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Disordered protein-graphene oxide co-assembly and supramolecular biofabrication of functional fluidic devices

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Supramolecular chemistry offers an exciting opportunity to assemble materials with molecular precision. However, there remains an unmet need to turn molecular self-assembly into functional materials and devices. Harnessing the inherent properties of both disordered proteins and graphene oxide (GO), we report a disordered protein-GO co-assembling system that through a diffusion-reaction process and disorder-to-order transitions generates hierarchically organized materials that exhibit high stability and access to non-equilibrium on demand. We use experimental approaches and molecular dynamics simulations to describe the underlying molecular mechanism of formation and establish key rules for its design and regulation. Through rapid prototyping techniques, we demonstrate the system's capacity to be controlled with spatio-temporal precision into well-defined capillary-like fluidic microstructures with a high level of biocompatibility and, importantly, the capacity to withstand flow. Our study presents an innovative approach to transform rational supramolecular design into functional engineering with potential widespread use in microfluidic systems and organon-a-chip platforms.

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here is an increasing interest to generate materials with bioinspired functions¹, such as the capacity to grow², self-replicate³, or controllably respond to specific stimuli⁴. Biological materials acquire most of these functionalities as a consequence of their ability to self-assemble various types of building blocks at multiple length scales. Engineering materials in this manner provides an opportunity to take advantage of the individual building blocks while enabling emergent properties as a result of their interactions^{5,6}. Consequently, multicomponent self-assembly represents an attractive route to develop more complex materials⁷ with enhanced modularity and tuneability of properties^{8,9}, such as structural hierarchy¹⁰, adhesion¹¹, electrical conductivity¹², or the capacity to grow¹³.

Proteins are the most functional building blocks of organisms¹⁴ and, as such, have been thoroughly explored to engineer intelligent materials^{15–17}. However, new discoveries are shaping our understanding of how proteins function and providing new insights for their utilization. For example, there is increasing evidence that both ordered (i.e., β-sheet and α-helix) and disordered (i.e., random coil) regions of proteins play a role in their functionality¹⁸ and growing acceptance that this functionality is regulated by their interaction with other molecules 19. Based on these principles, proteins are emerging as dynamic building blocks of multicomponent systems to engineer intelligent materials. We have recently reported on the possibility to exploit the disordered nature of elastin-like recombinamers (ELRs) to modulate their conformation and generate dynamic¹³ or hierarchically mineralizing²⁰ materials. ELRs, also known as the recombinantly produced elastin-like polypeptides, are based on the natural elastin motif Val-Pro-Gly-X-Gly (VPGXG), where X could be any amino acid apart from proline²¹. These molecules exhibit a reversible-phase transition with a change in temperature and have been used to create biocompatible materials²².

Multicomponent self-assembly also offers a unique opportunity to engineer complex hybrid systems. In particular, the controlled incorporation of graphene as a building-block could lead to the design of new biomaterials that benefit from its distinctive two-dimensional (2D) structure and outstanding electronic, thermal, and mechanical properties²³⁻²⁵. Toward this goal, graphene and its derivatives have been modified with biomacromolecules²⁶, such as DNA²⁷, proteins²⁸, and biopolymers²⁹ and used in, for example, implants and scaffolds for cell culture and regenerative medicine^{30,31}. Furthermore, graphene oxide (GO) is gaining significant interest and being used instead of graphene given its rich oxygen-containing functional groups (hydroxyl, epoxy, carbonyl, and carboxyl), which facilitate designed interactions with different molecules. However, both graphene and GO exhibit key limitations such as dose-dependent toxicity and issues associated with hierarchical organization and the ability to generate uniform and stable structures^{24–26}.

Material platforms that exploit the functionalities of both proteins and GO and enable their multiscale organization offer exciting possibilities for the engineering of advanced materials. We report a hierarchical self-assembling system that takes advantage of protein disorder-to-order transitions and supramolecular protein-GO interactions to enable both stable structures and access to non-equilibrium with spatial control to design functional materials and devices. Experimental approaches and molecular dynamics (MD) simulations were used to elucidate the underlying molecular mechanism and develop rules for its use. We show that the material can be combined with rapidprototyping techniques to assemble well-defined tubular microstructures embedded with cells and into fluidic devices. The study introduces an innovative way to biofabricate by self-assembly complex and functional devices such as microfluidic systems or organ-on-a-chip devices.

Results

System rationale. Previous studies have demonstrated the possibility to co-assemble peptides with large macromolecules to generate hierarchical membranes at a liquid-liquid interface 10,13. These systems rely on both molecular interactions, such as electrostatic and hydrophobic forces between the two components as well as their respective individual properties, such as molecular weight, charge, and 3D conformation. In particular, by coassembling ELRs and peptide amphiphiles (PAs), we have previously reported on a diffusion-reaction mechanism that relies on PA diffusion to give rise to a multilayer membrane that can access non-equilibrium¹³. However, this material is fragile and can only be assembled in and is stable under a narrow window of environmental conditions (pH, temperature, and salt concentration), which limits its functionality and widespread use. Giving the need for multicomponent approaches that can turn molecular design into functional systems¹, we envision the possibility to exploit the inherent properties of GO to work synergistically with disordered proteins to create materials with both emergent properties and functionality. Specifically, we reasoned that, unlike the PAs, the GO lamella conformation in aqueous environments³² would provide a supramolecular framework with high surface area for ELR interaction, reaching a level of integration far beyond that of the ELR-PA system. Furthermore, the GO's flat-sheet organization at air-liquid interfaces³³ would facilitate the generation of a diffusion-reaction process that takes advantage of the disordered nature of ELRs to diffuse, conform, and integrate with GO, generating a hierarchical process of assembly that can be manipulated on demand. In this way, we hypothesized that the co-assembly of GO and ELRs would lead to a robust supramolecular system where both strong molecular interactions and controllable access to non-equilibrium would lead to new materials with enhanced complexity and functionalities.

Materials rationale. We used GO sheets of two different average lateral sizes, including larger GO (GO-L) measuring $10.5 \pm 4.5 \,\mu m$ and smaller GO (GO-S) of $2.3 \pm 0.9 \,\mu\text{m}$, both exhibiting a typical hydrophobic surface and negatively charged carboxylic groups on their periphery (Fig. 1b and Supplementary Fig. 1). We chose ELRs as the protein component because of their modular and disordered nature³⁴ and the possibility to exhibit different molecular conformations at different temperatures³⁵. The ELK1 sequence (Fig. 1a) is a 51.9 kDa molecule consisting of 24 repeats of a single block made of four hydrophobic pentapeptides (VPGIG) and a positively charged (VPGKG) one. This relatively simple molecular design offers an accessible transition temperature (Tt) of 30 °C (at 2% ELK1 in MilliQ water) with clearly different ELR conformations above or below it, as well as medium molecular weight to enable both cooperative interactions between its charged and hydrophobic segments as well as with the anionic edge and hydrophobic surface of the GO (Fig. 1c). ELRs with similar molecular weight but different levels of charge and hydrophobicity (Fig. 1a), as well as a single repeat of an individual block of each of these three ELRs, were used as controls (Supplementary Figs. 2-4).

Co-assembly. When an ELK1 solution at its Tt (30 °C) is immersed in a larger volume of a GO solution, a multilayered membrane of up to 50 μ m in thickness develops at the interface around the immersed drop maintaining both solutions separated (Fig. 2a and Supplementary Movie 1). This membrane consists of layers made from both GO sheets and ELK1 (Fig. 2b, confocal), with GO sheets being present throughout the cross-section of the membrane (Fig. 2b, SEM) and ELK1 gradually decreasing in concentration from the inside (ELK1 side) to the outside (GO side)

a Molecular information of elastin-like recombinamers (ELRs)

ELR molecules		Sequence	Tt (°C) (2% water solution)	Molecular weight (kDa)
ELR-I120	(ELK0)	MESLLP-(VPGIG VPGIG VPGIG VPGIG)24	15	51.6
ELR-IK24	(ELK1)	MESLLP-(VPGIG VPGIG VPGIG VPGIG)24	30	51.9
ELR-(IK3I)2	4 (ELK3)	MESLLP-(VPGIG VPGKG VPGKG VPGIG)24	80	52.7

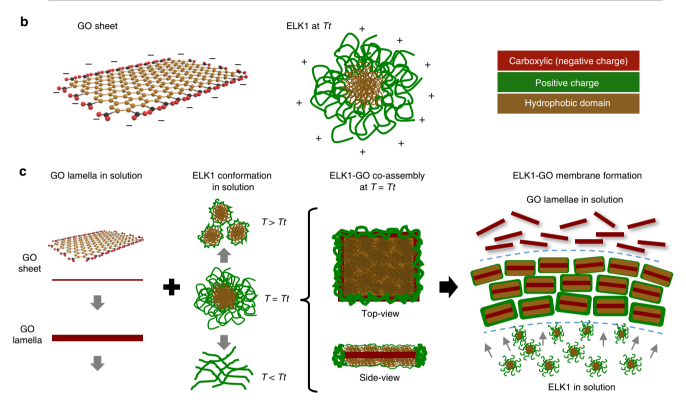


Fig. 1 Molecular building blocks and rationale for co-assembly. a Table summarizes the key information of the three elastin-like recombinamers (ELRs) used in the study comprising similar molecular weight but different levels of hydrophobicity (VPGIG) and positive charge (VPGKG). **b** Illustrations of the molecular structure of a GO sheet and the supramolecular organization of ELK1 at its transition temperature (*Tt*) (30 °C) indicating both the charged (red and green) and hydrophobic (brown) segments. **c** Schematic of the proposed mechanism of formation illustrating the molecular and supramolecular conformation of the GO and ELK1 before and after co-assembly at the ELK1's *Tt* as well as their interaction for membrane formation.

of the membrane (Fig. 2c and Supplementary Fig. 5). Multilayered structures are known to emerge from diffusion–reaction mechanisms³⁶. We have previously demonstrated that co-assembling PAs with ELRs, it is possible to trigger a diffusion–reaction mechanism, which generates multilayered membranes capable of exhibiting dynamic properties¹³. Similarly, by touching any surface within the first few seconds of formation, the ELK1–GO membrane adheres, spontaneously and reproducibly opens, and can be manipulated to grow into tubular structures with spatiotemporal control (Fig. 2a, d, and Supplementary Movie 2). However, in this case, the underlying ELR-GO mechanism of interaction and supramolecular assembly lead to the growth of a material with remarkably enhanced properties.

Material structure, properties, and biofabrication of devices. First, the ELK1–GO membrane is both dynamic and highly stable, permitting controlled anisotropic growth of tubular geometries that exhibit no apparent effects on their multilayered structure when the temperature drops below (down to 4 °C) or raises above (up to 70 °C) the Tt of ELK1. This enhanced stability is also evidenced by the capability to co-assemble capillary-like structures down to ~50 μ m in internal diameter, defined by the

size of the injecting tip. Moreover, the system also enables the capacity to decrease the thickness of the wall down to ~10 µm, achieved by removing the GO solution in order to stop the assembly after ~2 min (Fig. 2e). While smaller diameters may be possible by using smaller tips, wall-thicknesses below ~10 µm were found to be too fragile to be manipulated. In addition, these structures can be grown as tubular bridges between gaps by simply touching, adhering, opening, and sealing to a surface soon after co-assembly and continuing injecting ELK1 solution into the GO solution until the next surface is touched (Fig. 2f and Supplementary Movie 3). Moreover, the system works in saltcontaining solutions such as cell culture media, which enables coassembly and growth of capillary-like structures in the presence of cells, resulting in structures comprising cells embedded within and on the wall of the tube (Fig. 2g). Furthermore, given this versatility and robustness, we demonstrated the possibility to use rapid-prototyping techniques to guide the co-assembly process using an extrusion-based 3-D printer to print the ELK1 solution within a GO solution (Supplementary Movie 4), generating fluidic devices containing high-aspect ratio tubular structures of different internal diameters and comprising curves (Fig. 2h, i and Supplementary Movie 5), angles of different sizes (Fig. 2h, i), and bifurcations (Fig. 2h, k and Supplementary Movie 6). The fluidic

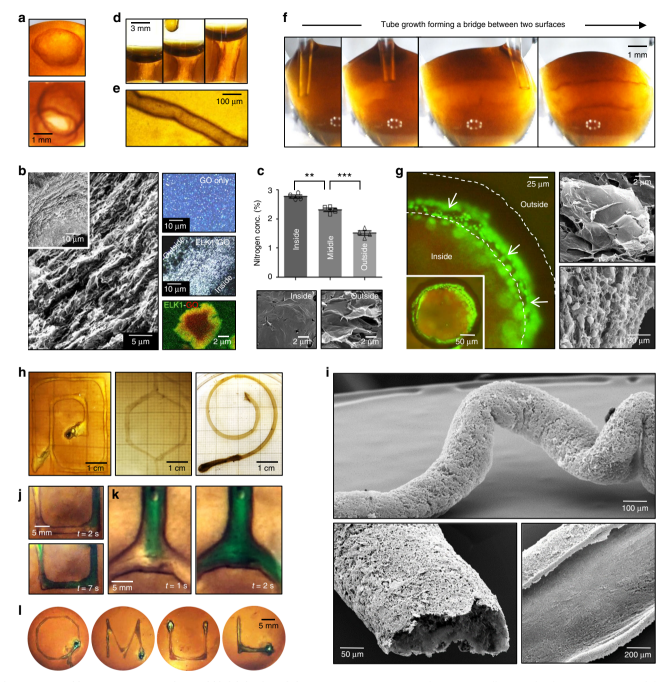


Fig. 2 Co-assembly, structure, properties, and biofabrication of the ELK1-GO system. a Time-lapse images illustrate the dynamic properties of the ELK1-GO membrane first (**a**-Top forming a closed sac when a drop of ELK1 solution is immersed in a larger GO solution and second (**a**-Bottom) opening upon touching an interface within the first seconds of formation. **b** The membrane exhibits a multi-layered architecture of about 50 μm thick comprising aligned GO sheets throughout (birefringence inset) interacting with ELK1 molecules (fluorescence image, green: ELK1, red: GO), **c** which are observed to decrease in concentration from the inside to the outside as evidenced by wavelength-dispersive spectroscopy (WDS). Only ELK1 comprises nitrogen in its molecular structure. ±s.d. for n = 3. *p < 0.05. t test. **d** The system enables growing the membranes into longer tubes on demand by displacing an interface. **e** The robustness of the system enables formation of capillaries down to about 50 μm in internal diameter with 10 μm thick walls, **f** bridging of surfaces simply by touching two interfaces while injecting one solution into the other, and **g** co-assembling in salt solutions, opening the possibility to embed cells (green identified by white arrows) within the membrane (outlined by dashed lines) as the tubes are formed. The images are taken after 24 h of culture and correspond to a live (green)/dead (red) assay. Scanning electron micrographs of cells embedded within layers of GO (top) and a cross-section of the ELK1-GO membrane comprising cells within different layers (bottom). **h-l** Images demonstrate the versatility of the co-assembly system by incorporating it with 3D printing to fabricate well-defined fluidic devices consisting of high-aspect ratio tubular structures (**h**) of different internal diameters and comprising curves, angles of different sizes, and bifurcations (**h**, **i**, **l**) capable of withstanding flow within a few minutes of formation (**j**, **k**).

devices were able to withstand aqueous flows of up to 12.5 mL/min for at least 24 h and within 60 min of formation (Supplementary Movies 7 and 8). The highest flows would generate 0.26 N/m² shear stress, which is within the range of mean shear stress values observed in common carotid arteries (0.7 N/m²)³7. Altogether, these capabilities suggest that the mechanism of formation exhibits both strong ELK1–GO interactions at the molecular scale and integrated organization at higher size scales (Supplementary Fig. 6 (ELK1)).

Underlying molecular mechanism of assembly: ELK1-GO molecular interactions. We first tested the presence of both electrostatic and hydrophobic forces by quantifying ELK-GObinding constants using ELRs with varying levels of charge and hydrophobicity. Tubes formed on application of ELK1 and ELK3 but not ELKO, confirming the need for electrostatic forces for its assembly. Interestingly, the highest binding constant (Ka), calculated by fluorescence emission titration, was obtained with ELK1 (1.3×10^6) compared to ELK0 (7.2×10^4) and ELK3 (3.2×10^4) 10⁵) (Fig. 3a, Supplementary Fig. 6b, c). Using this method, we also found that the optimum ELK1-GO concentration ratio to maximize the interaction between them is 15-40 (Supplementary Fig. 7). Therefore, keeping the ELK1 concentration at 2%, we developed tubes using GO concentrations between 0.05% (corresponding to an ELK1-GO ratio of 40) and 0.15% (corresponding to an ELK1-GO ratio of 15), and, as expected, qualitatively found that the best-defined and most robust tubes were made within this range (Fig. 3b). To quantify and identify the best ELK1-GO combination, we used an established nanotensile test³⁸ on tubes made of 2% ELK1 and increasing concentrations of GO (0.05, 0.10, and 0.15%) (Supplementary Section 14). As expected, the strength, the strain at break, and the toughness modulus increased on tubes formed with increasing concentrations of GO (Fig. 3c, Table). However, based on a Weibull statistical distribution, the results revealed that the elastic modulus was highest on tubes fabricated with 0.10% GO (212.90-247.15 kPa) compared to 0.05% (128.78-147.37 kPa) and 0.15% (159.57-208.16 kPa). This result is also visible from the stress-strain curves of the ELK1-GO (Fig. 3c, graph), where the samples made with 0.1% GO show a steeper slope, meaning that the material is stiffer.

In order to further investigate the role of electrostatic interactions, we formed tubes with ELR and GO solutions at varying pHs and again found that more robust membranes formed when the charge difference between both components was marginal (Fig. 3d, e). These results suggest that optimum coassembly does not solely depend on strong electrostatic forces but rather on a synergistic effect from different factors that we speculate to be electrostatic and hydrophobic forces, H-bonding, and 3D conformation. To confirm this premise, we first synthesized a single repeat of each individual ELR block. Using CD and MD simulations, we verified that these shorter molecules did not exhibit a Tt but have similar secondary structure with large amounts of random coil in aqueous environments (Supplementary Fig. 8). Upon mixing with GO, all three single repeat peptides exhibited similar levels of interaction as evidenced by calculation of the binding constant based on fluorescence emission titration (Supplementary Fig. 9). To further dissect the nature of the initial ELK1-GO interactions, we performed MD simulations at 30 °C and found that H-bonding between the ELK1 and GO plays a role and that these interactions can come from both the linear side chain of lysine and the backbone of the peptide (Supplementary Section 18). However, in addition to these ELK1-GO molecular interactions, we hypothesize that the 3D conformation of the full-length ELK1 protein and its ability to

cooperatively interact with the GO lamellae play a key role in the formation of the system. To test this hypothesis, and taking advantage of the ELR's capacity to change its conformation at different temperatures (Fig. 3f, graph, Supplementary Figs. 10 and 12), we assembled tubes using GO and ELK1 (2 wt%) at either below (4 °C), above (45 °C), or the ELK1's Tt (30 °C) (Fig. 3f). While tubes formed at all temperatures, they were more robust and exhibited better-defined multilayers (Supplementary Fig. 11e) and tubular geometry (Fig. 3f, images) at 30 °C, suggesting stronger interactions at this temperature. This enhanced interaction was also investigate by DLS, which revealed the presence of larger ELK1-GO aggregates at 30 °C compared to 4°C and 45°C (Fig. 3g) and further confirm that the 3D conformation of ELK1 at the different temperatures plays a key role in its interaction with the GO lamellae, which would in turn affect the diffusion-reaction mechanism and consequently determine the properties of the resulting ELK1-GO tubes (Fig. 3f, images).

Underlying molecular mechanism of assembly: ELK1-GO aggregates. To shed light on this enhanced ELK1-GO interaction at 30 °C, we used SANS and found that, as expected, ELK1 exhibited an expanded conformation at 4 °C and a collapsed aggregated conformation with a 74 nm radius of gyration of the core region at 45 °C (Supplementary Fig. 10b). Furthermore, at 30 °C, the molecule acquired a conformation that combined both an expanded structure and a collapsed aggregate core, consisting of a 60 nm radius of gyration of the core region surrounded by a larger 500 nm radius corona of expanded structures (Supplementary Fig. 10b). These different conformations were confirmed by cryo-transmission electron microscopy (cryo-TEM) (Supplementary Fig. 13). On the other hand, GO sheets are known to stack and form lamellae³² in aqueous environments. We hypothesized that the disordered nature of ELK1 would facilitate its interaction with the supramolecular framework provided by the GO lamella. To test this hypothesis, we used SANS to investigate the size and shape of the ELK1-GO aggregates upon co-assembly (Fig. 4a, b, and Supplementary Fig. 11a). We found that the scattering profile for the ELK1-GO aggregate formed at 30 °C is better fitted with a classical core-shell-bicelle-elliptical model³⁹ with a core measuring 7 nm in length, a thick_rim of 22 nm, and a thick_face of 16 nm (Supplementary Fig. 11d, e). According to this model, the core is formed by GO and the shell by ELK1. This core-shell conformation was confirmed by confocal microscopy (Fig. 4c). On the other hand, at 4 and 45 °C, the ELK1-GO aggregates acquire longer cores and thinner shells (Supplementary Fig. 11d, e), which suggests that at these temperatures the GO lamellae are less infiltrated by ELK1 molecules. In contrast, at 30 °C, the shorter core of the ELK1-GO aggregates indicates that the GO lamellae are more infiltrated by and likely interacting more with the ELK1 (Supplementary Fig. 11d, e).

Underlying molecular mechanism of assembly: disorder-to-order transitions to enhance integration. It is well-known that proteins rich in disordered regions change their conformation upon binding to other molecules or surfaces⁴⁰. We hypothesize that the enhanced infiltration by ELK1 within the GO lamellae at 30 °C is associated to the disordered nature of the ELK1 and its potential to acquire different secondary structures upon interaction with other molecules. We first used FT-IR amide III spectra to conduct a quantitative analysis of the ELK1's secondary structure (Fig. 4d)^{20,41}. At 30 °C and prior to co-assembly, ELK1 exhibits a high degree of random coil but also higher amounts of α -helix and lower amounts of β -sheet compared to 4 and 45 °C (Fig. 4d). Interestingly, upon binding with GO at 30 °C, ELK1

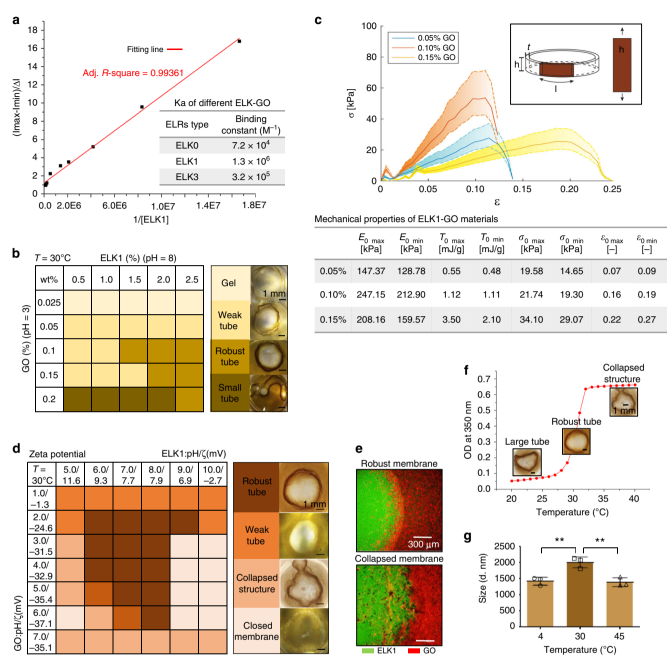


Fig. 3 Molecular interaction, composition, and mechanical properties of ELR-GO. a Binding constants (Ka) for the different ELR-GO combinations calculated by a Benesi-Hildebrand equation based on fluorescence emission titration of a mixture of GO (2.5×10^{-3} wt%) in MilliQ water solution and increasing concentrations of ELRs revealing higher Ka for ELK1-GO compared to ELK0-GO and ELK3-GO. **b** Table illustrating the role of building-block concentration ratio on the formation of ELK1-GO tubes. **c** Nanotensile test results described with the Weibull distribution and reported in the table reveal that the strength, the strain at break, and the toughness modulus increased on tubes formed with increasing concentrations of GO but the elastic modulus was highest on tubes made with medium level (0.10%) GO compared to lower (0.05%) and higher (0.15%) amounts. **d** Table illustrating the role of pH and ζ on the formation of the ELK1-GO tubes and their respective geometry. **e** Representative confocal microscopy qualitatively depicting the interface between ELK1 (green) and GO (red) during tube formation with different levels of definition as characterized in **d**. **f** Red line of the graph shows the turbidity changes of an ELK1 (2 wt%) solution in MilliQ water while inserted images depict the definition of tubes formed at specific temperatures. **g** Dynamic light scattering (DLS) revealing the presence of larger ELK1-GO aggregates at 30 °C compared to 4 °C and 45 °C. Error bars represent ±s.d. for n = 3. *p < 0.05. One-way ANOVA.

maintains its α -helix and increases in β -sheet (Fig. 4d). These results suggest that as ELK1 molecules diffuse through the GO lamellae (Fig. 4a–c) at 30 °C, they bind to and interact with GO maintaining their α -helix but increasing their levels of β -sheet (Fig. 4d and Supplementary Figs. 10 and 11). It is known that higher levels of β -sheet conformation generate denser aggregates 42 and that α -helix proteins lose entropy and increase the

system's stability by aggregating their helices⁴³. Therefore, it is possible that these secondary structures enhance the stability of the ELK1–GO complex and consequently lead to a better-defined diffusion barrier at the beginning of the co-assembly process, which is known to have an effect on interfacial membrane assembly⁴⁴. To confirm this, we attempted to form tubular structures using the GO-S, which instead lead to a gel-like

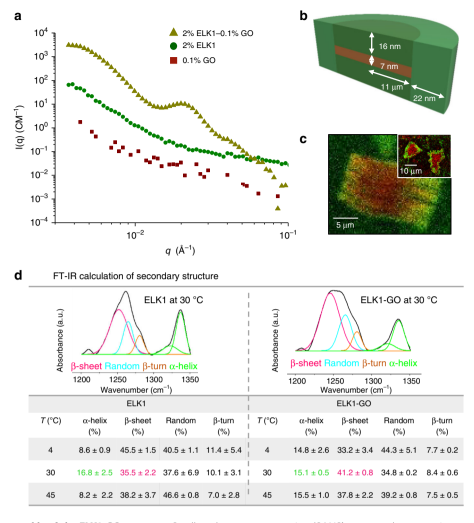


Fig. 4 Supramolecular assembly of the ELK1-GO system. a Small-angle neutron scattering (SANS) patterns demonstrating a resulting uniform microstructure formed when co-assembling ELK1-GO. Between the middle-q region (ca 0.007-0.04 Å⁻¹), the ELK1-GO structure (yellow triangle) exhibits a characteristic scattering peak associated with pure ELK1 (green square) and GO (red circle) at 30 °C, confirming the formation of a new order structure different from the individual components. **b** The classical core-shell-bicelle-elliptical model that was fitted to the ELK1-GO microstructure as measured by SANS at 30 °C (green: ELK1, brown: GO). **c** Confocal microscopy (green: ELK1, red: GO) corroborating the interaction between the ELK1 and the GO lamellae (inset depicts the top view of the ELK1-GO structure). **d** FT-IR calculation of secondary structure depicting the change and transition of conformation of the ELK1 molecule before and after binding with GO. At 30 °C and before binding with GO, ELK1 exhibits higher α-helix than at 4 and 45 °C, which is maintained after binding with GO and is complemented with an increase in β-sheet.

structure, suggesting the formation of a loser and more permeable diffusion barrier (Supplementary Fig. 1b).

Biological validation. The versatility and robustness of the ELK1-GO system offers an exciting possibility to develop complex and functional biohybrid devices with a high level of biological relevance by supramolecular processing. This potential was assessed by suspending human umbilical vascular endothelial cells (hUVECs) within the ELK1 solution prior to co-assembly and growing the tubes described here. Fluorescence microscopy revealed that cells were present both within the assembled ELK1-GO membrane as well as inside the lumen of the corresponding tubes right after co-assembly (Fig. 2g), which is likely a result of cells being either trapped within or adhered to the assembling membrane. Cells were observed to spread and grow for at least 7 days both within the membrane and on the lumen of the tubular structures, indicating that the material is able to support cell survival and growth. To confirm this finding, cell adhesion and proliferation assays were conducted on both sides of ELK1–GO wall of preformed tubes. Remarkably, cells were found to adhere and proliferate at similar levels as those growing on tissue culture plastic (TCP) (Fig. 5a, b), forming a confluent layer on both sides of the membrane (Fig. 5c). To further assess the cell behavior on the tubular structures, VE–cadherin (CD144) was labeled to observe the organization of the intercellular junctions, which are critical for the formation of an intact endothelial monolayer⁴⁵. Confocal images revealed that hUVECs were able to form an integral monolayer on both sides of the ELK1–GO membrane (Fig. 5d).

This notable cell growth and spread on the co-assembled membranes suggests that the hybrid material is cell friendly in vitro. While ELR materials have been shown to support cell growth 46,47, GO is known to be cytotoxic to endothelial cells in vitro at concentrations higher than 100 ng/mL as a result of plasma membrane damage and oxidative stress 48. It is important to keep in mind that GO cytotoxicity depends on the inherent properties of the specific GO used 49. Therefore, we assessed the cytotoxicity of the GO used in this study by conducting experiments using hUVECs in media containing varying GO

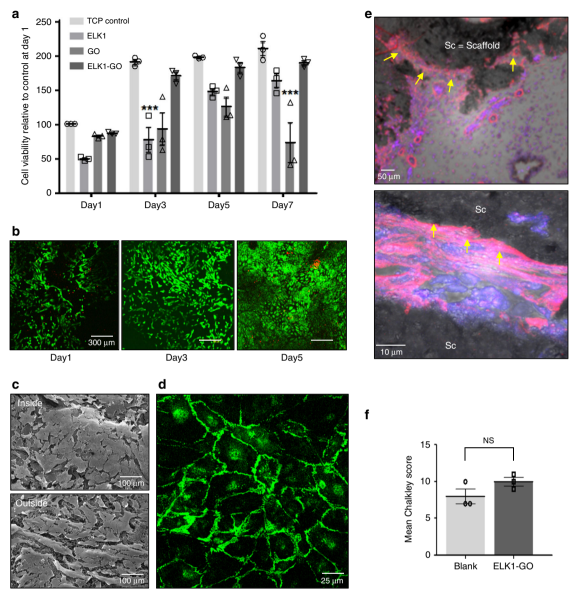


Fig. 5 In vitro biocompatibility and bioactivity of the ELK1-GO membrane. a The applicability of the material was assessed by an MTS assay to test cell viability and proliferation of hUVECS on both sides of the ELK1-GO membrane. The results revealed that cell viability and proliferation on ELK1-GO materials are at least similar to those of cells growing on tissue culture plastic (TCP) for 7 days. Error bars represent \pm s.d. for n = 3. \pm p < 0.05. Two-way ANOVA. **b** Live (green)/dead (red) assay confirmed the proliferation of hUVECs. **c** Scanning electron micrographs demonstrate the formation of an integral endothelial layer on both sides of the ELK1-GO membrane. **d** VE-cadherin (CD144) was labeled to observe the organization of the intercellular junctions and revealed that cells exhibited strong intercellular junction staining, also suggesting the formation of an integral endothelial layer on the ELK1-GO membrane. **e** Histological sections of the ELK1-GO tube implants within a chick chorioallantoic membrane (CAM) model for 7 days highlighting alpha smooth muscle actin (α-SMA, pink), and cell nuclei (blue). The results revealed endothelial cells forming capillary-like structures surrounding the ELK1-GO tubes (yellow arrows). **f** Chalkley count analysis showing a slightly higher level of angiogenesis on tube-containing samples compared to control (blank model) samples. \pm s.d. for n = 3. \pm p < 0.05. One-way ANOVA. NS no significance.

concentrations and found that our GO is toxic above 0.001% (10 μ g/mL) (Supplementary Fig. 14). However, when coassembling and growing ELK1–GO tubes in the presence of hUVECs, we used GO concentrations that are up to 400× higher than this cytotoxic limit, suggesting that the ELK1–GO complex considerably decreases the cytotoxic level of the GO. This decrease may be due to aggregation of GO sheets⁵⁰ and the localization of the ELK1 on the sharp edges of GO sheets⁵¹. To investigate potential cytotoxicity as a result of membrane degradation, we exposed tubes to cell culture media for different time points up to 15 days and used extracts from this media to culture hUVECs. In this case, no cytotoxicity was observed for all time points

(Supplementary Fig. 15a) compared to hUVECs growing in fresh culture media. Furthermore, by physically damaging ELK1–GO tubes suspended in culture media using strong agitation and exposing hUVECs to this suspension for 72 h, cells exhibited higher viability compared to cells exposed to GO sheets at similar levels of concentration (Supplementary Fig. 15b). To further confirm the biocompatibility of the material, we implanted ELK1–GO tubes directly on an ex vivo preclinical chick chorioallantoic membrane (CAM) model⁵² for 7 days and assessed their cytotoxicity and angiogenesis. Using a Chalkley count analysis, similar angiogenesis was observed on both tube-containing samples and control samples (blank model) (Fig. 5f,

Supplementary Fig. 16). Furthermore, immunohistochemistry revealed the presence of capillary-like structures, which in many cases appeared to develop and spread in the vicinity of the ELK1–GO membrane (Fig. 5e). These results are in alignment with previous studies demonstrating the angiogenic potential of GO⁴⁸. However, our approach permits this angiogenic potential while enabling the use of much higher concentrations of GO.

Discussion

We have demonstrated the possibility to exploit multicomponent self-assembly to hierarchically control the interactions between a disordered protein and GO and grow hybrid materials and devices with a spectrum of new functionalities. The system takes advantage of the inherent properties of both GO and ELRs to trigger a diffusion-reaction process that enables disorder-to-order transitions to facilitate GO-disordered protein interactions, supramolecular integration, and hierarchical assembly. Based on both experimental and simulation evidences, we have described the key steps of the underlying molecular mechanism and established rules to grow the material and easily assemble functional devices. The system exhibits a series of properties that emerge from the synergistic interaction between the two components, including remarkable stability, access to nonequilibrium for substantial periods of time, robustness of assembly, biocompatibility, and bioactivity. We have shown how these properties enable its integration with rapid-prototyping techniques to biofabricate functional microfluidic devices by directed self-assembly, opening new opportunities for engineering more complex and biologically relevant tissue engineered scaffolds, microfluidic systems, or organ-on-a-chip devices. Furthermore, our study addresses a major challenge in materials science by demonstrating the possibility to bridge the gap between supramolecular design and functional and robust biomedical engineering.

Method

Chemicals. 4-methylbenzhydrylamine (MBHA) rink amide resin and fluor-enylmethyloxycarbonyl (Fmoc)-protected amino acids were purchased from Merck Millipore. 1-hydroxybenzotriazole (HOBt) hydrate was purchased from Cambridge Bioscience. N,N'-diisopropylcarbodiimide (DIC), dimethylformamide (DMF), dichloromethane (DCM), piperidine, trifluoroacetic acid (TFA), triisopropylsilane (TIS), diethyl ether, acetonitrile (ACN), acetic anhydride and Kaiser Test Kit were purchased from Sigma-Aldrich and used without further purification. Rhodamine B (≥95%, HPLC grade) and paraformaldehyde (95%) were obtained from Sigma-Aldrich. Two kinds of GO (GO-L with product number-777676; GO-S with product number-763705) were obtained from Sigma-Aldrich. The GOs' size distribution can be found in Supplementary Fig. 1. Alexa Fluor™ 488 NHS Ester (Succinimidyl Ester) was obtained from Thermo Fisher Scientific.

Synthesis and characterization of ELRs. ELK0, ELK1, and ELK3 molecules were provided by TP Nanobiotechnology (Valladolid, Spain). Figure 1a shows the sequences, molecular weights, and inverse-phase Tt of the ELRs. ELRs were synthesized by $Escherichia\ coli\ recombinant\ expression\ system.$ The sequence and molecular weights of the polymers were verified using amino acid analysis. Sodium dodecyl sulfate–polyacrylamide gel electrophoresis and matrix-assisted laser desorption/ionization-time-of-flight SIMS were used to carry out the ELRs characterization.

Peptide synthesis and purification. Peptides representative of the single repeat of an individual ELR block were synthesized in a microwave-assisted automated peptide synthesizer (Liberty Blue, CEM) using the standard solid phase peptide synthesis method and Fmoc-protection chemistry. Because the individual blocks are linked together in the ELR molecule through peptide (amide) bonds, synthesized peptides were amidated at the C-terminus and acetylated at N-terminus to resemble the continuation of the amide backbone character and not introducing any additional charges. MBHA rink amide resin was used as the solid support. Amino acid couplings were performed with a mixture of Fmoc-amino acid/HOBt/DIC at a molar ratio of 4:4:4, relative to the resin. Fmoc deprotections were performed with 20% (v/v) piperidine in DMF for 10 min twice. Once all coupling and deprotection reactions were completed, the N-terminal of the peptides were manually capped with 20% acetic anhydride in DMF for 20 min twice. Acetylation

reaction was monitored with Kaiser test for free amines. Peptides were then cleaved from the resin with a mixture of TFA/TIS/H $_2$ O at a volume ratio of 95:2.5:2.5 for 2 h with simultaneous removal of side-chain protecting groups. The cleavage solution was then collected and the excess of TFA removed by rotary evaporation. Cold diethyl ether was added to precipitate the peptide product, which was then collected, washed again with cold diethyl ether and dried under vacuum overnight. Mass of the crude product was analyzed via electrospray ionization mass spectrometry (ESI–MS, Agilent).

Purification of the peptides was performed in an AutoPurification System (Waters) using a preparative reverse-phase C18 column (XBridge, 130 Å, 5 μM , 30 × 150 mm, Waters) and H2O/ACN (0.1% TFA) as mobile phase. Fractions containing the peptides were automatically collected when their exact mass was detected in the SQ Mass Detector (Waters). The collected peptide fractions were lyophilized and stored at $-20\,^{\circ}\mathrm{C}$ until further use.

Mass confirmation for all peptides was performed via ESI–MS and their purity analyzed in an Alliance HPLC system (Waters) equipped with an analytical reverse-phase C18 column (XBridge, 130 Å, 3.5 μ M, 4.6 × 150 mm, Waters) and monitored at 220 mm

Sample preparation (ELRs-GO system). Aqueous suspension of GO (0.1 wt%, $100 \,\mu\text{L}$) was added to a well of 96-well TCP and aqueous solution of the ELRs (2 wt %, $18 \,\mu\text{L}$) was slowly injected into the suspension of GO. The tip of the pipette was allowed to make contact with the bottom of the well before releasing the ELRs solution vertically at a constant speed. All samples were prepared in MilliO water.

Temperature-controlled spectrophotometry. The thermo-responsive behavior of ELK1 at certain concentration (2 wt%) and pH 8 was determined on a temperature-controlled UV-visible spectrophotometer (Agilent Technologies). ELR samples (2 wt%) were prepared in MilliQ water and the pH of the solutions was adjusted with HCl (0.5 M) and NH4OH (1.0 M) prior to heating at 1 °C/min ramping rate. Absorbance of the samples was obtained at $\lambda = 350$ nm.

Zeta potential (ζ). In order to optimize the formation of the ELK1–GO system, the zeta potential of both ELK1 and GO was measured on Zetasizer (Nano-ZS ZEN 3600, Malvern Instruments, UK) at 30 °C under various pH conditions. The concentration of ELK1 and GO used for the measurements is 0.025 and 0.00125 wt %, respectively. The pH values of the two component solutions were adjusted using 0.5 M HCl (at most 3 μ L into 1 mL ELK1 or GO solution) and 1.0 M NH4OH (at most 2 μ L into 1 mL ELK1 or GO solution) and the samples were equilibrated for 10 min at the set temperature prior to the measurement of zeta potential.

Dynamic light scattering (DLS). DLS was performed to measure changes in the particle size of ELK1–GO aggregates at 4 °C (below ELK1's *Tt*), 30 °C (at the *Tt*), and 45 °C (above the *Tt*). The ELK1 and GO were dissolved in MilliQ water at the concentrations of 0.2 and 0.01% separately. The two solutions were mixed in a 1:1 ratio and the particle sizes were measured using Zetasizer (Nano-ZS ZEN 3600, Malvern Instruments, UK). Samples were equilibrated for 10 min at the desired temperature before measurements.

Fluorescence emission. Fluorescence emission was measured on LS 55 spectro-fluorometer (Perkin Elmer). The aqueous solution of GO $(2.5 \times 10^{-3} \text{ wt\%}, 1.5 \text{ mL})$ and the solution of various concentrations of ELRs (1.5 mL) were mixed in a 10 mm path length cuvette at 30 °C. The excitation and emission slits were set at 10 nm. The GO was excited at 255 nm and the emission spectra were collected between 300 and 700 nm (200 nm/min). The fluorescence emission intensity was recorded at 518 nm. The data were fitted into the Benesi-Hildebrand equation (1) in order to determine the association/binding constant (Ka) between GO and ELRs.

$$1/\Delta I = 1/\Delta I_{\rm max} + (1/{\it Ka}[{\it C}])(1/\Delta I_{\rm max}) \eqno(1)$$

where [C] is the concentration of ELRs, $\Delta I = I - I_{\min}$ and $\Delta I_{\max} = I_{\max} - I_{\min}$, where I_{\min} , I, and I_{\max} are the emission intensities of GO considered in the absence of ELRs, at an intermediate ELRs concentration and a concentration of complete saturation, respectively. From the plot of $(I_{\max} - I_{\min})/(I - I_{\min})$ against $[C]^{-1}$ for GO, the value of Ka was determined from the slope.

Circular dichroism (CD). VT-CD measurements were carried out on Chirascan™ CD Spectrometer (Applied Photophysic Limited, UK) from 10 to 40 °C. The solutions of ELK1 (0.01 wt%) were prepared in MilliQ water and incubated at each temperature for 10 min before measurements. A quartz cuvette with 0.1 cm path length was used for the measurements and CD spectra were obtained by signal integrating 10 scans, from 190 to 260 nm at speed of 50 nm/min. Data were processed by a simple moving average and smoothing method.

Fourier transform infrared spectroscopy (FT-IR). FT-IR analysis was conducted on FT-IR spectrometer GX (PerkinElmer $^{\circ}$, Waltham, MA, USA). A solution of ELK1 (2 wt%) in a mixture of D₂O and H₂O (75/25 v/v) and the preformed ELK1–GO membranes prepared in the same solution were properly secured over the IR window before scanning. All samples were incubated and formed at 4, 30,

and 45 °C for 10 min before measurements. The program was set to take the average of 160 scans at a resolution of 2 cm $^{-1}$ after subtracting the background and spectra were obtained at wavenumber $4000-600~\text{cm}^{-1}$ with respect to the absorbance for all samples. In order to quantitatively determine the maximum absorption intensity corresponding to various secondary structures of the ELRs (α-helix, β-sheets, β-turns, and random coils) amide III region (1350–1200 cm $^{-1}$) was analyzed using second derivative of a Guassian and Lorentian curve fittings. The second derivative fingerprints for the secondary structures of the ELRs are as follows: 1220–1250 cm $^{-1}$ for β -sheets, 1250–1270 cm $^{-1}$ for random coils, $1270-1295~\text{cm}^{-1}$ for β -turns, $1295-1330~\text{cm}^{-1}$ for α -helix, as previously suggested by Cai et al 41 .

Scanning electron microscopy (SEM) and wavelength-dispersive spectroscopy (WDS). The microstructures of ELRs-GO and ELK1-GO membranes cocultured with HUVECs were examined by SEM. ELK1-GO membranes with HUVECs were fixed with 4% paraformaldehyde in MilliQ water for 20 min before dehydration while ELRs-GO membranes were dehydrated directly using increasing concentrations of ethanol (20, 50, 70, 90, 96, and 100%). All samples were subjected to critical point drying (K850, Quorum Technologies, UK) prior imaging. The SEM micrographs were captured on Inspect F50 (FEI Comp, the Netherlands) after sputter-coating with gold (10 nm thick). WDS elemental analyses were performed to study the molecular composition of both the inner and outer surfaces of the ELK1-GO membranes. Quantitative Nitrogen elements (nitrogen exists in ELRs not in GO.) were also analyzed using the Inspect F50 (FEI Comp, the Netherlands). All samples consisting only ELRs or GO were prepared for SEM imaging without a prior cross-linking process.

Cryo-transmission electron microscopy (Cryo-TEM). The ELK1 solutions were prepared at 2 wt% in MilliQ water. Grids were vitrified using the Vitrobot MK IV. The Vitrobot chamber was equilibrated to the desired temperature (4, 30, or 45 °C), at 95% relative humidity. Quantifoil R 1.2/1.3 grids were glow discharged in air using the Quorum GloQube* for 60 s, 40 mA. Grid tweezers, grid and pipette tips were preheated to protein aggregate temperature before using. Totally, 3 μ L of sample was applied to the grid, blotted for 5 or 6 s with a blot force of 6. Cryo-TEM imaging was performed on a Titan Krios microscope (Thermo Fisher Scientific, US) operating at 300 kV, using a Falcon III direct electron detector.

Confocal microscopy. The interaction and localization of ELK1 and GO was probed using laser scanning confocal and multiphoton microscopy (TCS SP2, Leica Microsystems, Germany). ELK1 (2 wt%) was dissolved in an aqueous solution of Alexa Fluor* 488 NHS Ester (10^{-6} wt%) and GO were diluted to 0.1 wt% with an aqueous solution of Rhodamine (10^{-6} wt%). All solutions were incubated for 20 min at 30 °C and protected from light. The tubes were fabricated with 50 μ L GO-Rhodamine solution and 10 μ L ELK1-Alexa Fluor solution in a 96-well Petri dish as previously described. Images were acquired at laser wavelengths of 488 and 543 nm which correspond to the excitation wavelength of Alexa Fluor and rhodamine, respectively. Images were further processed using ImageJ.

Small-angle neutron scattering (SANS). The GO suspension and ELK1 were dissolved in H₂O/D₂O (25%/75%) respectively with 0.1 and 2%. SANS measurements were performed on the fixed-geometry, time-of-flight LARMOR diffractometer (ISIS Neutron and Muon Source, Oxfordshire, UK). A white beam of radiation with neutron wavelengths spanning 2.2 to 10 Å was enabled access to Q $[Q = 4\pi \sin{(\theta/2)/\lambda}]$ range of 0.004–0.4 Å⁻¹ with a fixed-sample detector distance of 4.1 m. Solutions (0.4 mL) of individual components were contained in 1 mm path length UV spectrophotometer grade quartz cuvettes (Hellman) while the composite materials were prepared by mixing equal volume (0.2 mL) of both components in a demountable 1 mm path length cuvettes. The cuvettes were mounted in aluminum holders on top of an enclosed, computer-controlled sample chamber at 30 °C. For the variable temperatures (VTs) experiment (especially those involving ELK1 at 4, 30, and 45 °C), a thermostatted circulating water bath was fitted with the sample chamber. Time taken for each measurement was approximately 30 min. All scattering data were normalized for the sample transmission, the backgrounds was corrected using a quartz cell filled with D2O or H2O/D2O (25%/75%) and the linearity and efficiency of the detector response was corrected using the instrument-specific software.

In the present SANS experiments, we consider that the scattering length density (SLD) of the $H_2O/\ D_2O$ (25%/75%) is a volume fraction weighted average of the SLDs of the individual components. Given the SLDs for H_2O and D_2O are $-5.6\times 10^{-7}\ {\rm \AA}^{-1}$ and $6.3\times 10^{-6}\ {\rm \AA}^{-1}$, we determined the SLD of the H_2O/D_2O (25%/75%) is $4.653\times 10^{-6}\ {\rm \AA}^{-1}$. The neutron scattering length densities for the GO, H_2O/D_2O , and ELK1 are summarized in Supplementary Fig. 11c.

Cell culture. Human umbilical vein endothelial cells (hUVECs) (Lonza, Isolated in EGM*-2 Media, C2519A) were cultured in EGM*-2 Media (Lonza, CC-3156 and CC-4176). The medium was changed every 3 days until the cells reached 80% confluency. hUVECs between passage 2 and 4 were used for experiments. The tubes were first washed three times with phosphate-buffered saline (PBS) 8 h after assembly and sterilized with UV for 45 min. Then each tube was placed in a well of

48-well cell culture plate with inner or outer side facing up. The EGM**-2 Media (500 $\mu L)$ containing 50,000 cells was added to each well containing ELK1–GO membranes, coated with ELRs solution (18 μL , 2 wt%), GO (20 μL , 0.1 wt% GO) or on TCP (positive control). The coated wells were incubated for 8 h prior to cell seeding. The cells were incubated at 37 °C and 5% CO $_2$ for different time points for all tests (protocol shown below).

A LIVE-DEAD* Cytotoxicity Assay Kit (Invitrogen, USA) was used to measure the viability of hUVECs seeded on the ELK1–GO membranes. Five replicates of each condition were performed with each assay repeated in triplicate. A stock solution containing calcein AM (1 μ M) and ethidium homodimer (2 μ M) in PBS was prepared according to the assay instructions, and 200 μ L of stock solution was added to each well. Fluorescence images were captured on laser scanning confocal and multiphoton microscopy (TCS SP2, Leica Microsystems, Germany). Viable cells were stained green with calcein AM (ex 495 nm, em 530 \pm 12.5 nm), while dead cells red with ethidium homodimer (ex 528 nm, em 645 \pm 20 nm).

GFP-hUVECs (Fisher scientific, Angio Proteomie GFP-hUVECs, NC0601093, USA) were used to assess the cytotoxicity of GO by culturing >95% confluent GFP-hUVECs in media containing varying GO concentrations $(0,\,0.001,\,0.0025,\,0.005,\,0.01,\,0.25,\,0.05,\,$ and 0.1%) and cultured for 48 h. Fluorescent images were taken at 1, 8, 24, and 48 h to assess the cytotoxicity by morphology and confluency analysis.

GFP-hUVECs were also used to assess the potential cytotoxicity of ELK1–GO degradation products (extracts). We exposed 1 tube fabricated by $100~\mu L$ 0.1% GO and $18~\mu L$ 2% ELK1 to 1 mL cell culture media for different periods of time s (1, 3, 6, 9, 12, and 15 days) and used the extracts from this media to culture >95% confluent GFP-hUVECs for 7 days. Fluorescent images were taken at day 1 and day 7 to assess the cytotoxicity by morphology and confluency. To further assess the potential physical degradation of ELK1–GO, physically damaged ELK1–GO tubes using strong agitation were suspended in culture media and exposing >95% confluent GFP-hUVECs to this suspension for 72 h. Fluorescent images were taken to assess the cytotoxicity.

Immunofluorescence staining. hUVECs on the ELK1–GO membrane were fixed with 4% paraformaldehyde (Sigma, USA), washed and permeabilized with 0.5% Triton X-100 (Sigma, USA), and then rinsed 3 times with PBS. Nonspecific binding sites were blocked by PBS containing 1% bovine serum albumin. The CD144 marker was labeled by incubating the cells at room temperature for 1 h with antirabbit monoclonal VE-cadherin primary antibody (1:400, ab33168, Abcam, UK). Cells were then washed and incubated for 50 min at room temperature in Alexa 488 conjugated anti-Rabbit IgG as Secondary Antibody (1:1000, R37116, Invitrogen, USA). The stained ELK1–GO membranes were then transferred to slides and visualized on a laser scanning confocal and multiphoton microscopy (TCS SP2, Leica Microsystems, Germany) utilizing ×10 and ×40 objectives.

Chick chorioallantoic membrane (CAM) assay. Fertilized chick eggs (Gallus domesticus) were kept in a hatchmaster (Brinsea, UK) incubated at 37.5 °C and humidified with rotation. Twelve (six per group: blank control group and ELK1–GO group) day 1 fertilized eggs were maintained within the hatchmaster. After candling the egg to determine if the egg is fertilized a window was created at day 7 under sterile conditions. A window was created by scoring with a scalpel and an approximately 6 mm square opening created in the outer shell of the egg. The membrane was removed from the underlying CAM vascular membrane. ELK1–GO tube samples were inserted into the window and onto the chorioallantoic membrane. Eggs were transferred to a Hatchmaster incubator and incubated for a duration of 8 days at 37.5 °C 60% humidity without rotation. All procedures were performed in accordance with ethical approval and in accordance with the Animal (Scientific Procedures) Act 1986, UK (Project License number P3E01C456). After 8 days of the CAM culture the implanted samples were harvested.

Immunohistology staining. Slides were first deparaffinized by washing in two changes of xylene (Sigma, UK) and graded ethanol baths (absolute ethanol, 90%, 70%). Antigen retrieval was performed to unmask the antigenic epitope of the tissue sample by boiling the deparaffinized sections in citrate buffer (Vector laboratories, UK) at pH 6.0. Endogenous peroxidase activity was blocked by incubating sections in 3% H₂O₂ solution (Sigma, UK) in PBS at room temperature for 10 min followed by two rinses in PBS. To reduce background staining and any other immunostaining application, the samples were incubated with normal goat serum (5% in PBS, Vector laboratories, UK) to block nonspecific binding sites in a humidified chamber at room temperature for 1 h before staining. After draining the

blocking buffer, 100 μL of diluted primary anti- α -SMA antibody (1:500, ab5694, Abcam, UK) was added to the sections on the slides and incubated in a humidified chamber at room temperature for 1 h, after which the slides were washed twice in PBS. Then, 100 μL of ready-to-use biotinylated anti-mouse/rabbit 1gG secondary antibody (Ready-to-use, PK-7200, Vector laboratories, UK) was applied to the sections on the slides and incubated in a humidified chamber at room temperature for 30 min with the slides washed in PBS after that. Amplification of antigen was achieved using an Elite ABC-HRP Kit (PK-7200, Vector Laboratories, UK) and positive staining was visualized by incubating in a peroxidase substrate solution using a DAB Peroxidase (HRP) Substrate Kit (PK-7200, Vector laboratories, UK).

Analysis of goldner's trichrome staining by Chalkley count. The Chalkley point-overlap morphometric technique is a relative area estimate method to measure the abundance of microvessels in an immunohistochemical sample. A "Chalkley point array graticule" was used to fit onto the eyepiece of a microscope. This graticule consists of a grid that contain 25 random dots which can be rotated 360°. An observer can overlay these dots over structures that have stained positively with goldner's trichrome. The rotational position with the most dots that land on positively stained structures is described as the "Chalkley count" and samples have higher counts are considered to contain a greater abundance of blood vessels. A blank histological slide sample and three ELK1–GO histological slide samples were scoring by this technique.

Co-assembly of ELK1-GO-hUVECs. EGM**-2 Media containing hUVECs (10^5 cells/ml) was used to dissolved the ELK1 (2 wt%). The ELK1-hUVECs media ($10\,\mu\text{L}$) was added into GO ($50\,\mu\text{L}$, $0.4\,\text{wt}$ %) solution to make a tube as previously described. All these co-assembled ELK1-GO-hUVECs tubes were incubated at $37\,^\circ\text{C}$, $5\%\,\text{CO}_2$ for $24\,\text{h}$, $1\,\text{d}$, $3\,\text{d}$, $5\,\text{d}$ prior to LIVE-DEAD** cytotoxicity Assay and SEM procedures as described previously.

3-D printing of ELK1-GO materials. A PAM2 system (Centro Piaggio, Pisa University, Italy) was applied for the 3-D printing of ELK1–GO materials. Blue food dye (5 μ L) was added into aqueous solution of ELK1 (2 mL, 2 wt%) to make the printing procedure visible. For fabricating the different shapes of structures and the 60 μ m diameter small tube, a 65 μ m diameter glass tube tip was used as nozzle to release the solution of the ELK1 and the dye under 4 kPa pressure at a range of speed between 10 and 18 mm/s. The printing nozzle is merged in a container with 0.1% GO MilliQ water solution. All the 3-D pathway was controlled by the Repetier software. A peristaltic pump was used to perfuse 1 v/v green food dye in MilliQ water. For the vertical tube, the perfusion speed is from 4.7 to 8.3 mL/min. For other structures, the perfusion speed was 2 mL/min.

Statistical analysis. GraphPad Prism 5 was applied for data analysis. Student's *t*-test statistical analysis was applied for all the measured data.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The data that support the findings of this study are available from the authors on reasonable request, see author contributions for specific data sets.

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Author contributions

Y.W. and A.M. conceived the project, Y.W. carried out the experiments, A.M. and W.W. supervised the study, B.O. and Y.W. assisted in the performance of SANS experiments and interpretation of results, J.X. and W.L. performed biological characterization, I.K., S.K., D.N., and V.F. conducted computer simulations, A.B, I.C., and N.M.P. conducted the mechanical characterization, F.L.P.B. and G.V. performed the biofabrication, J.K. and R.O.C.O. performed the CAM experimentation, J.F. and M.T. provided the GO and performed characterization of dynamic properties, Y.S. and H.S.A. synthesized the peptides, R.F.T. helped perform the cryo-TEM experiments, and J.C.R.C. provided the ELRs, and Y.W. assisted with SANS.

Competing interests

The authors declare no competing interests.

Additional information

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Supplementary Information

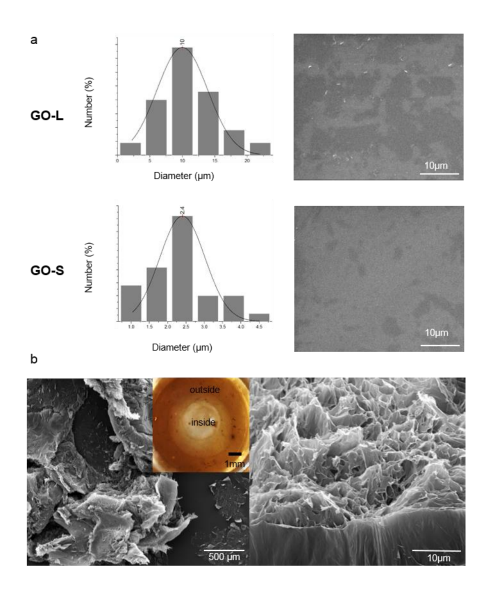
Disordered protein-graphene oxide co-assembly and supramolecular biofabrication of functional fluidic devices

Wu et al

Supplementary Information Contents:

- Supplementary Section 1: Lateral size distribution of the two GOs used with different dimensions and structure of ELK1-GO-S
- Supplementary Section 2: Single repeat of individual ELR block peptides information
- Supplementary Section 3: Gradient of ELK1 from inside to outside of tube
- Supplementary Section 4: Interaction between GO and ELRs with different levels of hydrophobicity and charge
- Supplementary Section 5: Optimal interaction concentration ratio between ELK1 and GO
- Supplementary Section 6: Secondary structure calculation of single repeat of individual ELR block peptides by Circular Dichroism (CD)
- Supplementary Section 7: Interaction between GO and single repeat of individual ELR block peptides
- Supplementary Section 8: Small-angle neutron scattering (SANS) measurements
- Supplementary Section 9: Secondary structure calculation of ELRs Circular Dichroism (CD)
- Supplementary Section 10: Cryo-transmission electron microscopy (TEM) image of ELK1
- Supplementary Section 11: Cytotoxicity measurement of GO concentration
- Supplementary Section 12: Cytotoxicity measurement of ELK1-GO degradation
- Supplementary Section 13: Chalkley Score
- Supplementary Section 14: Statistical analysis of nanotensile mechanical tests
- Supplementary Section 15: Zeta potential (ζ) of ELK1 and GO
- Supplementary Section 16: Ion strength affection of ELK1-GO system
- Supplementary Section 17: Polydispersity index (PDI) of Dynamic Light Scattering (DLS)
- Supplementary Section 18: Computer modelling
- References

Supplementary Section 1. Lateral size distribution of the two GOs used with different dimensions and structure of ELK1-GO-S



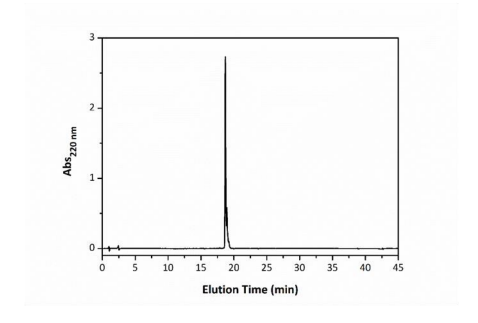
Supplementary Figure 1 a. Lateral size distribution of GO-L ($10.5 \pm 4.5 \,\mu\text{m}$) and GO-S ($2.3 \pm 0.9 \,\mu\text{m}$). b. SEM images of dried xerogel of ELK1-GO-S. Inset: Optical image of the gel of ELK1-GO-S.

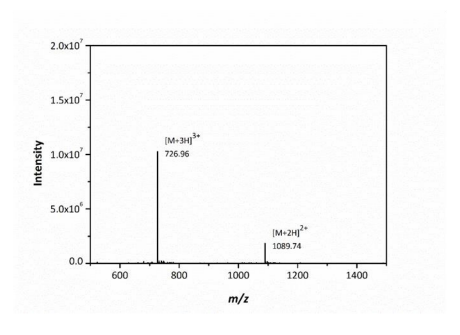
Supplementary Section 2. One repeat single block peptides information

а

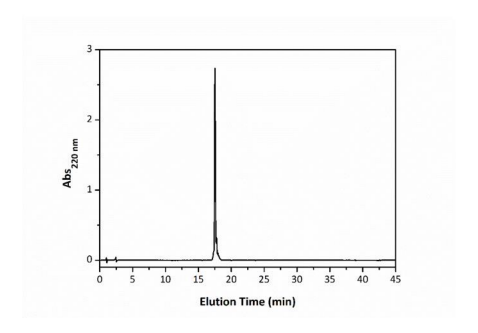
ELR peptides	Sequence	Molecular weight (Da)
ELK ₀₋₁	CH3CONH-(VPGIG-VPGIG-VPGIG-VPGIG)-CONH2	2176.61
ELK1-1	CH3CONH-(VPGIG-VPGIG-VPGIG-VPGIG)-CONH2	2191.63
ELK3-1	CH3CONH-(VPGIG-VPGKG-VPGKG-VPGIG)-CONH2	2221.63

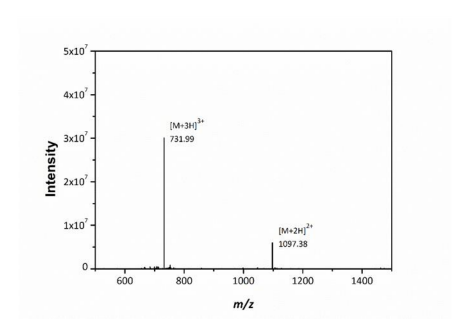
b





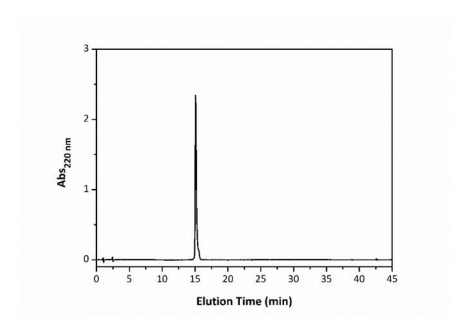
Supplementary Figure 2. Molecular information of ELR peptides. 2b. Analytical RP-HPLC chromatogram (top) of ELK0-1 under the gradient of 98% to 0% H_2O (2% to 100% ACN) with 0.1% TFA from 5 to 35 min showing high purity. ESI-MS spectrum (bottom) of ELK0-1 displaying the expected molecular mass ($C_{102}H_{17}ON_{26}O_{26}$, Mw: 2176.61g/mol)

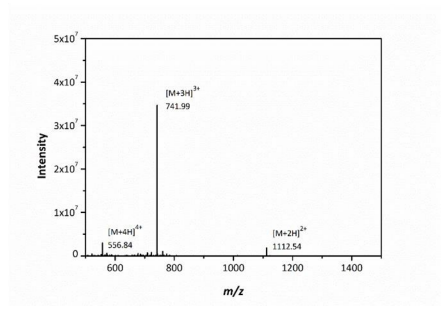




Supplementary Figure 3. Analytical RP-HPLC chromatogram (top) of ELK1-1 under the gradient of 98% to 0% H₂O (2% to 100% ACN) with 0.1% TFA from 5 to 35 min showing high purity. ESI-MS

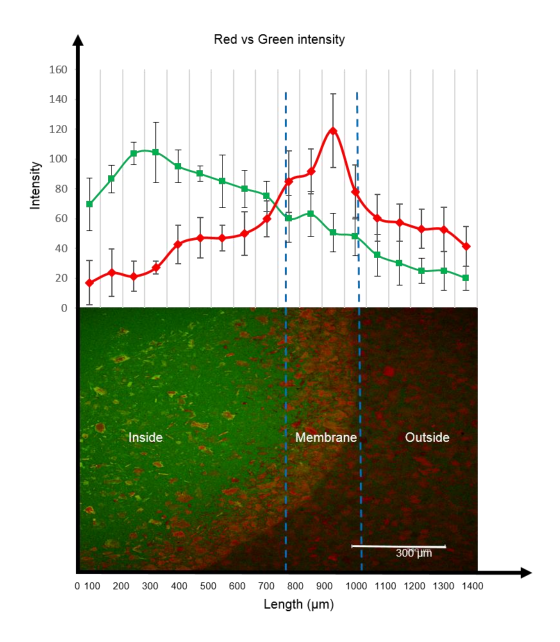
spectrum (bottom) of ELK1-1 showing the expected molecular mass ($C_{102}H_{171}N_{27}O_26$, Mw: 2191.63 g/mol).





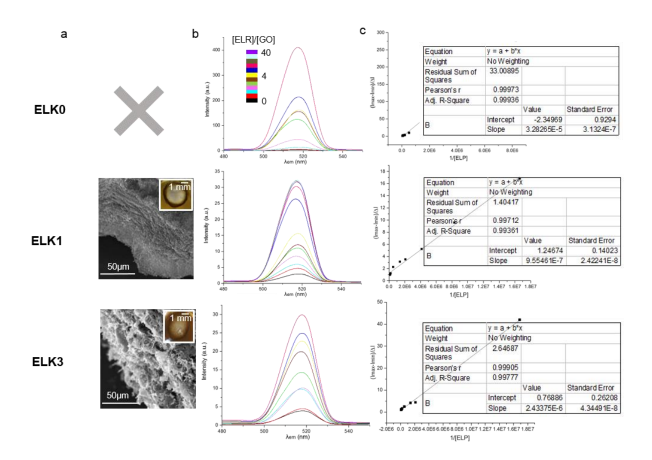
Supplementary Figure 4. Analytical RP-HPLC chromatogram (top) of ELK3-1 under the gradient of 98% to 0% H_2O (2% to 100% ACN) with 0.1% TFA from 5 to 35 min showing high purity. ESI-MS spectrum (bottom) of ELK3-1 showing the expected molecular mass ($C_{102}H_{173}N_{29}O_{26}$, Mw: 2221.63 g/mol).

Supplementary Section 3. Gradient of ELK1 from the tube inside to the outside



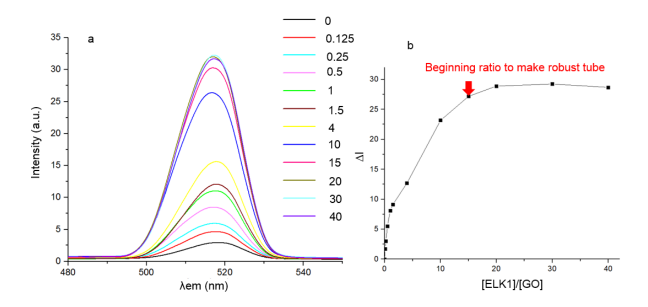
Supplementary Figure 5. Fluorescence intensity calculation of membrane (Green: ELK1. Red: GO) shows the gradual decrease of ELK1 from inside to outside.

Supplementary Section 4. Interaction between GO and ELRs with different levels of hydrophobicity and charge



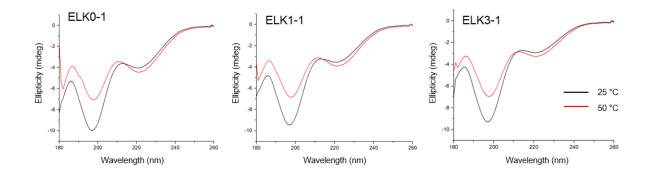
Supplementary Figure 6 a. (ELK0) No stable structure was obtained with ELK0-GO. (**ELK1**) Robust tube with a hierarchical architecture was obtained with ELK1-GO. (**ELK3**) A tube was obtained with ELK3-GO with a rough multilayer membrane at the interface. **4b**. Characterization of GO binding with ELRs by fluorescence emission titration of a fixed concentration of GO (2.5x10⁻³ wt%) and increasing concentrations of ELRs. **4c**. Associated binding constants (Ka) were calculate by Benesi-Hildebrand equation based on the titration results.

Supplementary Section 5. Optimal interaction concentration ratio between ELK1 and GO



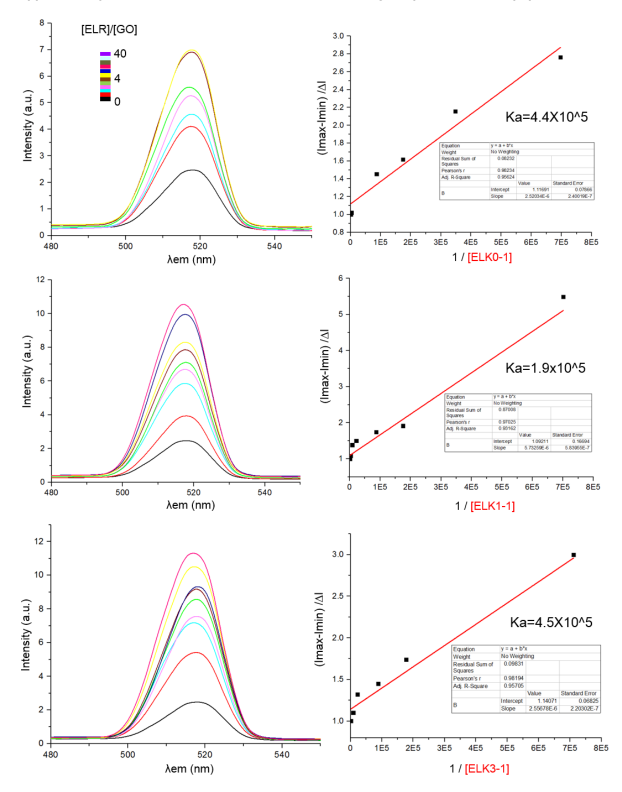
Supplementary Figure 7 a. Characterization of GO binding with ELK1 by fluorescence emission titration of a fixed concentration of GO (2.5x10⁻³ wt%) and increasing concentrations of ELRs. The fluorescent intensity of the peak at 518 nm will continue increasing with increments of the ELK1 concentration for more interaction between them until GO is saturated. **5b.** A graph was made between the [ELK1]/[GO] and peak intensity at 518 nm showing an inflection point at [ELK1]/[GO] equal to 15, the curve reach a plateau above 15.

Supplementary Section 6. Secondary structure measurement of single repeat ELR block peptides by circular dichroism (CD)



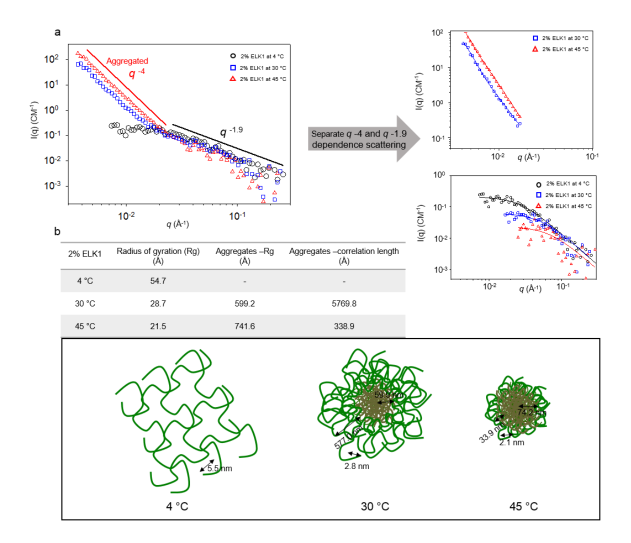
Supplementary Figure 8. CD spectra of the single repeat ELR block peptides show similar little increase of β -turn from 25 °C to 50 °C, while a large percent of random structure was kept in three peptides.

Supplementary Section 7. Interaction between GO and single repeat ELR block peptides

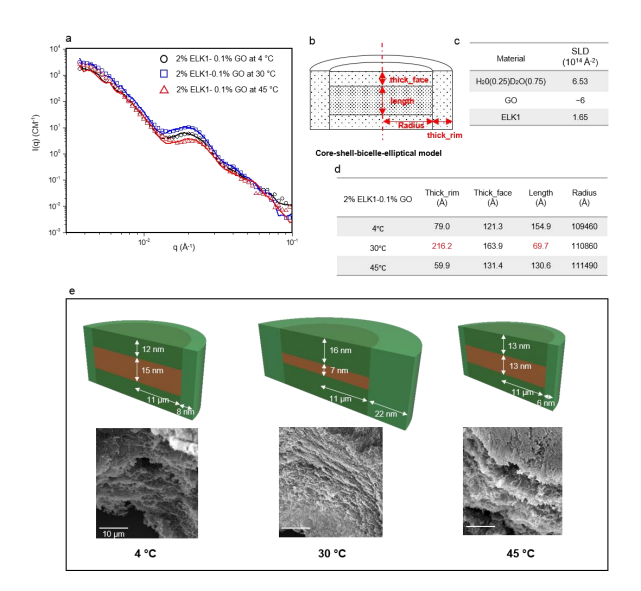


Supplementary Figure 9. Characterization of GO binding with ELRs by fluorescence emission titration of a fixed concentration of GO (2.5x10⁻³ wt%) and increasing concentrations of single repeat ELR block peptides (left). Associated binding constants (Ka) were calculate by Benesi-Hildebrand equation based on the titration results (right).

Supplementary Section 8. Small-angle neutron scattering (SANS) measurements



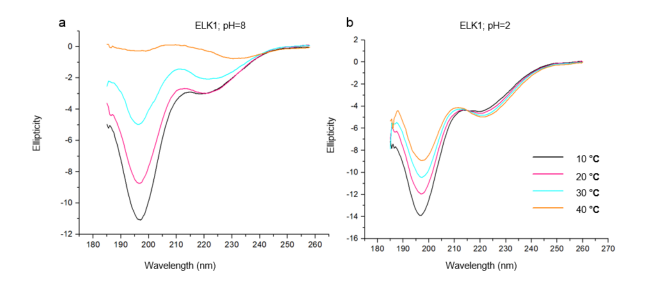
Supplementary Figure 10. SANS results (shapes), fittings (lines), and diagrams for ELK1 at different temperatures (4, 30 and 45 °C). **a (left).** Pronounced q^{-4} dependence scattering appear at 30 °C and 45 °C at low q (ca 0.0035-0.02 Å⁻¹), which demonstrates the scattering is from aggregates with smooth surface. At high q range, the l vs q curves at all temperatures follow a new power law of approximately $q^{1.9}$, which indicates the scattering is from gaussian chains in a dilute environment. **a (right)**. Furthermore, we separated the q^{-4} and $q^{-1.9}$ dependence scattering. A gel-fit model was used to fit q^{-4} porod scattering to obtain the Rg and correlation length of the aggregate structure at 30 °C (blue square) and 45 °C (red triangle) (**top**) and a poly-gauss-coil model was used to fit the ELK1 $q^{-1.9}$ dependence scattering to obtain the Rg of the gaussian chains at different temperatures (**bottom**). **b.** Parameters and diagram were given by the best fitting of poly-gauss-coil/ get-fit model for ELK1 at different temperatures (Green: ELK1 liner structure. Brown: ELK1 aggregates).



Supplementary Figure 11. SANS results (shapes), fittings (lines), fitting parameters, and diagrams for ELK1-GO at different temperatures. **a.** SANS measurement and fitted curves of the hybrid ELK1-GO structure at 4, 30 and 45 °C. Based on physical understanding, confocal microscopy results and using the SasView¹ software program, all of these three curves were fitted well by a core-shell-bicelle-elliptical model, as shown in **b.** The length and thick-rim of the core-shell-bicelle-elliptical model represent the length of the GO and thick_rim of the ELK1 in the ELK1-GO ordered structure, for which the scattering length density² is summarised in **c. d.** Critical parameters of the best fitting of the ELK1-GO structure at different temperatures using the core-shell-bicelle-elliptical model. **e.** Diagrams of the best fitting coreshell-bicelle-elliptical model for the ELK1-GO structures (green: ELK1, brown: GO) at different temperatures detailing the respective sizes and indicating more infiltration of ELK1 within GO at 30 °C

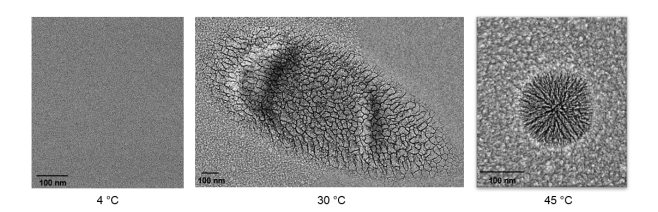
compared to 4 and 45 °C, evidenced by the shorter GO core. This result is supported by the thinner and more-defined layers of the ELK1-GO membrane cross-section at 30 °C as evidenced by SEM.

Supplementary Section 9. Secondary structure measurement of ELRs by CD



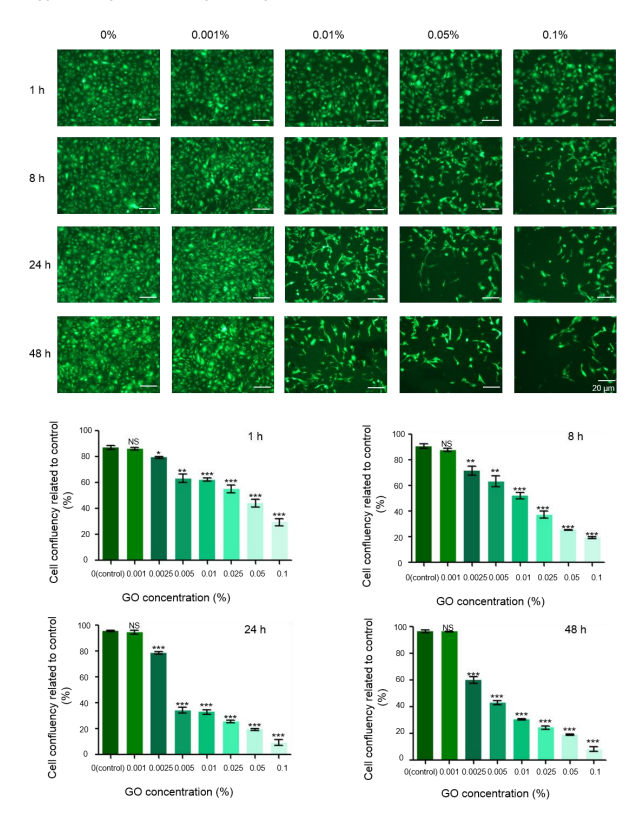
Supplementary Figure 12 a. CD spectra of 0.01% ELK1 in MilliQ water at pH 8 shows an increasing α -helix pattern. 9b. CD spectra of 0.01% ELK1 in MilliQ water at pH 2 shows an increasing β -turn pattern.

Supplementary Section 10. Cryo-transmission electron microscopy (TEM) imaging of ELK1 below, at and above Tt



Supplementary Figure 13. Cryo-TEM images of aqueous solution of ELK1 (2%) at 4°C, 30 °C, and 45 °C. At 4 °C, there are no visible protein aggregates. At 30 °C show the ELK1 form a loosely packed aggregate that is similar in size to the results obtained by SANS. At 45 °C, a more collapsed and densely packed ELK1 aggregate is observed. The presence of more collapsed and denser aggregates of ELK1.

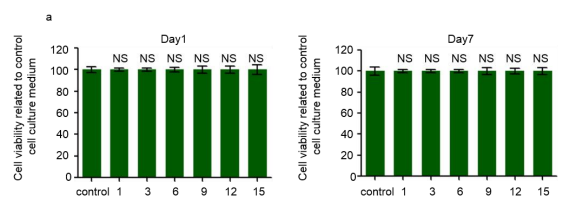
Supplementary Section 11. Cytotoxicity evaluation in function of GO concentration



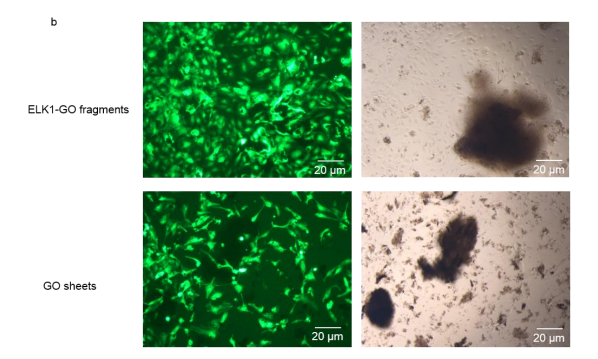
Supplementary Figure 14. Fluorescent images and confluency were achieved at 1, 8, 24 and 48 hr to assess the cytotoxicity of GO by conducting >95% confluent GFP-hUVECs using media containing

varying GO concentrations (0, 0.001, 0.0025, 0.005, 0.01, 0.25, 0.05 and 0.1%). ±s.d. for n=3. *p< 0.05. NS=no significance.

Supplementary Section 12. Cytotoxicity evaluation of ELK1-GO degradation products

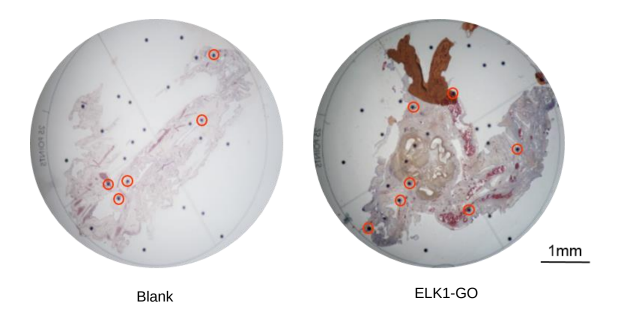


Cell culture medium collected at different time point (Day) Cell culture medium collected at different time point (Day)



Supplementary Figure 15 a. Confluency achieved at Day 1 and Day 7 by conducting >95% confluent GFP-hUVECs using extracted media co-cultured with ELK1-GO for different durations. **12b.** Fluorescent (left) and bright field (right) images of the same view taken from conducting >95% confluent GFP-hUVECs with ELK1-GO fragments and GO sheets only at similar concentration. ±s.d. for n=3. *p<0.05. NS=no significance.

Supplementary Section 13. Chalkley Score



Supplementary Figure 16. Quantification of blood vessel formation after implantation of the ELK1-GO tubes in an *ex vivo* preclinical chick chorioallantoic membrane (CAM) model by Chalkley count using a digital overlay. A digital grid that contained 25 random dots was aligned in 3-5 "hotspot" ROIs (areas covering the most blood vessels in the CAM sample tissue). The digital grid was rotated until the most dots landed on the positively stained vessels. The blank sample exhibited a Chalkley count of 5 while the ELK1-GO sample displayed a Chalkley count of 7 (shown by the red circles).

Supplementary Section 14. Statistical analysis of nanotensile mechanical tests

Tensile tests were performed on three kinds of different tube samples: 2% ELK1-0.05% GO, 2% ELK1-0.10% GO, and 2% ELK1-0.15% GO samples to test if the mechanical properties changed with increasing concentration of GO. 40 µL ELK1 solution was added into 200 µL GO solution in a well of 48-well Petri dish. Rectangular membranes were cut about 10 min after their preparation (Supplementary Figure 17). Ten samples for each composition were tested. Since the core of these structures is made of GO, density was assumed to be equal to 1.8 g/cm³ based on previous studies³. Samples were immediately tested in order to avoid their drying. The main purpose of this test was to characterize the mechanical properties of the bulk material through tensile test. Thus, every type of membrane was tested with a nanotensile machine Agilent Technologies T150 UTM (https://www.agilent.com/home): it is provided with high load resolution (about 50 nN), relatively high maximum load (500 mN) and high displacement resolution (about 0.1 nm).

Membranes were mounted one by one on rectangular paper holders. Both ends of the samples were attached using small amounts of super glue, then each sample holder was mounted in the machine and carefully blocked using two grips. The lateral parts of the paper holder were cut so the machine could perform a traction test and calculate samples mechanical characteristics. Samples had a gauge length of 5 mm and were tested with a strain speed of 0.05%/s. The output of the nanotensile test is a load-displacement curve, from which global properties can be derived. Experimental Young's modulus, strength, strain at break, and toughness modulus were analysed and consequently described by Weibull statistical distribution^{4,5}.

In the following, the Weibull distribution parameters are obtained by assuming the probability of failure F for a sample of volume V under uniaxial stress σ , and related Young's modulus E, strain at break ε , and toughness modulus T. The probability function is expressed as follows (1) (for x equal to σ , E, ε and T):

$$F(x) = 1 - e^{\left[-\frac{V}{V_0}\left(\frac{x}{x_0}\right)^{\alpha_x}\right]} \tag{1}$$

Where x_0 and α_x represent the Weibull's scale and shape parameters, respectively, and V_0 is a unit volume. For a generic quantity x, x_0 is obtained from the y-intercept βx of the best fit equation, being βx equal to $\alpha x^* ln(x_0)$ (from eq. 1), and has the same unit as x. The cumulative probability estimators for experimental tests can be obtained as (2)⁴:

$$F(x_i) = \frac{i - 0.5}{N} \tag{2}$$

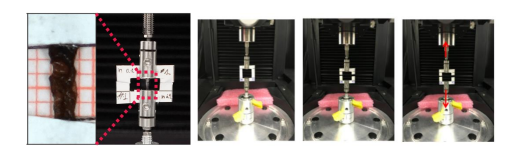
Where N is the number of tests. All the experimental quantities are ranked in ascending order.

For each quantity, σ_0 (or E_0 , ε_0 , T_0) and α_σ (or α_E , α_ε , α_τ) are the Weibull's scale and shape parameters respectively. σ_0 , E_0 , ε_0 and T_0 could be considered also as an index of the mean value of the distribution, while α_σ , α_E , α_ε and α_T are the Weibull modulus.

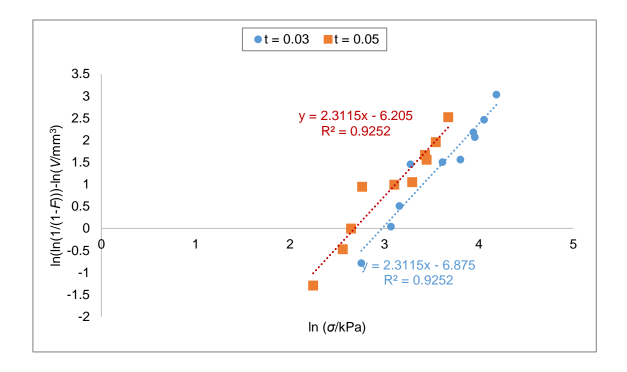
From the statistical analysis, it is possible to state that the Weibull distribution well describes the mechanical properties of samples, as it is possible to deduce from the high values of R^2 . Samples with a concentration of 0.10% of GO show a Young's modulus probability distribution that is smoother than the other types of samples. This means that in this case there is a major probability to have high values of E. Thus, it is possible to assert that 0.10% GO samples are stiffer than the others. Apart from the elastic modulus, the other mechanical properties increase when increasing the GO concentration. However, these preliminary results suggest that samples with 0.10% GO could exhibit higher stiffness without compromising its strain at break and dissipated energy.

Supplementary Table 1 | Sample dimensions, referring to Fig. 3c.

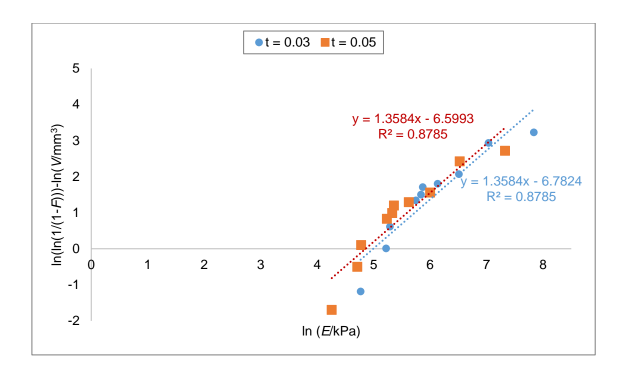
	0.05%		0.10%		0.15%	
h[mm]	0.98 ± 0.15		1.42 ± 0.37		1.67 ± 0.48	
I [mm]	5.00		5.00		5.00	
t [mm]	0.03	0.05	0.03	0.05	0.03	0.05
ρ [g/cm³]	1.8		1.8		1.8	



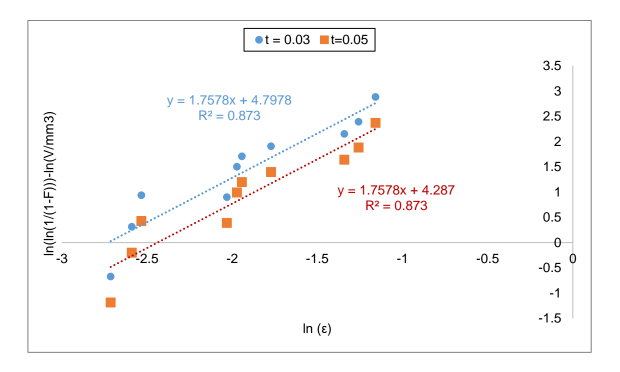
Supplementary Figure 17. Nanotensile test, details of the setup.



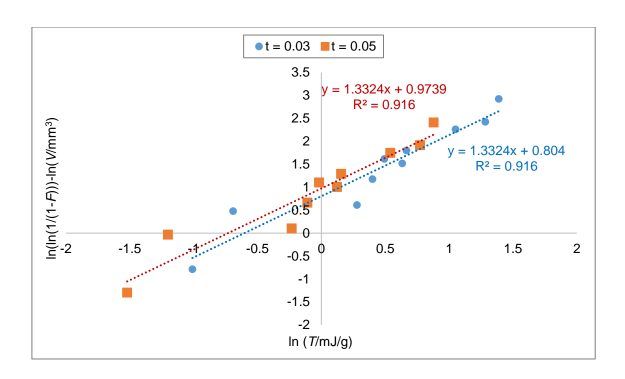
Supplementary Figure 18. Weibull statistics for strength of 0.05% GO samples, by adopting the minimum and maximum thickness (*t*) values.



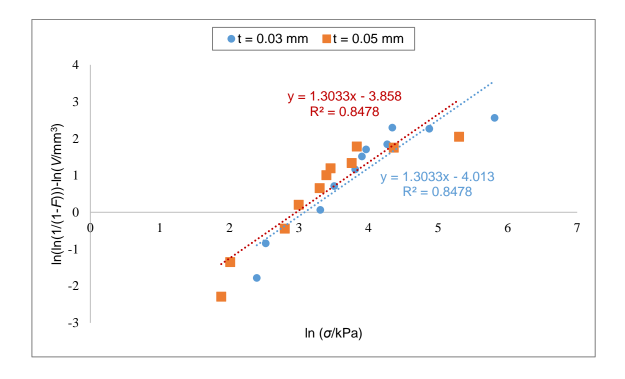
Supplementary Figure 19. Weibull statistics for Young's modulus of 0.05% GO samples, by adopting the minimum and maximum thickness (*t*) values.



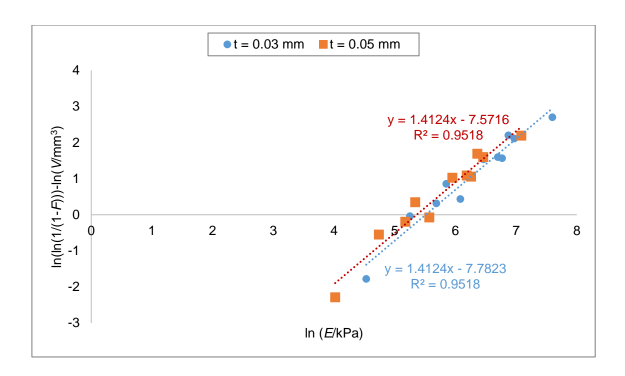
Supplementary Figure 20. Weibull statistics for strain at break of 0.05% GO samples, by adopting the minimum and maximum thickness (*t*) values.



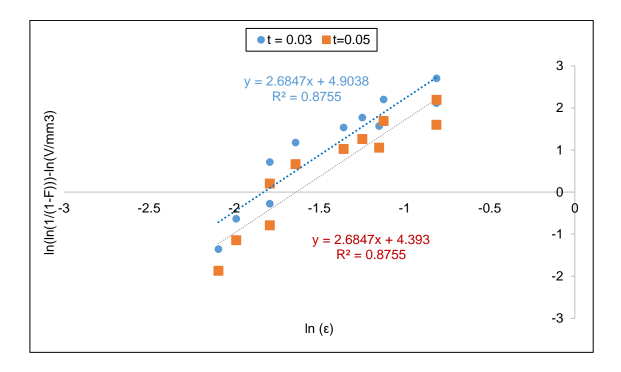
Supplementary Figure 21. Weibull statistics for toughness modulus of 0.05% GO samples, by adopting the minimum and maximum thickness (*t*) values.



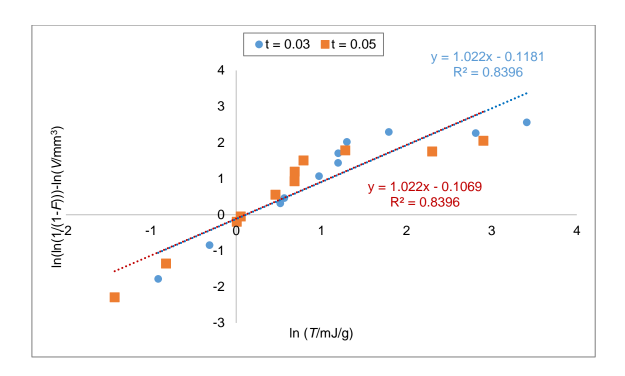
Supplementary Figure 22. Weibull statistics for strength of 0.10% GO samples, by adopting the minimum and maximum thickness (*t*) values.



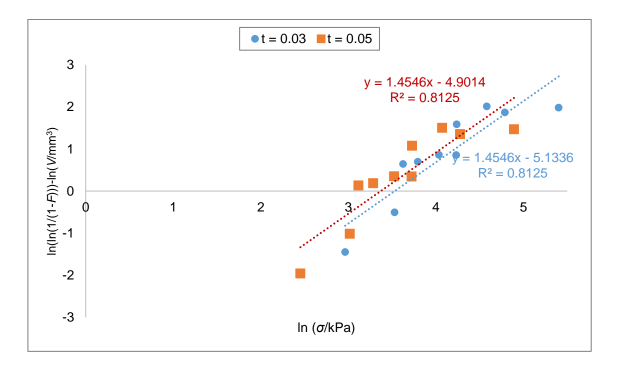
Supplementary Figure 23. Weibull statistics for Young's modulus of 0.10% GO samples, by adopting the minimum and maximum thickness (*t*) values.



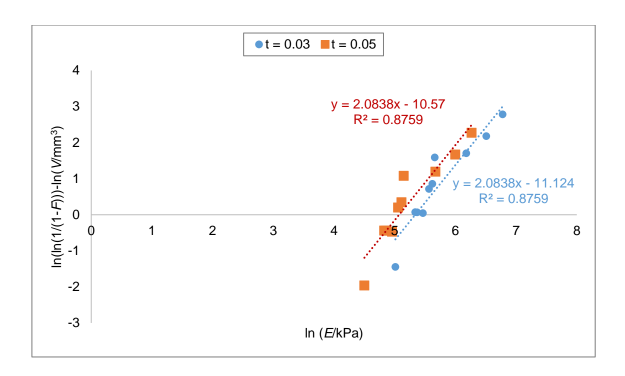
Supplementary Figure 24. Weibull statistics for strain at break of 0.10% GO samples, by adopting the minimum and maximum thickness (*t*) values.



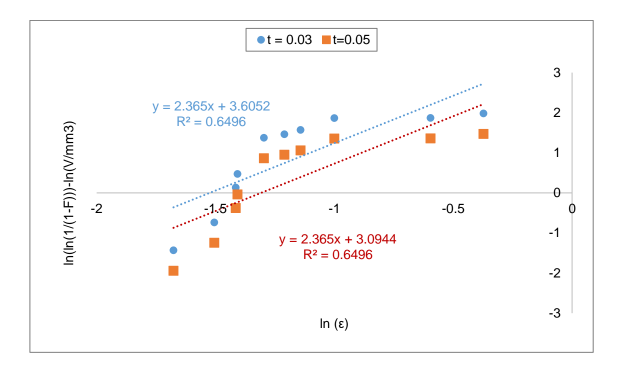
Supplementary Figure 25. Weibull statistics for toughness modulus of 0.10% GO samples, by adopting the minimum and maximum thickness (*t*) values.



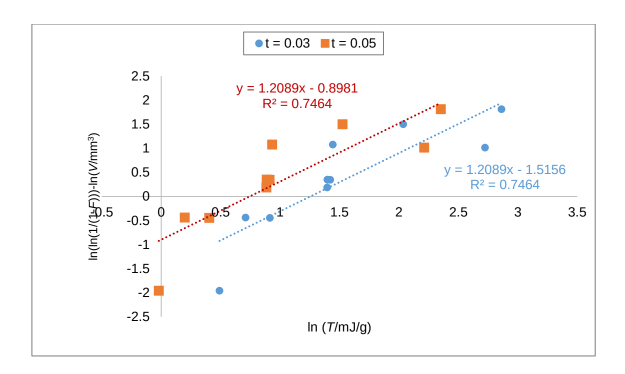
Supplementary Figure 26. Weibull statistics for strength of 0.15% GO samples, by adopting the minimum and maximum thickness (*t*) values.



Supplementary Figure 27. Weibull statistics for Young's modulus of 0.15% GO samples, by adopting the minimum and maximum thickness (*t*) values.



Supplementary Figure 28. Weibull statistics for strain at break of 0.15% GO samples, by adopting the minimum and maximum thickness (*t*) values.



Supplementary Figure 29. Weibull statistics for toughness modulus of 0.15% GO samples, by adopting the minimum and maximum thickness (*t*) values.

Supplementary Table 2 | Weibull statistics parameters for the strength of samples, obtained from the best fit curves. $\sigma_{0 \text{ max}}$ and $\sigma_{0 \text{ min}}$ refer to the minimum and maximum thickness t respectively. α and β are respectively the gradient and the intercept of the interpolant straight lines.

	σ _{0 max} [kPa]	σ _{0 min} [kPa]	α_{σ}	β_{σ} t = 0.03	eta_{σ} t = 0.05	R ²
0.05%	19.58	14.65	2.31	6.88	6.21	0.93
0.10%	21.74	19.30	1.30	4.01	3.86	0.85
0.15%	34.10	29.07	1.45	5.13	4.90	0.81

Supplementary Table 3 | Weibull statistics parameters for the Young's modulus of samples, obtained from the best fit curves. $E_{0 \text{ max}}$ and $E_{0 \text{ min}}$ refer to the minimum and maximum thickness t respectively. α and β are respectively the gradient and the intercept of the interpolant straight lines.

	<i>E_{0 max}</i> [kPa]	<i>E₀ min</i> [kPa]	αε	β_E t = 0.03	β _E t = 0.05	R²
0.05%	147.37	128.78	1.36	6.78	6.60	0.88
0.10%	247.15	212.90	1.41	7.78	7.57	0.95
0.15%	208.16	159.57	2.08	11.12	10.57	0.88

Supplementary Table 4 | Weibull statistics parameters for the strain at break of samples, obtained from the best fit curves. $\epsilon_{0 \text{ max}}$ and $\epsilon_{0 \text{ min}}$ refer to the minimum and maximum thickness t respectively. α and β are respectively the gradient and the intercept of the interpolant straight lines.

	€ _{0 max} [-]	€0 min [-]	αε	β_{ε} $t = 0.03$	β_{ε} t = 0.05	R²
0.05%	0.07	0.09	1.76	4.80	4.29	0.87
0.10%	0.16	0.19	2.68	4.90	4.39	0.88
0.15%	0.22	0.27	2.37	3.61	3.09	0.65

Supplementary Table 5 | Weibull statistics parameters for the toughness modulus of samples, obtained from the best fit curves. $T_{0 \text{ max}}$ and $T_{0 \text{ min}}$ refer to the minimum and maximum thickness t respectively. α and β are respectively the gradient and the intercept of the interpolant straight lines.

	<i>T₀ _{max}</i> [mJ/g]	<i>T₀ _{min}</i> [mJ/g]	α_T	$\beta \tau$ t = 0.03	$\beta \tau$ t = 0.05	R^2
0.05%	0.55	0.48	1.33	-0.80	-0.97	0.92
0.10%	1.12	1.11	1.02	0.12	0.11	0.84
0.15%	3.50	2.10	1.21	1.12	0.90	0.75

Experimental results obtained from tensile tests

Supplementary Table 6 | Mechanical tests results for samples with 0.05% of GO.

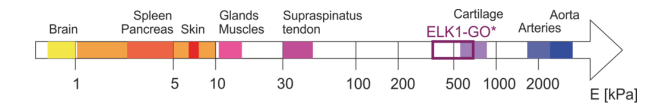
	E _{max} [kPa]	<i>E_{min}</i> [kPa]	σ _{max} [kPa]	σ _{min} [kPa]	ε _m [-]	ε _u [-]	T _{max} [mJ/g]	T _{min} [mJ/g]
Test 1	461.63	276.98	23.51	14.11	0.21	0.28	1.95	1.17
Test 2	117.89	70.74	44.87	26.92	0.28	0.31	3.61	2.17
Test 4	198.36	119.02	21.49	12.90	0.13	0.17	1.32	0.79
Test 5	185.30	111.18	57.66	34.60	0.23	0.26	4.01	2.41
Test 6	343.85	206.31	37.11	22.27	0.11	0.14	1.49	0.90
Test 7	673.10	403.86	52.33	31.40	0.07	0.13	1.88	1.13
Test 9	1137.03	682.22	26.43	15.86	0.05	0.07	0.50	0.30
Test 10	354.08	212.45	65.76	39.45	0.13	0.14	2.86	1.72
Test 11	313.29	187.98	15.72	9.43	0.07	0.08	0.36	0.22
Test 12	2533.44	1520.06	51.47	30.88	0.06	0.08	1.64	0.98

Supplementary Table 7 | Mechanical tests results for samples with 0.10% of GO.

	E _{max} [kPa]	E _{min} [kPa]	σ _{max} [kPa]	σ _{min} [kPa]	ε _m [-]	ε _u [-]	T _{max} [mJ/g]	T _{min} [mJ/g]
Test 1	190.71	114.42	49.59	29.75	0.24	0.29	3.67	2.20
Test 2	1063.90	638.34	27.31	16.39	0.15	0.44	1.76	1.06
Test 3	964.28	578.57	45.10	27.06	0.03	0.33	3.31	1.98
Test 4	870.28	522.17	334.08	200.45	0.22	0.32	30.36	18.21
Test 6	437.63	262.58	12.46	7.48	0.13	0.17	0.73	0.44
Test 7	810.44	486.26	71.44	42.87	0.11	0.12	2.64	1.59
Test 8	347.09	208.25	52.71	31.63	0.15	0.19	3.31	1.99
Test 9	1991.04	1194.62	130.71	78.43	0.08	0.44	16.66	10.00
Test 10	639.44	383.67	76.94	46.16	0.18	0.26	6.00	3.60
Test 11	294.72	176.83	33.34	20.01	0.12	0.17	1.68	1.01
Test 12	92.73	55.64	10.94	6.56	0.12	0.14	0.40	0.24

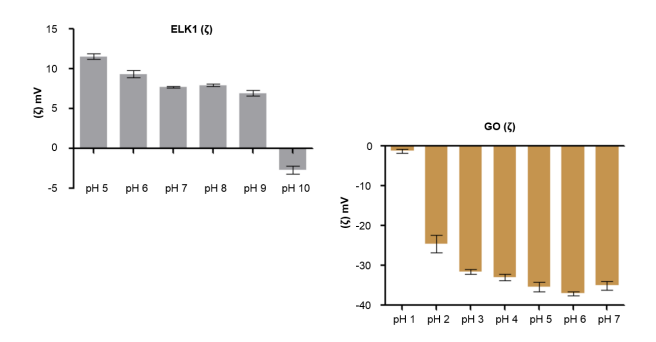
Supplementary Table 8 | Mechanical tests results for samples with 0.15% of GO.

	E _{max} [kPa]	E _{min} [kPa]	σ _{max} [kPa]	σ _{min} [kPa]	ε _m [-]	ε _u [-]	T _{max} [mJ/g]	T _{min} [mJ/g]
Test 1	207.86	124.72	37.52	22.51	0.26	0.27	2.03	1.22
Test 2	286.85	172.11	44.40	26.64	0.29	0.37	4.04	2.42
Test 3	261.16	156.70	119.87	71.92	0.39	0.55	17.52	10.51
Test 4	236.31	141.79	221.89	133.13	0.68	0.69	15.23	9.14
Test 5	670.22	402.13	69.22	41.53	0.12	0.19	4.24	2.54
Test 6	877.23	526.34	97.59	58.55	0.20	0.30	7.66	4.60
Test 7	276.14	165.68	68.86	41.32	0.17	0.22	4.15	2.49
Test 8	482.86	289.72	56.48	33.89	0.18	0.24	4.05	2.43
Test 9	150.05	90.03	19.35	11.61	0.29	0.32	1.63	0.98
Test 11	214.86	128.91	34.04	20.42	0.19	0.24	2.50	1.50



Supplementary Figure 30. (*) Mean values of the Young's modulus for ELK1-GO samples ($E_{0.05\%}$ = 561 kPa, $E_{0.10\%}$ = 683 kPa, $E_{0.15\%}$ = 356 kPa), described by a Weibull distribution with scale parameters E_0 reported in Fig.3c compared with human native tissues^{6–11}.

Supplementary Section 15. Zeta potential (ζ) of ELK1 and GO solutions at different pHs



Supplementary Figure 31. Zeta potential of ELK1 and GO. Measurements were performed at 30 °C in MilliQ water. ELK1 molecules were found to be positively charged at pH values from 5 to 9, but negatively charged at pH 10. GO was found to be negatively charged at all measured pHs.

Supplementary Section 16. Effect of ionic strength on the formation of ELK1-GO system

	C	Concentration of NaCl (g/L) in 2% ELK1 water solution							on	
	T=Tt	8	4	2	1	0.5	0.25	0.125	0	
J c	8									
aCI (g/L solution	4									Robust tube
VaC r sol	2									
n of N water	1									Weak
ratior GO v	0.5									tube
ncentra 0.1% (0.25									
Concentration of NaCl (g/L) in 0.1% GO water solution	0.125									Closed
0	0									membrane

Supplementary Figure 32. Well-defined ELK1-GO tubes can be formed with concentrations of NaCl no more than 1g/L.

Supplementary Section 17. Polydispersity index (PDI) of dynamic light scattering (DLS)

Supplementary Table 9 | PDI of ELK1-GO's DLS

Temperature (°C)	PDI
4	0.39±0.14
30	0.52±0.26
45	0.55±0.09

Supplementary Section 18. Computer modelling.

Molecular models of simulated systems

Our modelling was focused on the interactions between Graphene Oxide (GO) and protein molecules (ELK1) in water. The instrument of choice was classical all-atom Molecular Dynamics (MD) because it allows for precise definition of the system composition and very detailed examination of its behavior at considerable (hundreds of nanoseconds) time.

Therefore, we assume the ELK₁₋₁ (VPGIG VPGIG VPGIG VPGIG VPGIG) are able to reveal the functional groups contributing to the GO–ELK1 interactions, it is enough to simulate a single repeat block. This makes the system of interest readily feasible for meaningful MD. The GO sheets are of µm-s size, which makes their edge length negligibly small compared to their area. We, therefore, used periodic boundary conditions (PBC) to model an "infinitely large" sheets. The sheets homogenously oxidized across the plane. According to the known C: O: H ratio 2:1:1, each second C atom holds a hydroxyl group. Thus, in simulation we can use sheets of the same chemical composition but reduced in size without loss of accuracy, with the help of PBC.

Examined molecular systems and simulation details

MD simulations were performed using free open source software GROMACS 5.1 ¹⁵ on the HPC Midlands supercomputer cluster. The initial configuration of ELK₁₋₁ was generated from the sequence using an on-line platform implementing the I-TASSER based algorithms for protein structure predictions¹⁶. The termini were charged and the lysine residues were protonated; Cl⁻ was chosen as the counter-ion for overall system electroneutrality. The initial GO structure was reconstructed in two stages: first, a graphene sheet was generated using the VMD Nanotube builder plugin¹⁷, then OH groups were attached to randomly chosen atoms using own scripts. Finally, three sheets were placed next to each other to form a stack mimicking the GO lamellas. According to MD simulations of Shih *et*

 $al.^{18}$, the C - C distance between neighbouring sheets was set to 0.8 nm, so that a layer of water molecules can be fitted between the sheets.

Following previous studies of graphene and graphene oxide^{18,19}, the OPLS-AA forcefield²⁰ was employed together with SPC/E water model. In GO, the non-oxidized C atoms were described by the #147 atomtype (naphthalene rings junction) and had no partial charge. For C–O–H groups, the parameters of triatomic alcohols were adopted: atomtypes #175, #171, #172, and partial charges 0.265, -0.73, and 0.465, respectively.

The simulation parameters, common for all the simulated systems, are as follows: 3D periodic boundary conditions; Berendsen thermostat with time constant 2 ps, applied individually to protein and water; particle mesh Ewald method for computing electrostatic interactions; cut-off of van der Waals interactions at 1 nm; constraints on all bonds (LINCS algorithm).

Two kinds of systems were considered:

- 1) A single ELK₁₋₁ peptide in water was simulated with the aim to reach the equilibrium conformation at the given temperature, Supplementary Figure 33a. The parameters were: cell size 8x8x8 nm³, run time 440 ns, temperatures 4°C, 30 °C, 45 °C, pressure of 1 bar, maintained with Berendsen barostat with time constant 1.5 ps. The initial peptide conformation was generated using I-TASSER.
- 2) Three ELK₁₋₁ peptides in water with an 8x8 nm² GO stack were simulated to identify functional groups responsible for the interactions, Supplementary Figure 33b. The parameters were: cell size 8x8x16 nm³, run time 100 ns, temperatures 30 °C, and constant volume. The GO stack was fixed during the simulations, only H atoms were allowed to move. Two initial configurations were used for simulation: i) the ELK₁₋₁ peptides distributed in solution, and ii) the ELK₁₋₁ peptides placed near the GO surface, Supplementary Figure 33c.

Simulation results for an ELK1-1 peptide in water

The secondary structure of an ELK1-1 peptide was computed using DSSP 2.0.4 software²¹, Supplementary Figure 34. The structure does not change considerably during the last 40 ns of the runs, thus it can be considered as converged. No alpha-helices were identified, which agrees with the experimental results (Supplementary Figure 8). Although at equilibrium, the secondary structure does change slowly, it can be seen that the conformational transitions happen at the time scale of hundreds of nanoseconds – microseconds.

The hydrophilic and hydrophobic solvent-accessible surface areas (as computed by VMD on the last 40 ns of the trajectory) at 4 °C were equal to 7.4 nm² and 12.3 nm², respectively, while at 30 °C the values were 6.5 nm² and 13.8 nm², respectively. The values at 45 °C were similar to those at 30 °C. Therefore, the ratio of hydrophilic/hydrophobic parts of the protein surface was approximately the same for all three temperatures, showing no considerable change of hydrophobicity occurring.

Simulation results for three ELK1-1 peptides with GO

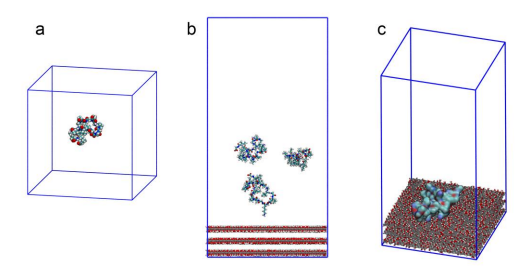
First, consider the runs where ELK₁₋₁ peptides were initially located near the GO surface. The aggregate of peptides did not detach from GO during the whole simulation period. It stayed connected to GO by several hydrogen bonds (Supplementary Figure 35b). This proves that MD correctly reproduces the presence of strong interactions between the peptides and GO.

Now, turn to the runs, started with peptides in the bulk solution. One peptide became adsorbed on the GO surface during the simulation. During 100 ns, sticking of one peptide molecule to another, previously adsorbed on GO, occurred. Thus, we extended the run for additional 100 ns, but no changes occured. The final result was an aggregate of three peptides attached to GO by several hydrogen bonds.

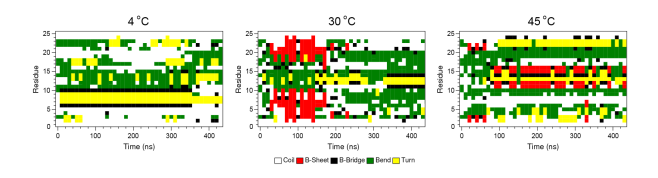
Visual examination of the trajectories clearly shows the main factor ensuring the adhesion of the ELK₁
peptides to GO. It is the protonated amino group of lysine residues, which forms hydrogen bonds with
the OH groups of GO. The rest of the peptide usually almost does not make contacts with GO and is
separated from it by water, which is possible because the lysine's amino group is bonded to a long
hydrocarbon chain that can stretch out. Occasionally, the –CO–NH– peptide bonds of the protein's
backbone also make contacts with GO and form hydrogen bonds, but this is usually not the main factor.

The average numbers of hydrogen bonds between the species (in the runs with initially adhered

peptides) are listed in Supplementary Figure 35a and the typical snapshots from the simulations are shown in Supplementary Figure 35b. Further, the charged termini (COO⁻ and NH₃+) also actively take part in adsorption.



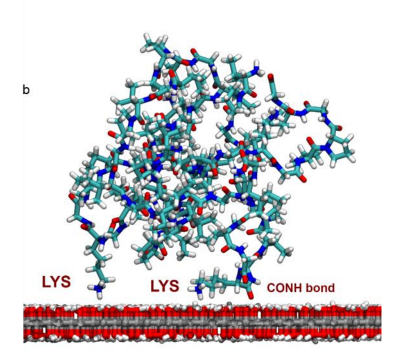
Supplementary Figure 33. Initial configurations for simulated systems. **a.** An ELK1-1 peptide was put in water. This system aimed to simulate the first part including one ELK1-1 peptide in water at 3 different temperatures. **b.** Three ELK1-1 peptides and a stack of GO sheets in water together before running the simulation to identify interactions between them. This system aimed to simulate the second part. **c.** To check the dependence of MD results from initial conditions, three ELK1-1 peptides were initially located near the GO surface to test whether the interaction between ELK1-1 and GO is strong enough to keep this binding status. The system corresponds to the second part, too. The aggregate did not detach from the GO during the whole simulation period being connected with GO by several hydrogen bonds.



Supplementary Figure 34. Investigation of changes in the secondary structure of an ELK1-1 peptide alone in water at different temperatures. The simulation did not show a similar structure change as that of the full-length ELK1.

а

Types of H-bonds	Number of H-bonds
Lysine-GO	2.68
Peptide bonds-GO	2.47
Termini–GO	0.28



Supplementary Figure 35. Hydrogen bond interactions between ELK1-1 and GO. **a.** The average number of hydrogen bonds between ELK1-1-GO. **b.** Instantaneous snapshot from MD simulations of three ELK1-1 peptides with GO indicating the groups responsible for hydrogen bonding.

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