Thixotropic Peptide-Based Physical Hydrogels Applied to Three-Dimensional Cell Culture

Nicola Zanna,‡ Stefano Focaroli,§ Andrea Merlettini,‡ Luca Gentilucci,† Gabriella Teti,‡ Mirella Falconi,‡ and Claudia Tomasini*†‡

†Dipartimento di Chimica Ciamician, Alma Mater Studiorum Università di Bologna, Via Selmi, 2, 40126 Bologna, Italy
‡Dipartimento di Scienze Biomediche e Neuromotorie, Alma Mater Studiorum Università di Bologna, Via Ugo Foscolo, 7, 40123 Bologna, Italy

Supporting Information

ABSTRACT: Pseudopeptides containing the D-Oxd or the D-pGlu [Oxd = (4R,5S)-4-methyl-5-carboxyl-oxazolidin-2-one, pGlu = pyroglutamic acid] moiety and selected amino acids were used as low-molecular-weight gelators to prepare strong and thixotropic hydrogels at physiological pH. The addition of calcium chloride to the gelator solutions induces the formation of insoluble salts that get organized in fibers at a pH close to the physiological one. Physical characterization of hydrogels was carried out by morphologic evaluation and rheological measurements and demonstrated that the analyzed hydrogels are thixotropic, as they have the capability to recover their gel-like behavior. As these hydrogels are easily injectable and may be used for regenerative medicine, they were biologically assessed by cell seeding and viability tests. Human gingival fibroblasts were embedded in 2% hydrogels; all of the hydrogels allow the growth of encapsulated cells with a very good viability. The gelator toxicity may be correlated with their tendency to self-assemble and is totally absent when the hydrogel is formed.

Several techniques may promote the gelation process of these molecules, salt addition, pH variation enzymatic cleavage dissolution in solvent mixtures, ultrasound sonication, although some of these methods may cause cell death.

As LMWGs self-assemble by physical interactions, the hydrogels may show a thixotropic behavior, which means that the gel becomes liquid if a shear stress is applied and then it quickly recovers the solid form on resting. This property allows us to easily inject by syringe the molten hydrogel, which self-adapts in the space inside the injection site and quickly recovers the solid form. Currently, there is no way to predict whether a peptide will be a good hydrogelator; nevertheless, a variety of systematic studies show that amphiphilic peptides and N-protected peptides with aromatic groups are able to form gels in water.

INTRODUCTION

Regenerative medicine is a field of increasing interest as it promotes tissue healing after injuries and diseases. Tissue engineering involves the use of biomaterial scaffolds to create in vitro three-dimensional (3-D) tissue-like structures that simulate the extracellular matrix (ECM) where cells can grow, as often typical bidimensional cell cultures lack the ability to encapsulate extensively studied as materials for cell culture and cell encapsulation and may be injected to act locally in the specific region to be treated, avoiding surgical procedures.

For these applications, the most studied hydrogels are based on natural biopolymers, such as collagen, fibrin, hyaluronic acid, gelatin, chitosan, cellulose, alginate, and agarose. These biomaterials do not meet simultaneously all of the design parameters of an ideal injectable hydrogel (cell adhesion, lifetime, body compatibility, and mechanical strength); moreover, their gelation and mechanical properties cannot be tuned, as they should be used without any chemical modification to maintain their biocompatibility.

In contrast, short peptide chains, that act as low-molecular-weight gelators (LMWGs) as they contain several aromatic rings that favor the formation of hydrogels by means of π-π interactions, are biocompatible and may form hydrogels under several conditions; thus, they enable us to finely tune all of the properties to reach the optimal condition for 3-D cell cultures and good mechanical strength.

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pGlu moieties, together with the presence of aromatic rings, allows the formation of intramolecular interactions that lead to the creation of fibers,\textsuperscript{14,45} which self-assemble to yield a gel.

The pH variation method\textsuperscript{23} used to trigger the gelation process of these compounds allows the formation of gels with low pH values that are not suitable for cell proliferation.

In the present study, we discuss the preparation, characterization, and biological assessment of hydrogels prepared using pseudopeptide gelators, containing the D-Oxd or D-pGlu moiety and selected amino acids. Calcium chloride is used as a trigger to form strong and thixotropic hydrogels at physiological pH, emulating what is usually done to form alginate moieties and selected amino acids. Calcium chloride is used as a coupling and protection/deprotection procedures, as previously reported.\textsuperscript{21,30,39,42}

Amount of CaCl$_2$ to the solutions to test its effect on the hydrogel is crucial to obtain good and reproducible results: if either 1 or 2\% w/w concentration and a stoichiometric amount of aqueous 1 M NaOH. The procedure adopted to form the hydrogel is:

\[ \text{gelators} + \text{CaCl}_2 + \text{NaOH} \]

Thixotropic gels (Figure 1) reached a maximum at a ratio of calcium to carboxylic acid of approximately 2:1 but that good results may also be obtained with different ratios. Thus, we tested the ability of gelators A–C to form hydrogels with different calcium/carboxylic acid ratios: excellent results were obtained with substoichiometric amounts of calcium chloride (Table 1), with the formation of self-supporting and strong hydrogels, even with 1\% w/w gelator concentration, which is the minimum gelator concentration (Figure 2).

| gelator (w/w) | hydrogel | NaOH (equiv) | CaCl$_2$ (equiv) | $T_{gel}$ (°C) | final pH | notes         |
|-------------|---------|-------------|-----------------|---------------|----------|-------------|
| A (1)       | 7       | 2           | 0.6             | 85           | 7.5      | thixotropic |
| A (2)       | 8       | 2           | 0.6             | 100          | 7.5      | thixotropic |
| B (1)       | 9       | 1           | 0.3             | 75           | 7.5      | thixotropic |
| B (2)       | 10      | 1           | 0.3             | 75           | 7.5      | thixotropic |
| C (1)       | 11      | 1           | 0.3             | 50           | 7.5      | thixotropic |
| C (2)       | 12      | 1           | 0.3             | 60           | 7.5      | thixotropic |

*Syneresis occurs on heating.

Hydrogels 7, 8, 11, and 12 show also a thixotropic behavior, as they become liquid if a shear stress is applied and then they quickly recover the solid form on resting, with no variation of the $T_{gel}$.

More information on the nature of hydrogels 7–12 was obtained by scanning electron microscopy (SEM) analysis of aerogels prepared by freeze-drying these samples (Figure 3). Thixotropic gels 7, 8, 11, and 12 furnish aerogels characterized by complex patterns with a rough orientation, whereas hydrogels 9 and 10 furnish aerogels characterized by dense fibrous networks. These observations suggest that when the hydrogel results from the formation of dense fibrous networks, the application of a shear stress destroys these networks, which hardly get quickly reorganized; thus, it inhibits the hydrogel reformation.

To analyze the viscoelastic behavior of the most promising hydrogels, 8, 10, and 12, rheological analyses have been performed to evaluate them in terms of storage and loss moduli ($G'$ and $G''$, respectively) (Table 2 and Figure S2). All of the analyzed hydrogels are characterized by a "solidlike" behavior, that is, the storage modulus is approximately an order of magnitude higher than that of the loss component. Furthermore, the values of $G'$ and $G''$ obtained through strain sweep experiments well-correlate with the previously performed $T_{gel}$ analysis, confirming that the stiffer gel among the obtained ones is hydrogel 8, followed by hydrogels 10 and 12.

Frequency sweep analysis (Figure S2) pointed out that for all of the obtained hydrogels both $G'$ and $G''$ were almost independent from the frequency in the range from 0.1 to 100 rad/s (always with $G' > G''$), confirming the previously discussed solidlike rheological behavior for the analyzed hydrogels.

Step strain experiments were performed to check the thixotropic behavior of 8, 10, and 12 at the molecular level. The strain values within and above the linear viscoelastic (LVE) region were consecutively applied to the hydrogels, which lose their solidlike behavior ($G' < G''$) when the strain is applied.
above their LVE region and quickly go back to a solidlike state \((G' > G'')\) when the strain is applied to the LVE region of the hydrogels (Figure 4).

The results observed for hydrogels 8 and 12 show that they are characterized by a great capability to recover the gel-like behavior and confirm their thixotropic properties at the molecular level. In addition, hydrogel 10 is a thixotropic hydrogel from a molecular point of view, although it does not fully recover the solidlike behavior, when the strain level goes back within the LVE region.

Cytotoxicity and cytocompatibility studies were carried out on diluted solutions of gelators A–C. Human fibroblasts were grown for 24 h in the presence of gelators A–C at increasing concentrations up to 5 mM (about 0.3 w/w concentration), which is always much lower than the concentration needed to form hydrogels.

The toxicity of gelators A–C was evaluated by the lactate dehydrogenase (LDH) assay. The results show a low toxicity of B solutions up to a concentration of 5 mM with a cell viability of 95% with respect to the control. In contrast, the cell viability decreases upon using C and A solutions (72.2 and 72.7% of living cells at a concentration of 5 mM, respectively) after 24 h of incubation (Figure 5). We previously noticed that gelator B tends to form dense fibrous networks, so the very low toxicity could be correlated with the tendency of the gelator to self-assemble. This effect has been recently observed for amyloid precursor proteins (APPs) that exist as soluble oligomers and are extremely neurotoxic. Moreover, several studies have shown that the soluble pool of Aβ is better correlated to cognitive decline than the insoluble pool.

Bearing this preliminary result in mind, we evaluated the behavior of living cells trapped in the hydrogels. The fibroblasts were embedded in 2% hydrogels prepared in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 1% fetal calf serum (FCS), penicillin (50 UI/mL), and streptomycin (0.05 mg/mL) under the same conditions as those for 8, 10, and 12. Gelators A–C efficiently promoted the formation of hydrogels from this medium, although it is not pure water, thus showing their versatility in hydrogel formation. This operation was simplified by the hydrogels’ thixotropic behavior: the gels were prepared and shaken to recover the sol state; thereafter, the cells were rapidly embedded and the resulting solution was allowed to stand for 6 h to reform the hydrogels. The overall results show a reduced toxicity of all of the hydrogels, which well-correlate with the 3-D network formation. Thus, the viability did not statistically differ in hydrogels 8 and 10 (compared to that in the previous experiments) after 24 h of culture, whereas the toxicity of 12 drastically decreased. After 7 days of incubation, when the 3-D networks are completely formed and the hydrogels still maintain their shape, the number of living cells increases for the three samples (Figure 6).

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These results are in agreement with our previous observations on gelator solutions and indicate that the toxicity of gelators is completely eliminated when the molecules are self-assembled, showing no substantial differences with the behavior of the most commonly used gelators for pharmaceutical and medical applications (such as gelatin,55−57 alginate,5,7 chitosan,58,59 agarose,14,60,61 and hyaluronate62,63).

Finally, the suitability of the hydrogels for long-term cell culture and therapeutic approaches has been confirmed by the evaluation of the NAD(P)H-dependent cellular oxidoreductase mitochondrial enzymes’ activity using 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) as a substrate: the presence of formazan crystals in the cells indicates a good cell viability after 7 days of culture (Figure 7), thus confirming that these hydrogels may be used for 3-D cell culture useful in regenerative medicine.

■ CONCLUSIONS

The preparation, characterization, and biological assessment of hydrogels prepared using pseudopeptide gelators, containing the D-Oxd or the D-pGlu moiety and selected amino acids, have been studied and reported. A substoichiometric amount of calcium chloride was used as a trigger, as it allows the formation of strong and thixotropic hydrogels at physiological pH. This behavior was observed at the macroscopic level and confirmed at a molecular level by rheological analyses, which were used to assess the viscoelastic behavior of the most promising hydrogels in terms of storage and loss moduli.

Then, cytotoxicity and cytocompatibility studies were carried out on diluted solutions (up to 5 mM) of gelators A−C, followed by the biocompatibility evaluation of hydrogels 8, 10, and 12. The results show low toxicity of both the gelator solutions and the hydrogels. In addition, after 7 days, when all of the 3-D networks are completely formed, the number of living cells increases in all of the samples and the gelator toxicity is totally eliminated. This result is in agreement with
what has been recently observed for the APPs, as the soluble pool of Aβ better correlates with the cognitive decline than the insoluble pool.

**EXPERIMENTAL SECTION**

**Materials.** All chemicals and solvents were purchased from Sigma-Aldrich, VWR, or Iris Biotech and were used as received. Acetonitrile was distilled under inert atmosphere before use. MilliQ water (Millipore, resistivity = 18.2 mΩ cm) was used throughout. DMEM supplemented with 10% FCS, penicillin (50 UI/mL), and streptomycin (0.05 mg/mL) was purchased by Life Technologies. The LDH kit was purchased from Thermo Fisher Scientific.

**Synthesis of HO-d-Oxd-L-Phe-CO(CH2)7CO-l-Phe-d-Oxd-OH A.** Compound A was synthesized from d-Thr, azelaic acid, and Boc-l-Phe-OH following a multistep procedure in solution, reported in ref 26.

**Synthesis of Fmoc-l-Tyr-d-Oxd-OH B.** Compound B was synthesized from d-Thr and Fmoc-l-Tyr(t-Bu)-OH following a multistep procedure in solution, reported in ref 34.

**Synthesis of Fmoc-l-Phe-d-pGlu-OH C.** Compound C was synthesized from d-pGlu and Fmoc-l-Phe-OH following a multistep procedure in solution, reported in ref 17.

**Conditions for Gel Formation with Gelator A.** A portion of gelator A (5–10 mg, depending on the final concentration, ranging from 1 to 2% w/w) was placed in a test tube (diameter: 8 mm), and then, MilliQ water (0.5 mL) and 2 equiv of 1 M aqueous NaOH were added. The mixture was stirred until complete compound dissolution. Different amounts of CaCl2 were added to the solution under rapid stirring (see Tables 1 and S1 for details), and then, the tubes were allowed to stand quiescently until gel formation, which occurred after about 10 min.

**Conditions for Gel Formation with Gelator B or C.** A portion of gelator B or C (5–10 mg, depending on the final concentration, ranging from 1 to 2% w/w) was placed in a test tube (diameter: 8 mm), then MilliQ water (0.5 mL) and 1 M aqueous NaOH (1 equiv) were added, and the mixture was stirred until sample dissolution. Different amounts of CaCl2 were added to the solution under rapid stirring (see Tables 1 and S1 for details), and then, the tubes were allowed to stand quiescently until gel formation, which occurred after about 10 min.

**Conditions for \( T_{gel} \) Determination.** \( T_{gel} \) was determined by heating the test tube (diameter: 8 mm) containing the hydrogel sample and a glass ball (diameter: 5 mm, weight: 165 mg) on the top of it. When the hydrogel is formed, the ball is suspended atop. The \( T_{gel} \) is the temperature at which the ball starts to penetrate inside the gel. Some hydrogel samples melt, producing a clear solution, whereas in other cases, the gelator shrinks and water is ejected, as synerysis occurs.

**Aerogel Preparation.** Some samples of hydrogels 7–12 were freeze-dried using a Benchtop Freeze Dry System LABCONCO 7740030 with the following procedure: the hydrogel (0.5 mL) was prepared in an Eppendorf test tube at room temperature. After 16 h, the samples were immersed in liquid nitrogen for 10 min and then freeze-dried for 24 h in vacuo (0.2 mBar) at −50 °C.

**Morphological Analysis.** Scanning electron micrographs of the samples were recorded using a Hitachi 6400 field emission gun scanning electron microscope.

**Rheology.** Rheology experiments were carried out on an Anton Paar rheometer MCR 102 using a parallel plate configuration (25 mm diameter). Experiments were performed at a constant temperature of 23 °C controlled by the integrated Peltier system and a Julabo AWC100 cooling system. To keep the sample hydrated, a solvent trap was used (H-PTD200). The amplitude and frequency sweep analyses were performed with a fixed gap value of 0.5 mm on hydrogel samples prepared directly on the upper plate of the rheometer once the gelation reaction was completed. The samples were prepared the day before the analysis and left overnight at a controlled temperature of 20 °C to complete the gelation process. Oscillatory amplitude sweep experiments (\( \gamma \): 0.01–100%) were carried out to determine the LVE range at a fixed frequency of 1 rad/s. Once the LVE range of each hydrogel was established, frequency sweep tests were performed (\( \omega \): 0.1–100 rad/s) at constant strain within the LVE region of each sample. Thixotropic experiments were conducted on hydrogels 8, 10, and 12 by applying consecutive deformation and recovery steps. The deformation step was performed by applying to the gels a constant strain above the LVE region of each sample for a period of 7 min. The recovery step was performed by keeping the sample at a constant strain within the LVE region for 7 min. The cycles were performed multiple times at a fixed frequency of 1 rad/s.

**Isolation and Culture of HGFs.** HGFs were obtained from healthy patients subjected to gingivectomy of the molar region. Informed consent was obtained from each patient. Immediately after the removal, the tissues were washed in phosphate buffer, cut in small pieces, and placed in DMEM, supplemented with 10% FCS, penicillin (50 UI/mL), and streptomycin (0.05 mg/mL), at 37 °C in a 5% humidified CO2 atmosphere. After the first passage, the HGFs were routinely cultured in DMEM supplemented with 10% FCS and were not used beyond the fifth passage.

**Evaluation of Gelator Cytocompatibility.** Cells were seeded in a 96-well plate at a density of 20,000 cells/cm². The gelators were solubilized at concentrations from 5 μM to 5 mM
in culture media, and cytotoxicity of the compounds was evaluated by measuring the cytosolic LDH activity in the culture supernatants.

The LDH assay was performed after 24 h of incubation using a commercial LDH kit following the manufacturer’s recommendations. LDH is a cytosolic enzyme present in many different cell types. The plasma membrane damage releases LDH into the cell culture media. Extracellular LDH in the media can be quantified by a coupled enzymatic reaction in which LDH catalyzes the conversion of lactate to pyruvate via NAD$^+$ reduction to NADH. Diaphorase then uses NADH to reduce a tetrazolium salt to a red formazan product that can be measured at 490 nm. The level of formazan formation is directly proportional to the amount of LDH released into the medium, which is indicative of the cytotoxicity. To determine total LDH activity, cells from the positive control group were treated with 1% Triton X-100. The optical density in each well was measured using a spectrophotometer microplate reader (model 680; Bio-Rad Lab. Inc., CA) at a wavelength of 490 nm. Each experiment was performed three times, and four replicate cell cultures were analyzed in each experiment.

**Cell Seeding and Cytotoxicity Test.** Gelators A, B, or C (10 mg) were solubilized in 0.5 mL of DMEM supplemented with 1% FCS, penicillin (50 UI/mL), and streptomycin (0.05 mg/mL) under sonication at room temperature during 15 min. Once liquid solutions were obtained, 1 M NaOH (2 equiv for A and 1 equiv for B or C) followed by 0.2 M CaCl$_2$ (0.6 equiv for A and 0.3 equiv for B or C) were added to the solutions, with the immediate formation of hydrogels that rested at room temperature without agitation for 30 min. To allow cell seeding, the gel was heated at 37°C and thoroughly mixed with 100 μL of cell suspension at a concentration of 2 × 10$^5$ cells/mL of gel and the gel/cell systems were transferred into different wells of a 12-well plate and maintained under controlled atmosphere (5% CO$_2$, 37°C) for 24 h and 7 days. After each experimental point, the gel/cell constructs were transferred into 1.5 mL vials, centrifuged at 160 g for 10 min, and the supernatants were used to check the cytotoxicity by the LDH assay described above. Light microscopy observation of living cells was performed using the MTT assay. After each experimental point, 100 μL of a MTT stock solution was added into each well to reach a concentration of 1 mM. After that, the samples were incubated for 4 h at 37°C in a controlled atmosphere and the resulting blue formazan crystals were observed in living cells.

### ASSOCIATED CONTENT

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.7b00322.

Physical properties of the hydrogels obtained with gelators A–C and a stoichiometric amount of CaCl$_2$; photographs of the samples of hydrogels prepared with gelators A–C and CaCl$_2$ in different concentrations; strain dependence and frequency dependence of the storage modulus and loss modulus for hydrogels 8, 10, and 12 (PDF)

### AUTHOR INFORMATION

**Corresponding Author**

*E-mail: claudia.tomasini@unibo.it.*

**ORCID**

Luca Gentilucci: 0000-0001-9134-3161

Claudia Tomasinii: 0000-0002-6310-2704

**Author Contributions**

*N.Z. and S.F. contributed equally.

**Author Contributions**

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

**Notes**

The authors declare no competing financial interest.

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