institut de Biologie Paris-Seine (Sorbonne University, Paris). Prior to observations, samples were sandwiched between two cupules then cryo fixed by immersion in liquid ethane using a cryofixation station (CPC, Leica). Freeze-fracture was processed at −150°C using a high vacuum station (ACE 600, Leica). FE-SEM observations were performed at −120°C at low voltage (0.790 kV) using an in lens secondary electron (SE) detector, with a 20 µm objective aperture diameter and a working distance at / around 2.5 mm (GeminiSEM 500, Zeiss; Cryo station VCT100, Leica). A sublimation step at −90°C was applied for 15 min in the SEM chamber to remove ice. Scan speed and line or drift compensation integrations were adjusted during observations.

Cell culture experiments. Normal Human Dermal Fibroblasts (NHDF) were cultured in complete cell culture medium: Dulbecco’s Modified Eagle’s Medium (DMEM) supplemented with 10% fetal serum, 100 U.mL⁻¹ penicillin, 100 µg.mL⁻¹ streptomycin, 0.25 µg.mL⁻¹ Fungizone, and glutamax. Tissue culture flasks (75 cm²) were kept at 37°C in a 95% air : 5% CO₂ atmosphere. Before confluence, cells were removed from culture flasks by treatment with 0.1% trypsin and 0.02% EDTA. Cells were rinsed and suspended in complete culture medium before use.

After pre-fibrillation of the collagen membranes (either electrospun or casted, see below section S3), NHDF were cultured for 24 hours on the surface of membranes at a cell seeding density of 2
Cells were incubated at 37°C in a 95% air : 5% CO₂ atmosphere. Cell culture experiments were performed three times in triplicate (n=9).

**Long term cytotoxicity of collagen membranes on NHDF.** Collagen membranes (casted, n = 3, or electrospun, n = 3) were set into a 48 well plate. Then, 100,000 NHDF were seeded per well. Cells were incubated for 7 days in contact with membranes in 200 µL of colorless DMEM complete medium at 37°C in a 95% air - 5% CO₂ atmosphere. Cell metabolic activity was monitored at day 7 using Alamar Blue assay. For this purpose, fibroblasts were incubated with 300 µL of a rezazurin solution at 0.1% (w/v) for 4 hours. The supernatant in each well was then collected, diluted with 700 µL of fresh colorless medium and the absorbance measured at λ = 570 nm and λ = 600 nm. The percentage of rezazurin reduction was calculated following the formula provided by the supplier. Cell metabolic activity of each well was compared to control samples, i.e. cells cultivated without membranes. The arbitrary value 100% was given to control samples.

**Fluorescence microscopy (evaluation of cytocompatibility).** Collagen membranes were fixed with 4% paraformaldehyde in PBS for 1 hour, rinsed three times with PBS for 5 min each and incubated with PBS 0.2% Triton for 20 min. They were then incubated with Alexa Fluor 488 Phalloidin (Molecular Probe®) diluted 1/200 (v/v) in PBS in a dark chamber for 45 min. After rinsing three times for 5 min each in PBS, the samples were incubated with DAPI (Molecular Probe®) diluted 1/50,000 in PBS in a dark chamber for 10 min. Finally, the membranes were washed three times with PBS and mounted between slides and coverslips with the antifading solution AF3 (Biovalley) for fluorescence microscopy (ZEISS Axioplan microscope). Fibroblasts were observed at magnification ×100 and ×400.

RESULTS AND DISCUSSION.
A self-supported membrane

Electrospinning is particularly relevant for the design of 3D fibrillary scaffold. In the ES process, high voltage is applied to a polymer solution that is ejected through a needle towards a grounded collector as dried fibers. Upon voltage application, the liquid droplet deforms into a cone, the “Taylor cone”, and emits a charged jet of liquid. Because of the low charge of collagen, obtaining a self-supported membrane is not an easy task. It requires adjusting many parameters to get a stable jet and ensure nanofiber formation (Fig. S1 and S2).\(^3\) Solvent is an important factor for biopolymer stability and stretching during ES and for the drying of the mesh on the collector. Collagen was dissolved in a mixture of hydrochloric acid (3 mM HCl, pH 2.5) and ethanol (75% vol) with a final concentration of 1.7 mg.mL\(^{-1}\). In those carefully adjusted conditions, a self-supported membrane was obtained with a thickness of 40 µm exhibiting randomly oriented and interconnected fibrils with diameter in the nanometer range (58 ± 10 nm) (Fig. 1 B-D, and Fig. S2).

The stabilization of the membrane structure in a hydrated form is crucial for biomedical applications. When immersing the electrospun collagen in cell culture medium, the membrane falls apart. To tackle this issue, works reported in the literature often use chemical cross-linkers that can be detrimental for cell survival, while hampering collagen hierarchical self-assembly. Alternative strategies have to be developed to strengthen the hierarchical structure of collagen. To this aim, we exposed ES membranes to ammonia vapors, to take advantage of the increase in pH to stabilize collagen fibrils. This strategy is known to trigger fibrillogenesis and gelling in collagen solution\(^36\)–\(^38\) but is here adapted to dry materials. After a 20 minutes exposure of the dry membrane, the macroscopic structure of membranes was unchanged. The stabilized biomaterials could be stored
in culture medium over months (Fig. S3-S5) with mechanical properties characterized by a mean Young’s modulus of ca 6 kPa (Fig. S6 and Supplementary experimental section).

Whether collagen may be denatured during ES remains an open question in literature. This can involve both the solvent used for improving the electrospinnability of the collagen or the spinning process by itself and the high voltage applied. This question is addressed in the following sections through a multiscale characterization from the micron to the molecular scale.

**Hierarchical organization of collagen**

In living tissues, collagen self-assembles into fibrils, with triple helices all oriented in the same direction with the same polarity. SHG is a coherent process specific for non-centrosymmetric materials. At the molecular level, it relies on nonlinear dipole excitation along the peptide bonds. It builds up efficiently in highly anisotropic fibrillar collagen because of the tight unidirectional alignment (parallel) of peptide bonds along the collagen triple helix and within fibrils both in isolated collagen fibrils\(^{39,40}\) and fibrillary networks.\(^{41}\) This makes SHG microscopy the gold standard technique for 3D characterization of collagen-rich tissues in ambient conditions and without exogeneous labelling, with a structural specificity that cannot be reached with any other optical and fluorescence microscope, being specific to (i) the native state of collagen (over gelatin) and (ii) the unidirectional alignment of collagen triple helices.\(^{42}\)
Figure 2. (1) SHG, (2) pSHG and (3) cryo-FE-SEM images of (A) a dry collagen membrane, (B) after exposure to ammonia vapors and (C) rehydration in culture medium. The look-up table (on the right in the A1-C1 line) corresponds to arbitrary units for each SHG image. The orientation in the pSHG image is displayed on the colored circular scale (on the right in the A2-C2 line) ([collagen] = 1.7 mg.mL\(^{-1}\) in 30 mM HCl:EtOH (25:75 v/v)). Scale bar for SHG and pSHG is 20 µm. (D) Histogram of bundle width distribution according to cryo-FE-SEM.
Measurements on dry membranes showed a high SHG signal (Fig. 2A1). This is a proof of the native state of collagen that corresponds to a non-centrosymmetric packing of the triple helices. In addition, the grainy-like pattern with a myriad of dots indicates the presence of small structures that are highly entangled, in agreement with SEM observations (Fig. 1C,D). Indeed, given that the diameter of the fibrils (58 ± 10 nm according to SEM images) is small relatively to the focal volume (typically 0.4 μm (lateral) × 1.7 μm (axial)), many fibrils may be simultaneously observed in one pixel. At the intersection of two fibrils, molecules in the focal volume are not all oriented in the same direction, which results in a centrosymmetric distribution and leads to a local decrease of SHG signal (Fig. 3A1). If the network is highly entangled, SHG decrease is widespread throughout the focal volume leading to a grainy-like image (Fig. 3A2-A3). The structural specificity of SHG images has been further improved by acquiring polarization-resolved images (pSHG), which measures the main orientation of collagen in the imaging plane, in every pixel of the SHG image.\textsuperscript{34} Given that each color codes for a given direction, the presence of dots with many different colors throughout the whole image shows that the highly packed structures of native collagen are randomly distributed with respect to one another (Fig. 2A2). Again, at fibrils intersection, all molecules are not oriented in the same direction, which impedes the determination of a main orientation, resulting in black pixels (Fig. 3B).
**Figure 3.** Schematic representation of the local decrease in (A) SHG and (B) pSHG intensity: (1) at the intersection of two fibrils, and (2) in highly entangled network of small diameter-fibers. (3) Corresponding SHG images with a grainy-like pattern ([collagen] = 1.7 mg.mL$^{-1}$ in 30 mM HCl:EtOH (25:75 v/v)).

After 20 min exposure to NH$_3$ vapors, no variation of the SHG and pSHG profiles could be detected (Fig. 2B1-2). This shows the preservation of the triple helix structure and packing within small entangled structures. No important variation at the scale probed by SHG can be detected despite the very different behaviors of the dry and stabilized membranes when immersed in culture medium.

In contrary, SHG observations of the hydrated membrane showed very important variations of the membrane structure that appears made up of large micrometer-long fibrils forming ca. 100 µm-
large domains (Fig. 2C1). This is reminiscent of collagen matrices in dense fibrillated media. pSHG very nicely reveals the unidirectional ordering of the triple helices in each collagen fibril that appears as monocolored structures (Fig. 2C2). In this case, the absence of grainy-like profile is attributed to the fact that collagen reorganization leads to the formation of large structures. Indeed, one of the major features of type I collagen is its ability to self-assemble into highly organized supramolecular structures, which in turn form larger highly oriented fibers and fascicles. This prevents the observation of multiple collagen fibrils with random organization with respect to each other within the focal volume, hence precluding local SHG extinction.

Further characterization of collagen triple helix arrangement can be obtained at the nanometer scale by use of electron microscopy. We used cryo-FE-SEM that preserves the hydrated state of collagen in hydrated biomaterials. The dry membrane exhibits the same structure as observed under conventional SEM (Fig. 1) due to the absence of solvent within the initial biomaterial: randomly entangled fibrils are clearly visible (Fig. 2A3). Once stabilized under NH$_3$ vapors, the overall structure of the sample remains unchanged, exhibiting an entangled network but with slightly melted fibrils (Fig. 2B3). This shows that ammonia exposure does not affect the 3D organization of the collagen nanofibers. This is a very interesting alternative to crosslinking strategies usually used to stabilize collagen-based membranes. Unlike previous works, in this case no gelation process is induced: ammonia exposure allows for the membrane storage in a pre-fibrillated state. This can be attributed to the limited extent of fibrillation at the solid-vapor interface hampered by the low amount of water as diffusing medium. After hydration, large fibrils could be observed exhibiting the 67-nm periodic pattern characteristic of collagen organization in living tissues (Fig. 2C3). Additional images can be found in the photo gallery of the Supplementary Materials to better highlight the nanostructure of collagen fibrils with the periodic pattern of 67
nm (Fig. S7). These fibrils further organized into bundles exhibiting large width and polydispersity: from 200 nm to 2 µm, with a mean width of 560 ± 290 nm over a population of 61 bundles (see histogram of bundle width distribution in Fig. 2D). This very nicely confirms the variations of the collagen network observed by SHG/pSHG and again proves the native state of collagen within ES membranes given that non-native collagen cannot form cross-striated fibrils. Very importantly, this ascertains the possibility to recover the self-assembling processes and hierarchical organization of collagen as observed in living tissues. This is the first time that such organization could be obtained after ES.

However, SHG and cryo-FE-SEM give a rather local view of the sample and we require at this stage to have the confirmation of collagen native state over the whole sample. To do so, we used synchrotron-radiation circular dichroism (SRCD) that gives the molecular signature of native collagen at the scale of the triple helix over the whole sample. Self-supported membranes were dissolved in 3 mM HCl. A large negative band at 198 nm and a small positive band at 223 nm were observed at 20°C (Fig. 4A solid line). This corresponds to the spectral signature of the collagen triple helix after extraction and before ES (Fig. 4A, dotted line). These bands are characteristic of the polyproline II (PPII) conformation of collagen triple helix. This CD spectrum strikingly deviates from previous studies that showed collagen denaturation,10,11 and confirms that collagen fully preserves its native state after ES. The melting temperature (Tm) of the soluble triple helix was measured by thermal denaturation with increasing temperature up to 74°C in steps of 3°C. As expected, the intensity of both bands decreases with the temperature increase (Fig. 4B from dark blue to dark red). This is attributed to the increased fraction of unfolded structures upon thermal denaturation. Tm was determined by plotting the ellipticity at maxima (198 and 223 nm) as a function of temperature using the sigmoidal fit in “Igor Pro” spectral analysis program. After
ES, the Tm was found to be around 31°C at 198 nm and 30.8°C at 223 nm (Fig. 4C), in agreement with previously reported values for native collagen. Above 31°C, the structure is dominated by unordered conformations, and collagen completely unfolds above 40°C. To estimate the impact of the ES process, we compared the thermal denaturation of the electrospun triple helix with that of native collagen after extraction. Figure 4D shows that Tm of pure collagen is similar to the one of the ES membrane (31.5°C at 198 nm and 31.7°C at 223 nm). This shows that the fraction of PPII at room temperature in both samples is the same and further confirms that collagen fully preserves its native state over the whole sample.
Figure 4. SRCD spectra of (A) collagen in 3 mM HCl before (dotted line) and after (solid line) ES, and of (B) thermal unfolding of ES collagen from 10 to 74 °C in steps of 3°C. (C-D) Ellipticity as a function of temperature: Tm at 198 and 223 nm for ES and native collagen respectively.

**Biofabrication for a broad range of applications**

To assess the cytocompatibility and bioactivity of electrospun collagen nanofibers, cell attachment and spreading of normal human dermal fibroblast cells (NHDF) seeded onto the
collagen matrix were studied. Immunostaining of actin and nuclei revealed the colonization of the scaffold, where the fibroblasts were spread all over the membrane and exhibited a spindle-like shape morphology with stress fibers clearly visible inside the cellular body (Fig. 5A, white arrows), showing that electrospun membrane are perfectly suitable for cell adhesion and spreading. No significant difference in terms of cell morphology and area could be observed when compared to control non-electrospun biomaterials, ie casted membranes obtained by solvent evaporation (Fig. 5B). Finally, the metabolic activity after one week was assessed revealing no significant difference between electrospun, casted and control support (Fig. 5C).

**Figure 5.** (A1-A2) Fluorescence microscopy images at different magnifications of NHDF cultured for 24 h on ES membranes. Actin (green: phalloidin) and nucleus (blue: DAPI) staining. White
arrows show stress fibers. (B) Cell area of NHDFs seeded on electrospun and casted membranes (n=3). Statistical analysis performed using Mann-Whitney non parametric test (*:p< 0.05). (C) Metabolic activity of NHDFs measured by Alamar blue assay after a one week culture on different supports (ES and casted collagen membrane, and control plastic support). The column represents Mean with SEM.

The processing of biopolymers with preserved native molecular conformation, hierarchical structure and interactions with cells is necessary to replicate biological microenvironments. This remains a huge challenge and a real need for the advancement of basic research in biology and biomedical research. With this challenge ahead, we first worked on the engineering of hybrid _ or composite _ biomaterials obtained by mixing collagen with an inorganic phase. The composite approach is an efficient strategy to improve the functionality of bio-inspired matrices by synergistically combining biopolymers with colloids of different chemical nature. In this context, we added carbon nanotubes (0.125% in weight) to collagen before ES. Self-supported membranes could be obtained, in identical conditions to that of pure collagen, with a very good homogeneity in terms of carbon nanotube dispersion within the final biomaterials, as observed both macroscopically and under SEM (Fig. 6A, where carbon nanotubes are hardly distinguishable among collagen nanofibers). Alternatively, silica particles could successfully be electrospun and incorporated within a collagen-based self-supported membrane with good dispersion (Fig. 6B, white arrows). This shows that the ES of collagen is a robust process that allows adjusting the composition of the biopolymer solution for the electrospinning of inorganic colloids. In addition, and very importantly, the ES of collagen ensures a good dispersion of the two phases that would not be reached using a casting strategy based on solvent evaporation. Given the strength of the
composite approach to design conductive, optical, magnetic devices, or tune their mechanical properties,\textsuperscript{43,44} collagen ES represents an important step for the engineering of collagen-based biomaterials.

Figure 6. Photos (inserts) and SEM images of hybrid biomaterials combining collagen with (A) carbon nanotubes and (B) silica particles.

Another advantage in using ES is the possibility to easily control the thickness of the deposited membrane for the coating of a support of interest. This is particularly interesting in nano- and microtechnologies typically for the fabrication of microfluidic chips. Microfluidic chips are becoming crucial tools in biological and biomedical research by implementing molecular exchange and flux to \textit{in vitro} models. It is thus of utmost importance to provide micro biochips with scaffolding that recapitulates the architecture and cell interactions of living systems. In this context, we used microfabricated frames to be incorporated into microfluidic devices as a support
for collagen ES (Fig. 7A). These microframes have been shown to be highly suitable for cell culture and stem cell differentiation.\textsuperscript{45-48} In this case, collagen is electrospun directly onto the microframe as described above, and the microframe prefibrillated under ammonia vapor and hydrated in culture medium. In those conditions, SHG observations on top of the microframe showed the formation of submicron thick collagen layers with native hierarchical structure (Fig. 7B). This again shows the robustness of our protocol to produce ECM-like scaffolds made up of native type I collagen. This makes the ES of collagen a precious alternative to synthetic or non-native biopolymers or to the use of hydrogels with variable thickness.

\textbf{Figure 7.} (A) Photo of the microfabricated frame and bio-chip before ES, and (B) SHG image of the coating of the microframe by collagen ES.
CONCLUSION. Native type I collagen can be electrospun into a self-supported membrane in absence of cross-linker and polymer support. This is a prerequisite to improve in vitro models providing cells with the necessary biochemical and biophysical cues. We propose a methodology to thoroughly characterize collagen-based biomaterials using SHG, which is the most powerful optical probe for self-assembled type I collagen given its high specificity to triple helix packing within collagen fibrils, combined with cryo electron microscopy to preserve the hydrated state of the collagen membrane and SRCD as the gold biochemical standard to investigate the conformation of biomolecules. Altogether, this shows that collagen preserves its native conformation and ability to self-assemble into fibrils after ES. Dry electrospun collagen can be stabilized by pre-fibrillation in vapor state to avoid collapse upon hydration. This procedure allows to store dry membranes as pre-fibrillated scaffolds. This provides very easy to handle biomaterials of pure collagen, ready to use for biomedical applications including implantations, with preserved hierarchical structure and biological activity. Because of the final processing of ES materials into thin coatings and self-supported films, a broad range of applications can be targeted, including wound healing, cornea repair, heart patches and neural regeneration. We also show that this process can be applied to the coating of microchips for biological microfluidics with the possibility to adjust the composition of the biopolymer solution to be electrospun, notably towards hybrid materials. This opens many developments for the reconstruction of microenvironments with improved biological relevance for future advances in biological and biomedical research.

ASSOCIATED CONTENT

Supporting Information. Optimization of the solution conditions to improve collagen electrospinnability; Macroscopic structure of the fibrillary self-supported membrane; Passive
degradation after one month in cell culture medium; Mechanical properties of the ES membrane; Photo gallery of the hydrated electrospun membrane by cryo-FE-SEM; Supplementary experimental section for passive degradation test and AFM nanoindentation and imaging.

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Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

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Notes

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ABBREVIATIONS
CCR2, CC chemokine receptor 2; CCL2, CC chemokine ligand 2; CCR5, CC chemokine receptor 5; TLC, thin layer chromatography.

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SYNOPSIS (Word Style “SN_Synopsis_TOC”).
