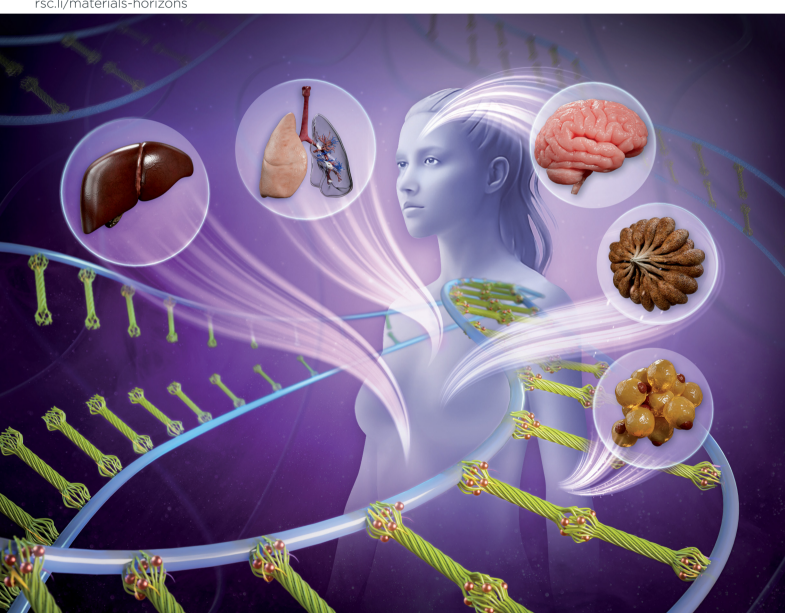
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Nano-enabled dynamically responsive living acellular hydrogels†

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As a key building block of mammalian tissues, extracellular matrices (ECMs) stiffen under shear deformation and undergo cell-imparted healing after damage, features that regulate cell fate, communication, and survival. The shear-stiffening behavior is attributed to semi-flexible biopolymeric ECM networks. Inspired by the mechanical behavior of ECMs, we develop acellular nanocomposite living hydrogels (LivGels), comprising network-forming biopolymers and anisotropic hairy nanoparticle linkers that mimic the dynamic mechanical properties of living counterparts. We show that a bifunctional dynamic linker nanoparticle (nLinker), bearing semi-flexible aldehyde- and carboxylate-modified cellulose chains attached to rigid cellulose nanocrystals converts bulk hydrogels to ECM-like analogues via ionic and dynamic covalent hydrazone bonds. The nLinker not only enables the manipulation of nonlinear mechanics and stiffness within the biological window, but also imparts self-healing to the LivGels. This work is a step forward in designing living acellular soft materials with complex dynamic properties using bio-based nanotechnology.

1. Introduction

Engineering dynamically responsive hydrogels that mimic the mechanical and biological properties of tissues¹⁻³ may enable

characterization, aldehyde titration curve of DAMC, time sweep rheological measurements of NCH, LivGels pore size, differential modulus (K') versus oscillatory shear stress for LivGels, containing 2.25 wt% nLinker, and strain amplitude tests of LivGels, containing varying nLinker and Ca²⁺ concentrations. See DOI: https://doi.org/10.1039/d4mh00922c

New concepts

It is non-trivial to convert conventional, static hydrogels to those that mimic the dynamic properties of extracellular matrices (ECMs) and tissues. The recent decade has witnessed the quest to engineer "living materials" that mimic the characteristic features of biological systems, such as the strain-stiffening behavior, anisotropy, and self-healing, driven by the need for innovations in biomedicine and beyond. Acellular living materials are an emerging class of engineered living materials, aiming to replicate the complexity of biological materials without using any living cells. Our research presents a new concept in converting static gels to those that partially mimic the ECM and tissue properties via anisotropic hairy nanoparticle linkers (nLinker). We develop novel acellular living hydrogels (LivGels) that are engineered using the nLinker and a biopolymer via tailoring polymer-polymer and polymer-nLinker dynamic interactions. Unlike conventional hydrogels, the LivGels, inspired by the structural organization of ECM networks and the healing potential of cells, enable shear-stiffening and self-healing traits. This conceptually novel design allows precise control of the nonlinear mechanics and stiffness of self-healing LivGels. The LivGels may open a new avenue for developing dynamic soft materials for regenerative engineering, tissue and disease models, organs-on-a-chip platforms, additive manufacturing, soft robotics, and active materials.

living soft materials. The majority of native mammalian tissues comprise non-living extracellular matrices (ECMs) and living cell ensembles. The ECM dynamically responds to external physical stimuli and stiffens under shear. 4,5 Accordingly, nonlinear shear-stiffening mechanics are crucial for tissue structural support and cell functions, such as adhesion, migration, and differentiation.6

Tuning biopolymer concentration has been the main approach to tailor the nonlinear mechanics of ECM-mimetic soft materials, 7-10 which may inevitably alter other hydrogel properties, including mechanical strength, swelling behavior, degradation rate, and biocompatibility. Additionally, synthetic hydrogels that mimic the mechanical properties of ECMs are undermined by the concerns associated with biocompatibility, immune response, and toxic degradation byproducts. 11 As a mechanical-biological trait, self-healing is also highly desirable

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in fabricating living hydrogels to enable autonomous morphological and mechanical restorations following reoccurring deformation and defects.10

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Developing multifunctional living acellular hydrogels with ECM-like mechanics and self-healing properties remains an ongoing challenge. Most living tissues have hierarchical and anisotropic structures, partially arising from the multiscale three-dimensional (3D) organization of ECM constituents. 12,13 The isotropy of bulk polymeric hydrogels limits their ability to faithfully replicate the structural and mechanical features of ECMs. To address this shortcoming, nanoparticles have been included in 3D polymeric networks, yielding anisotropic nanocomposite hydrogels (NCH). 14,15 Besides using anisotropic nanoparticles, nanoparticle-polymer crosslinking has also been tailored via non-covalent and/or covalent bond formation. Polymer-nanoparticle inclusions have widely been investigated in terms of microscale network dynamics and macroscale mechanical and rheological properties; 16-18 however, a limited number of NCH with tunable ECM-mimetic mechanical characteristics, particularly strain-stiffening, exist. As an example, NCH with strain-stiffening and self-healing properties have been engineered via incorporating tannic acid-coated cellulose nanocrystals (CNC) into synthetic poly(vinyl alcohol)-borax dynamic networks. 19 Other kinds of NCH with tunable strainstiffening properties have been developed via manipulating the nature of polymer-nanoparticle crosslinks and density, albeit using synthetic polyisocyanide and anisotropic iron oxide nanorods.20,21

To the best of our knowledge, there has been no report of naturally derived NCH with anisotropic particles that mimic the dynamic characteristics of native ECM (i.e., strainstiffening) or tissues (i.e., self-healing) and enable the control of these properties. Hairy CNC (HCNC) are a newly emerged member of the nanocellulose family, comprising a rod-like crystalline body with disordered cellulose chains (hairs) at both ends. 22,23 The hairs contribute to the unique anisotropic surface chemistry and architecture. We hypothesize that chemically tailored HCNC, bearing a combination of carboxylate and aldehyde moieties on the hairs, can link polymer chains in NCH matrices via reversible ionic and covalent bonds, resulting in controlled strain-stiffening and self-healing properties. These bifunctional nanoparticle linkers are called nLinker.

Here, we aim to develop acellular nanocomposite living hydrogels (LivGels) with nonlinear mechanics and selfhealing properties. Dihydrazide-modified alginate (DH-ALG) is synthesized as the matrix biopolymer to provide two types of active sites, i.e., carboxylate groups and hydrazide for ionic and dynamic covalent bond formation, respectively. To impart dynamic properties to bulk alginate hydrogels, we develop an anisotropic nLinker, bearing aldehyde and carboxylate groups. The network is engineered via varying the nLinker concentration and binding mechanisms (dynamic covalent bonding and/or ionic crosslinking). Structure-property relationships in the NCH are investigated to develop a phase diagram, identifying NCH compositions that yield LivGels.

2. Experimental section

2.1. Materials

Bleached, delignified softwood kraft pulp sheets were provided by Resolute Forest Products Inc., Canada, and used as the starting material for the nLinker synthesis. Medium viscosity (MV) alginic acid sodium salt from brown algae was obtained from MilliporeSigma, USA. Sodium (meta)periodate (NaIO4, >99%), sodium chlorite (NaClO₂, 80%), sodium chloride (NaCl, >99.5%), hydrogen peroxide (H₂O₂, 30 wt%), sodium hydroxide (NaOH, ACS reagent, >97%), hydrochloric acid (HCl, ACS reagent, 37%), ethylene glycol (Reagent plus, >99%), hydroxylamine-hydrochloride (NH2OH·HCl, Reagent plus, 99%), adipic acid dihydrazide (ADH, NH₂NHCO(CH₂)₄CONHNH₂, \geq 98%), calcium chloride dihydride (CaCl₂·2H₂O, ≥99%), poly-_L-lysin (PLL, 0.1 w/v% in water), Whatman filter papers (Grade 1, circles), and mineral oil (heavy) were purchased from MilliporeSigma, USA. AFM stainless steel disks and mica sheets (V1 grade) were obtained from SPI supplies (USA) and Ted Pella Inc. (USA), respectively. Deuterium oxide (D₂O, 99.9%) was supplied by Cambridge Isotope Laboratories, Inc., USA. Anhydrous ethanol (200 proof) was purchased from KOPTEC, USA. All material synthesis and experiments were conducted using either ultrapure Milli-Q water (resistivity \sim 18.2 M Ω cm at T = 25 °C) or deionized (DI) water, as specified.

2.2. nLinker synthesis

The nLinker was prepared according to our previously established method for HCNC synthesis with minor modifications.^{24,25} The softwood pulp sheets (5 g) were torn into thin pieces with approximate dimensions of 2 cm \times 2 cm, and pulp fragments were soaked in DI water (500 mL) while stirring with an overhead stirrer (SH-II-6C, Faithful Instrument CO., China) overnight. Afterwards, water was removed via vacuum filtration using a nylon cloth (pore size: 20 µm), and the collected wet pulps were suspended in a NaCl solution (325 mL, 1 M). Following 5 min mixing and thoroughly wrapping the beaker with aluminium foil, NaIO₄ (6.60 g) was added to selectively oxidize the vicinal diols of cellulose to dialdehyde groups. The reaction mixture was stirred at ambient temperature for 42 h. Once completed, ethylene glycol (5 mL) was added and stirred for 10 min to deactivate unreacted NaIO₄. The oxidized pulps (dialdehyde modified cellulose, DAMC) were then vacuum filtered and rinsed with DI water at least 5 times (250 mL each time) to remove residual chemicals and byproducts.

To convert half of the DAMC aldehyde groups to carboxylate groups and isolate nLinker, DAMC was further oxidized with NaClO₂. To this end, DAMC fibrils were suspended in DI water (250 mL, including the water in the wet DAMC), followed by adding NaClO₂ (4.22 g) and NaCl (14.12 g). H₂O₂ (4.22 g) was then added dropwise within 2 min after which the mixture was stirred at 100 rpm for 12 h. The pH was adjusted to 5.0 \pm 0.2 using a NaOH solution (0.5 M) during the reaction until pH remained almost constant at 5. The bifunctional fibrils were collected via centrifugation at 8000 \times g for 20 min and rinsed with an ethanol solution in DI water (70 v/v%, 100 mL) four times. The fibrils were suspended in DI water (500 mL) and heated at 80 °C for 2 h, followed by centrifugation at 8000 \times g for 20 min to remove the unfibrillated cellulose. The nLinker particles in the supernatant were precipitated via adding a nonsolvent (ethanol, volume ~1.5 times the supernatant volume), collected by centrifugation at 8000 \times g for 20 min, and purified via dialysis (Spectra/Por membranes, $M_{\rm W}$ cutoff = 8-10 kDa) against DI water for 1 day. The nLinker concentration was increased to 5 wt% via evaporation at 40 °C under constant airflow for further use. The concentration was determined by oven-drying a known volume of nLinker dispersion at 60 °C, followed by weighing.

2.3. Matrix biopolymer (DH-ALG) synthesis

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DH-ALG was synthesized via successive periodate oxidation and Schiff base reactions. A periodate-mediated oxidation reaction was conducted to functionalize alginate (ALG) with dialdehyde groups, yielding dialdehyde-modified alginate (DA-ALG, a reactive intermediate).²⁶ DA-ALG synthesis was started by dispersing MV sodium ALG (5 g) in ethanol (50 mL), followed by adding DI water (200 mL). Ethanol, as a non-solvent, breaks up agglomerates, resulting in a homogenous distribution of ALG in water.²⁷ Once fully dissolved, NaIO₄ (5.7 g) was added to the ALG solution in an aluminium foil-wrapped beaker while stirring by the overhead stirrer (speed = 100 rpm) at room temperature. Given that NaIO₄ is a photosensitive oxidizer, the reaction was performed in the dark.²⁸ After 1 h, the reaction was quenched by adding ethylene glycol (5 mL) and stirring for 10 min. To purify the resulting DA-ALG, NaCl (5 g) was added to the solution to screen the negatively charged carboxylate groups, facilitating precipitation. The DA-ALG was subsequently rinsed with an ethanol solution in DI water (70 v/v%, 800 mL each time) three times, and the precipitated DA-ALG was dissolved in DI water (250 mL). Further purification was conducted via dialysis (Spectra/Por membranes, Mw cutoff = 8-10 kDa) against DI water for 1 day. To prevent microbial contamination, DA-ALG was stored at 4 °C until further use.

Next, the dialdehyde groups were reacted with hydrazide via a Schiff base reaction to form hydrazone bonds, yielding DH-ALG.²⁹ ADH (0.9 g, ~5 mmol) was dissolved in a DA-ALG solution (1 wt%, 100 mL, ~5 mmol aldehyde), and a Schiff base reaction was carried out for 5 h at room temperature. The ADH amount was calculated based on a 1:1 molar ratio of the aldehyde group: ADH. The resulting DH-ALG solution was then dialyzed (Spectra/Por membranes, $M_{\rm W}$ cutoff = 8-10 kDa) against DI water for 12 h. DH-ALG was stored at 4 $^{\circ}\text{C}$ and used within 3 days to minimize hydrazone bond hydrolysis.³⁰

2.4. NCH preparation

NCH were formed via (i) nLinker-mediated dynamic covalent crosslinking of DH-ALG or (ii) simultaneous nLinker-mediated and ionic crosslinking of DH-ALG. The nLinker-crosslinked NCH (1 mL) were prepared via mixing a DH-ALG (5 wt%) solution, the nLinker (5 wt%) dispersion, and Milli-Q water at varying ratios to yield a constant final DH-ALG concertation of 2 wt% and nLinker concentrations ranging from 1.25 to 2.5 wt%. The biopolymer-nLinker NCH were mixed using a

positive displacement pipette (Microman E M1000E, Gilson, OH, USA) for ~ 30 s and maintained at room temperature for 5 min prior to any characterization. To prepare the nLinkerionically-crosslinked NCH, the nLinker (5 wt%, 10 mL) dispersion was mixed with 0.0875 mL, 0.175 mL, or 0.350 mL of a CaCl₂ solution (5 M) to obtain a carboxylate: Ca²⁺ molar ratio of 1:0.5, 1:1, or 1:2, respectively, resulting in 4.95, 4.91, or 4.83 wt% of final nLinker concentrations. After 5 min of incubation. Ca²⁺-loaded nLinker dispersions were mixed with the DH-ALG solutions (5 wt%) to reach a total volume of 1 mL. The final DH-ALG concentration was maintained constant at 2 wt%, and the final nLinker concentration was varied (1.25, 1.75, 2, or 2.5 wt%) by adding Milli-O water, similar to those in the nLinker-crosslinked NCH. Note that NCH with a 1:0.5, 1:1, or 1:2 molar ratio of carboxylate: Ca²⁺ are referred to as 9, 18, or 36 mM Ca²⁺, respectively. For the NCH preparation, Milli-Q water was used to dilute the nLinker dispersion when required.

2.5. Materials characterization

2.5.1. Functional group content measurements. Conductometric titration was performed to quantify the aldehyde (CHO) group content of DAMC, nLinker, and DH-ALG, as well as the carboxylate (COO⁻) group content of nLinker and DH-ALG using the automated Metrohm 907 Titrando (Metrohm AG, Switzerland). The concentration of aldehyde groups was measured via an oxime formation method.31 Briefly, a known amount of DAMC (50 mg), nLinker (30 mg), or DH-ALG (30 mg) was added to DI water (100 mL) while being magnetically stirred. The pH of DAMC suspension and a NH2OH·HCl solution (20 mL, final concentration = 5 wt%) was independently adjusted to 3.5 using NaOH or HCl solutions. The oxime reaction was initiated by mixing the DAMC suspension and NH₂OH·HCl solution, producing HCl, which was titrated using a NaOH solution (10 mM) at a rate of 0.1 mL min⁻¹. The endpoint was obtained when the pH was stabilized at the initial pH (3.5). The aldehyde group content was calculated based on the volume of NaOH consumed to neutralize the released HCl during the reaction of aldehyde groups with NH2OH·HCl according to eqn (1).

$$C_{\rm CHO} = \frac{V_{\rm NaOH} \times M}{m_{\rm DAMC}} \tag{1}$$

where C_{CHO} is the aldehyde group content per unit mass of DAMC (mmol g^{-1}), V_{NaOH} is the NaOH volume (mL), M is the molarity of NaOH (M), and m_{DAMC} is the DAMC dry mass (g).

To measure the carboxylate group content, a known amount of nLinker or DH-ALG (20 mg dry mass) and NaCl solution (2 mL, 20 mM) were added to 100 mL of Milli-Q water. The pH of nLinker dispersion or DH-ALG solution was then decreased to 3.2 using a HCl solution (0.1 M). The dispersion was titrated by a NaOH solution (10 mM) up to pH ~ 11 at a titration rate of 0.1 mL min⁻¹. The carboxylate group content was calculated based on the volume of NaOH required to neutralize the weak acid (i.e., carboxylate groups) using eqn (2).

$$C_{\text{COO}^{-}} = \frac{V_{\text{NaOH}} \times N}{m} \tag{2}$$

where C_{COO}^{-} is the carboxylate content per unit mass of nLinker or DH-ALG (mmol g^{-1}), and m is the nLinker or DH-ALG dry mass (g).

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2.5.2. Atomic force microscopy (AFM) imaging. The morphology and dimensions of nLinker were investigated using AFM. Imaging was carried out using an atomic force microscope (Bruker Dimension Icon I, USA), equipped with a silicon nitride probe (Bruker ScanAsyst-Air, USA). The experiments were conducted using the PeakForce Tapping mode. To prepare AFM specimens, a freshly cleaved mica sheet (diameter = 10 mm) was adhered to a stainless-steel disc (diameter = 15 mm). The negatively charged mica was then covered with a PLL solution (0.1% w/v) to minimize the electrostatic repulsion between the nLinker and mica. After ~15 min, the excess PLL was rinsed with Milli-Q water (100 µL) five times, followed by drying at room temperature for ~ 30 min. Once dried, a small droplet of nLinker dispersion (~10 μL, 0.1 wt%) was deposited onto the mica sheet and left to dry overnight. The nLinkercoated mica sheet was then rinsed with Milli-Q water (100 µL) five times to eliminate any non-adsorbed particles or debris and air-dried prior to imaging. The acquired images were processed using NanoScope Analysis software (Version 1.8, Bruker, accessed via Penn State Materials Characterization Lab, MCL). The nLinker length and width was determined via manually assessing over 50 particles using the Gwyddion software (Version 2.49, accessed via the MCL).

2.5.3. Attenuated total reflection-Fourier-transform infrared (ATR-FTIR) spectroscopy. The characteristic chemical bonds of nLinker, DA-ALG, DH-ALG, and NCH were identified using a FTIR spectrometer (Fisher Scientific, USA) according to the Bouguer-Beer-Lambert law.³² The spectrometer was equipped with a single bounce diamond ATR accessory. Prior to the analysis, samples were frozen at -80 °C, followed by sublimating the ice at 0.01 mbar overnight using a FreeZone benchtop freeze dryer (Labconco, USA). The dried materials were directly placed on the ATR crystal, and the maximum pressure was applied by lowering the pressure clamp while the incident angle was 45°. The spectra were obtained by averaging 100 scans in a wavenumber range of 4000 to 500 cm⁻¹ (resolution = 6 cm^{-1}). The absorbance was obtained *via* referencing the spectrum of clean bare diamond and then converting to transmittance.

2.5.4. Hydrodynamic size and ζ -potential measurements. To determine the hydrodynamic equivalent size of nLinkers, our established method was followed.³³ Measurements were carried out on a dynamic light scattering (DLS) instrument (Malvern Zetasizer Nano series, UK) at a fixed scattering angle of 90° and 25 °C. The nLinker concentration was adjusted to 0.1% w/v using Milli-Q water, followed by pipetting 70 μL of it into a low-volume quartz cuvette (ZEN2112, Malvern, UK). The Z-average values, i.e., cumulants mean, were reported as the hydrodynamic equivalent size.

To determine ζ -potential, the electrophoretic mobility of nLinker was measured using a Nano ZS Zetasizer (Malvern instrument, UK) at room temperature and pH = 6.5. Before initiating the measurements, the nLinker dispersion was

diluted to 0.1% w/v using Milli-Q water, followed by pipetting a desired volume (900 µL) into disposable folded capillary cells (Malvern, UK). To apply the effect of nLinker rod shape to the ζ-potential value, Oshima's mobility expression was used.³⁴ Since the $\kappa a \sim 1$ (κ is the Debye-Hückel parameter, and a is the rod-like nLinker width, ~5 nm, obtained from the AFM image analysis), the Henry's function $f(\kappa a) \sim 0.5$, 35 and the following equation was used to calculate the ζ-potential using the electrophoretic mobility. Note that the ionic strength was calculated by considering the counter ions (Na+) of nLinker carboxylate groups.

$$\mu_{\rm av} = \frac{\mu_{\parallel} + 2\mu_{\perp}}{3} = \frac{\varepsilon_{\rm r}\varepsilon_0}{3\eta} \zeta[1 + 2f(\kappa a)] \tag{3}$$

where μ_{\parallel} and μ_{\perp} are the electrophoretic mobility of cylindrical particles in the parallel and perpendicular direction to the electric field, respectively, ε_r denotes the relative dielectric permittivity of solvent, ε_0 is the dielectric permittivity in a vacuum (vacuum permittivity), and η denotes the electrolyte solution viscosity.

2.5.5. Proton (¹H) nuclear magnetic resonance (NMR) spectroscopy. To verify and quantify the aldehyde and hydrazide groups on DH-ALG, ¹H NMR spectroscopy was performed using an NMR instrument (Bruker Avance III, MA, USA), operated at 600 MHz and equipped with a triple resonance inverse (TCI) cryoprobe. Freeze-dried ALG, DA-ALG, or DH-ALG (15 mg) were separately dissolved in D₂O (1 mL) for 2 h at room temperature. The spectra were acquired via averaging a total of 128 scans. The characteristic peaks were assigned to the respective functional groups. The area under the corresponding peaks, which was proportional to the number of hydrogen atoms, was measured using TopSpin software (Bruker, Version 4.0.7).³⁶

2.6. Rheological analyses

Rheological properties were measured using an AR-G2 rheometer (TA Instruments, DE, USA), equipped with a parallel plate geometry with an upper plate diameter of 20 mm and a truncation gap of 1000 µm. Freshly prepared NCH were sandwiched between two parallel plates. After trimming the excess hydrogel, the air-exposed area was covered with a mineral oil to prevent water evaporation from the gel during the experiments. To assess the strain-induced stiffening of NCH, an oscillatory amplitude sweep test was conducted at a constant frequency of 1 rad s^{-1} while varying the strain from 0.1 to 200%. To investigate the self-healing of NCH, recovery tests were conducted, wherein the dynamic moduli (i.e., storage modulus, G', and loss modulus, G'') were measured by applying alternating strain of 1% and 500% at time intervals of 100 s and a constant oscillation frequency (1 rad s^{-1}). The time sweep test was performed on freshly prepared NCH at a constant frequency of 1 rad s⁻¹ and strain of 0.1% for 15 min. Temperature was maintained at $\sim\!20~^\circ\text{C}$ in all measurements.

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2.7. Statistical analyses

Data were reported as the average of three independent samples $(n = 3) \pm \text{standard deviation (SD)}$, unless otherwise specified. GraphPad Prism (Version 9.4.1, MA, USA) was used for all statistical analyses. One-way (Fig. 4b, c and Fig. S4a, ESI†) or two-way (Fig. 4g, h and Fig. S4b, ESI†) analysis of variance (ANOVA), followed by the Tukey post hoc test was conducted to determine the statistical significance. The pvalues < 0.05 were considered statistically significant (*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001).

3. Results and discussion

3.1. LivGel design

LivGels are dynamically responsive bio-based NCH inspired by the mechanical responsiveness of ECM, specifically nonlinear strain-stiffening, as well as the self-healing capability of tissues. Fig. 1 presents the design rationale for LivGels, their building blocks, and strategies to control the nonlinear mechanical properties while imparting other functionalities, such as selfhealing. Fig. 1a shows examples of human soft tissues and their main components, including ECM and cell ensembles, which have strain-stiffening and self-healing properties, respectively. LivGels are prepared via incorporating the bifunctional nLinker in reactive DH-ALG, prepared from semi-flexible ALG, as shown in Fig. 1b. ALG, the matrix biopolymer of LivGels, has a semiflexible backbone with carboxylate groups and vicinal diols in repeating units, 37,38 allowing for NaIO4-mediated selective oxidation and further functionalization with ADH (i.e., DH-ALG formation). The nLinker contains carboxylate and aldehyde groups, which are able to undergo Ca²⁺-mediated ionic crosslinking and Schiff base formation with the carboxylate and hydrazide groups of DH-ALG, respectively. The nLinker may enable not only dynamic covalent bonding (i.e., hydrazone bonds) with DH-ALG through a Schiff base reaction but also nanoparticle-polymer assembly via the Ca²⁺-mediated bridging of carboxylate groups in the DH-ALG matrix.

The chemical modification of ALG is schematically shown in Fig. S1a (ESI†). The vicinal diols are converted to aldehyde groups via NaIO₄-mediated oxidation, yielding DA-ALG (Fig. S1a-i, ESI†). Aldehyde groups are then converted to hydrazide via a Schiff base reaction, yielding DH-ALG (Fig. S1a-ii, ESI†). Hydrazides are commonly used for developing dynamic covalent bonds with aldehyde groups. 39,40 ALG chemical modification is assessed via ¹H NMR spectroscopy, and the spectra of ALG, DA-ALG, and DH-ALG are shown in Fig. S1b (ESI†). In the DA-ALG spectrum, two discrete peaks at the chemical shifts of ~ 5.35 ppm and 5.63 ppm were associated with the aldehyde groups in their hemiacetal forms. 41 Aldehyde groups of DA-ALG (5.0 \pm 0.2 mmol g⁻¹, Fig. S1c, ESI†), are reacted with hydrazide to form DH-ALG. In the DH-ALG spectrum, peak (a) ranging from 7 to 8 ppm corresponds to the -NH-, originated from the Schiff base reaction of DA-ALG dialdehyde groups with the hydrazide. In addition, peaks (b) and (c) at 1.5 ppm and 2.25 ppm, respectively, are attributed to the -CH₂- groups.⁴² Fig. S1d (ESI†) presents the content of remaining DH-ALG aldehyde groups after the reaction with hydrazide, which was only 0.4 ± 0.2 mmol g⁻¹. Additionally, the carboxylate group content of DH-ALG is $3.2 \pm 0.3 \text{ mmol g}^{-1}$ (Fig. S1e, ESI†), which agrees with the previously reported values (e.g., 3 mmol g^{-1}) for alginic acid.43

The schematic of DH-ALG/nLinker networks within the LivGel matrix is presented in Fig. 1c, indicating nLinker/nLinker and DH-ALG/nLinker crosslinks. Fig. 1d shows proposed mechanisms governing the strain-stiffening behavior of these crosslinked networks. In silico studies on random networks comprising crosslinked biopolymer chains have shown that a large-strain response is followed by the development of a stress path for axially stressed fibers and crosslinks.44 Strainstiffening in the LivGels may result from two mechanisms: (i) pulling out of stress-path undulations on DH-ALG/DH-ALG and DH-ALG/nLinker conjunctions and (ii) the reorientation of stress path on DH-ALG. The pulling out of stress-path undulations is bending-dominated in which case shear modulus (G) is correlated with stress (σ) to the power of 3/2, mainly resulting from the entropic behavior of biopolymers. When the biopolymer chains are crosslinked within a network, the interplay between the entropic behavior of chains and the constraints imposed by the crosslinks (i.e., network fluctuations) may lead to nonlinear mechanical behaviors. As σ increases, the polymer network becomes stiffer because of the entropic effects of biopolymer chains. The reorientation of the stress path is stretch dominated in which case G is correlated with σ to the power of 1/2, originating from single filament stretching.⁴⁴

3.2. nLinker synthesis and characterization

The nLinker, an integral component of LivGels responsible for imparting strain-stiffening and self-healing properties, was synthesized and characterized in terms of functional groups (specifically aldehyde and carboxylate groups), particle size, and morphology. Fig. 2a shows nLinker synthesis steps, involving the oxidation of cellulose fibrils and DAMC. Via the controlled partial oxidation of cellulose fibrils using NaIO4, DAMC was yielded and used as an intermediate precursor for nLinker synthesis. Fig. S2a (ESI†) shows a representative pH titration curve for DAMC fibrils. The NaOH amount required to neutralize the HCl released from the NH2OH·HCl-aldehyde reaction, is used to measure the aldehyde content (6.8 \pm 0.3 mmol g^{-1}).

After subjecting DAMC to a controlled chlorite oxidation reaction, whereby almost half of the aldehyde groups were transformed to carboxylate groups, the bifunctional nLinker was isolated following successive thermal treatment and ethanol-mediated precipitation. Fig. 2b and c show the representative titration curves of nLinker, bearing 3.0 \pm 0.2 mmol g^{-1} aldehyde and 3.5 \pm 0.5 mmol g^{-1} carboxylate groups, respectively. The coexistence of carboxylate and aldehyde groups enables concurrent physical (ionic) and chemical (dynamic covalent) bond formation with DH-ALG.

Fig. 2d shows the ATR-FTIR spectra of cellulose, DAMC fibrils, and the nLinker. In the DAMC spectrum, an increased

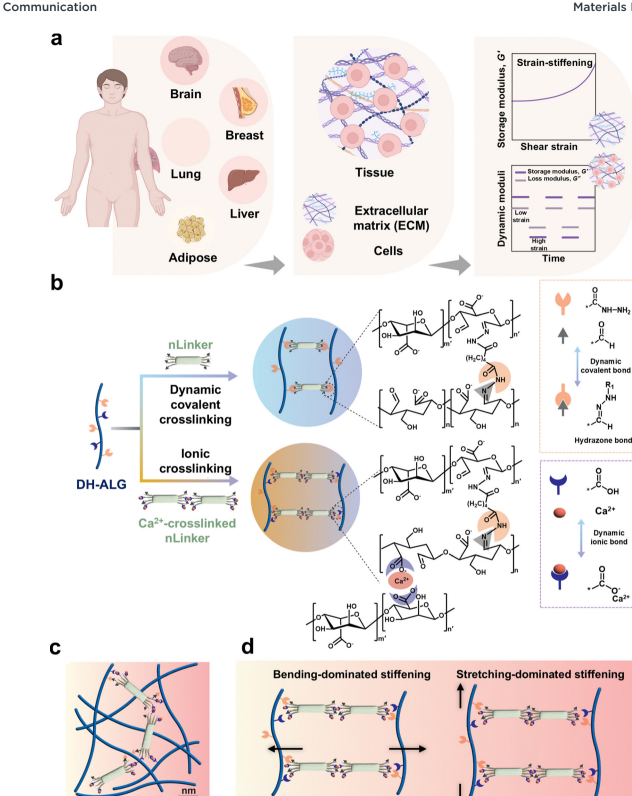


Fig. 1 Design of dynamically responsive, self-healing LivGels. (a) Examples of human soft tissues, consisting of shear-stiffening ECM and cell ensembles conferring self-healing. (b) Materials, engineering pathways, and physicochemical interactions, designed to enable dynamically responsive LivGels. DH-ALG, a semi-flexible bifunctional biopolymer undergoes ionic crosslinking and dynamic covalent hydrazone bond formation with bifunctional nLinkers. (c) Proposed mechanisms governing the large deformation-induced stiffening of crosslinked LivGels, involving (i) pulling out of stress-path undulations on polymer-polymer and polymer-nLinker networks, which is bending dominated, and (ii) stress path reorientation of DH-ALG chains, which is dominated by stretching. Note that Ca²⁺-mediated DH-ALG/DH-ALG crosslinks are not shown in panels (b)-(d).

 $G \propto \sigma^{3/2}$

Network fluctuation

Strain-stiffening self-healing matrix $G \propto \sigma^{1/2}$

Single filament stretching

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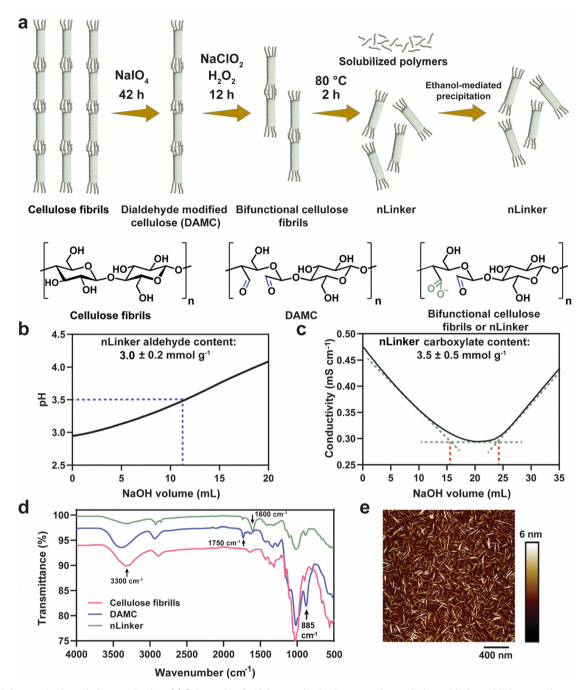


Fig. 2 nLinker synthesis and characterization. (a) Schematic of nLinker synthesis via successive periodate-chlorite oxidation reactions, as well as the corresponding changes in cellulose chemical structure. (b) A representative aldehyde titration curve of nLinker. (c) A representative carboxylate titration curve of nLinker using a strong base (10 mM NaOH). (d) ATR-FTIR spectra of cellulose fibrils, DAMC, and nLinker. (e) An AFM image of the nLinker.

intensity of the peak at $\sim 885 \text{ cm}^{-1}$ compared with cellulose was correlated to the hemiacetal linkages, arising from the interactions of aldehyde groups with the neighboring hydroxyl groups.31 The small peak at ~1750 cm-1 and broad peak at \sim 3300 cm⁻¹ were assigned to the stretching vibrations of carbonyl and hydroxyl groups, respectively. In the nLinker spectrum, the aldehyde and carboxylate groups were confirmed *via* the stretching vibrations of carbonyl groups at $\sim 1750 \text{ cm}^{-1}$ and $\sim 1600 \text{ cm}^{-1}$, respectively.

AFM images (see Fig. 2e as an example) showed that nLinker is a rod-shaped nanoparticle with a length of \sim 103 \pm 24 nm and a width of 5 \pm 2 nm. The hydrodynamic equivalent size of nLinker was 132 \pm 11 nm at pH \sim 6 and ionic strength \sim 3.5 mM, which was consistent with the previously reported values for bifunctional HCNC (~108 nm) and trimmed anionic HCNC (~ 140 nm, carboxylate content ~ 3.4 mmol g⁻¹). 45,46 The ζ -potential of nLinker at the same condition was – 31 \pm 6 mV, confirming the negatively charged carboxylate groups.

3.3. Mechanical characterization of LivGels

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An overview of the strain-stiffening and self-healing properties of common gels in the literature is provided in Fig. 3a. Conventional gels comprising crosslinked, flexible polymers are typically inert to shear strain.⁴⁷ However, gels made of semiflexible polymers, regardless of their biological or synthetic nature, may be shear-stiffening, 7,8 albeit without self-healing. Similar to semiflexible polymer gels, ECM-mimetic analogs have been designed not only to undergo stiffening under strain, 1,48 but also to enable other tissue-mimetic behaviors, such as anisotropic mechanics and shear-thinning. We aim to develop all-biopolymer-based LivGels that have both strainstiffening and self-healing properties and enable the adjustment of stiffness and stiffening response.

Before conducting mechanical characterization, lowamplitude oscillatory rheology at a constant frequency of 1 rad s⁻¹ and strain of 0.1% was performed on gels \sim 5 min after preparation, to examine their rheological behavior over time, as shown in Fig. S3 (ESI \dagger). G' did not significantly change within 15 min after NCH formation at most of the experimented nLinker or Ca2+ concentrations, indicating no significant evolution in network formation. To investigate the effect of nLinker-mediated dynamic hydrazone bonds on the linear and nonlinear mechanical properties of NCH, oscillatory strain amplitude sweep tests were conducted. Fig. 3b shows the G'of NCH versus shear strain at varying nLinker concentrations. At an nLinker concentration of 1.75 wt%, 2 wt%, or 2.25 wt%, the NCH underwent stiffening in the strain range of \sim 56 \pm 28% to 277 \pm 54%, \sim 66 \pm 10% to 188 \pm 88%, or \sim 54 \pm 1% to $95 \pm 6\%$ (n = 3), respectively, yielding LivGels. To determine the stiffening range of LivGels, we considered the stiffening onset as the strain point at which a 2% increase in G' occurred. The G' of LivGels, containing 1.75 wt%, 2 wt%, or 2.25 wt% nLinker, in the linear viscoelastic region (LVR) at a constant strain of 1%, was 30 \pm 15 Pa, 165 \pm 19 Pa, or 447 \pm 119 Pa (n = 3), respectively, falling in the stiffness range of reconstituted ECM $(G' \sim 30-1400 \text{ Pa})$. Type-1 collagen $(G' \sim 30-1000 \text{ Pa})$, fibrin ($G' \sim 300 \text{ Pa}$), and reconstituted basement membrane matrix $(G' \sim 60-1400 \text{ Pa})$ are examples of reconstituted ECM.⁵

In the NCH, the nLinker form dynamic hydrazone bonds via a Schiff base reaction between the nLinker aldehyde and ALG-DH hydrazide groups. By increasing the nLinker concentration from 1.75 wt% to 2.25 wt% at a constant ALG-DH concentration (2 wt%), the G' increased in the LVE region and NCH became stiffer as a result of the increased number of hydrazone bonds and nLinker-mediated network reinforcement. The mechanical properties of NCH strongly depend on the connectivity between polymer networks and particles. 20 The NCH G' within the LVR was used to estimate the average pore size $(\xi) \approx (G'/k_{\rm B}T)^{-1/3},^{49,50}$ where $k_{\rm B}$ is the Boltzmann constant, and T is the absolute temperature. Fig. S4a (ESI†) shows that the NCH pore size monotonically decreased from 52 \pm 6 nm to 21 \pm 1 nm when the nLinker concentration increased from 1.75 wt% to 2.25 wt%, implying that the crosslinking density increased. Importantly, when the nLinker concentration was ≤ 1.25 wt%, no NCH was formed. The molar ratio of nLinker aldehyde groups to DH-ALG

(2 wt%) hydrazide in the NCH containing 1.25 wt% nLinker was 1:0.4, which might not provide a sufficient number of dynamic hydrazone bonds for gel formation.

The effects of Ca²⁺-mediated ionic crosslinking along with the dynamic covalent bonds on the linear and nonlinear mechanics of NCH are presented in Fig. 3c-f. Note that 9 mM, 18 mM, or 36 mM of Ca²⁺ correspond to the saturation of $\sim 25\%$, $\sim 50\%$, or $\sim 100\%$ of nLinker carboxylate groups, respectively. Fig. 3c shows the G' and stiffening response of NCH, containing 1.25 wt% nLinker and varying Ca2+ concentrations. The nLinker (1.25 wt%) loaded with 9 mM or 18 mM of Ca²⁺ formed LivGels with a nonlinear stiffening behavior in the strain range of \sim 59 \pm 9% to 242 \pm 33% or \sim 91 \pm 1% to 188 \pm 28% (n = 3), respectively. However, NCH, containing the nLinker loaded with 36 mM of Ca²⁺ underwent stiffening over the whole strain range until the network was disrupted at a high strain. Increasing the Ca²⁺ concentration may shrink nLinker polymer chains via the neutralization of negatively charged carboxylates (stoichiometrically, at 36 mM, nLinker COO⁻: Ca²⁺ = 2:1 mol mol⁻¹), resulting in an increase in crosslinking density and G'. In another study, the addition of salt to the ALG-chitosan complexes increased G', which was attributed to the loss of water as the charges were screened by the counterions.51 Datapoints at which the gels undergo a significant drop in G' have not been presented in our figures. By increasing the Ca²⁺ concentration from 0 to 36 mM, the NCH stiffness increased because of the additional ionic bond formation within the NCH network. The G' of NCH, containing nLinker (1.25 wt%) with 9 mM or 18 mM of Ca^{2+} , was 44 ± 6 Pa or 94 \pm 20 Pa (within LVR, at strain \sim 1%), respectively, mimicking the stiffness of type-1 collagen or reconstituted basement membrane matrix.5

Fig. 3d presents the G' and stiffening of NCH, containing 1.75 wt% nLinker and varying Ca²⁺ concentrations. These gels, containing 9 or 18 mM of Ca2+, had a strain-stiffening behavior in the strain range of \sim 58 \pm 31% to 164 \pm 62% or \sim 31 \pm 8% to 71 \pm 26% (n = 3), respectively. The stiffness of LivGels, containing 1.75 wt% nLinker with 9 or 18 mM Ca^{2+} , was 155 \pm 43 Pa or 508 \pm 138 Pa (at strain \sim 1%, LVR), respectively, close to that of the reconstituted ECM.5 Besides increasing NCH stiffness by increasing the Ca²⁺ concentration (0 to 36 mM) at a constant nLinker concentration, increasing the Ca2+-loaded nLinker from 1.25 wt% to 1.75 wt% resulted in stiffer hydrogels, implying an increase in the crosslinking density.

The strain-stiffening mechanism of LivGels can be explained by stress localization at the hairy junction points where the majority of dynamic and ionic crosslinks exist. When the strain increases in NCH, stress is applied to the hairs. As a result, the nLinker can reorient locally along the strain direction, stretching the entire network and causing a stiffening response. Followed by nLinker reorientation, their extremely stiff crystalline body (Young's modulus ~ 120-135 GPa⁵²) may be connected to each other via the protruding hairs, forming semiflexible fibrils with finite bending stiffness. At an extremely large strain, some fraction of hairy junction points may break, initiating the yield (failure) process.

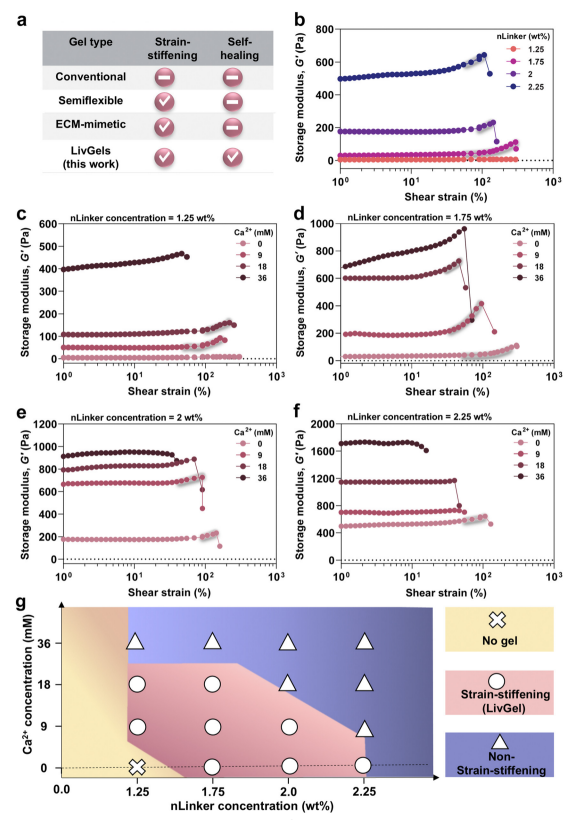


Fig. 3 Strain-stiffening properties of LivGels are regulated by the nLinker and Ca²⁺-concentrations. (a) An overview of strain-stiffening and self-healing properties of conventional gels, comprising flexible chains, networks of semiflexible polymers, or ECM-mimetic biopolymer networks, compared with the LivGels. The LivGels mimic the ECM mechanics and are self-healing. Storage modulus (G') versus oscillatory shear strain for the NCH, containing (b) varying Ca²⁺-free nLinker concentrations, or varying Ca2+ concentrations at a constant nLinker concentration ((c) 1.25 wt%, (d) 1.75 wt%, (e) 2 wt%, or (f) 2.25 wt%) and a constant frequency of 1 rad s^{-1} . The matrix biopolymer (DH-ALG) concentration was 2 wt% in all the NCH. (g) The phase diagram of NCH behavior at varying nLinker and Ca²⁺ concentrations and a constant DH-ALG concentration (2 wt%). The LivGels are the NCH that exhibit a strain-stiffening behavior.

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Fig. 3e and f show the G' and stiffening response of NCH, containing 2 wt% and 2.25 wt% nLinker at varying Ca2+ concentrations, respectively. The NCH without Ca2+ (at 2 or 2.25 wt% nLinker) or with 9 mM of Ca²⁺ (at 2 wt% nLinker) had a stiffening behavior, shown in shades; however, at higher Ca²⁺ concentrations (18 and 36 mM), the NCH did not have a nonlinear response. At a constant Ca²⁺ concentration (9 mM), NCH with 2.25 wt% nLinker showed no stiffening, while the analogs with 2 wt% nLinker underwent $\sim 5\%$ increase in G', indicating that ionic crosslinking via nLinker further tailors the stiffening behavior. Additionally, the stiffness of LivGels, containing 2 wt% nLinker without Ca^{2+} (165 \pm 19 Pa) or with 9 mM Ca^{2+} (592 ± 128 Pa), were in the range of reconstituted ECM. When the Ca²⁺ concentration was significantly high (36 mM), all the NCH containing 1.25 wt% to 2.25 wt% nLinker did not have any nonlinear response.

At any nLinker concentration, Ca2+ increased the NCH stiffness as a result of ionic crosslinking, reducing the nonlinear stiffening capability. At high Ca²⁺ concentrations (e.g., 18 mM for the NCH containing 2 or 2.25 wt% nLinker and 36 mM for all the NCH), the loss of strain-stiffening behavior in the gels may be a result of increased network connectivity beyond the isostatic threshold, rendering the otherwise semiflexible networks stiff. This may impair network fluctuations through changes in the nLinker-nLinker or DH-ALG-nLinker connections along the strain direction. Thus, the deformation mechanism is mainly filament stretching ($G \propto \sigma^{1/2}$). In contrast, if the average connectivity is below the isostatic threshold, the network is floppy and the deformation is through changes in the angle between the rods ($G \propto \sigma^{3/2}$), allowing the alignment of rods along the strain direction. 9,53,54

Fig. S4b (ESI†) shows the estimated NCH pore size at varying nLinker and Ca²⁺ concentrations. When the NCH, containing 1.25 wt% nLinker, were ionically crosslinked with 9 mM of Ca^{2+} , the pore size was 43.5 \pm 3 nm, which significantly decreased to 34 \pm 2 nm at 18 mM of Ca $^{2+}$ and to 22 \pm 1 nm at 36 mM of Ca²⁺. In the NCH, containing 1.75 wt% nLinker, when the Ca²⁺ concentration increased from 0 to 9 mM, the pore size significantly decreased from 52 \pm 6 nm to 27 \pm 1 nm, and the pore size changes were insignificant by increasing the Ca^{2+} concentration from 18 mM (19 \pm 1 nm) to 36 mM (18 \pm 1 nm). The evolution of pore size suggests that the maximum crosslinking via ionic bonding in the NCH matrix was obtained at \sim 18 mM of Ca²⁺. Stoichiometrically, 32 mM of Ca²⁺ is required for the saturation of all the carboxylate groups $(3.2 \pm 0.3 \text{ mmol g}^{-1})$ of DH-ALG, considering that I mol of Ca²⁺ binds to 2 mol of COO-. However, not all the nLinker loaded with Ca²⁺ may be available for ionic interactions with DH-ALG carboxylate groups, likely as a result of Ca²⁺-mediated internal crosslinking of carboxylates within the nLinker. In the NCH containing 2 wt% nLinker (Fig. S4b, ESI†), the pore size significantly decreased from 28 \pm 1 nm (Ca²⁺ concentration = 0) to 17 \pm 1 nm at 9 mM of Ca²⁺, followed by no significant changes as the Ca2+ concentration was increased up to 36 mM. Similarly, increasing the Ca²⁺ concentration no longer affected the pore sizes of NCH, containing 2.25 wt% of nLinker. Fig. 3g

shows the phase diagram of NCH, including strain-stiffening (LivGels) and non-strain-stiffening gels yielded at varying nLinker and Ca2+ concentrations. At a low (e.g., 1.25 wt%) nLinker concentration, no NCH was formed without Ca2+-mediated ionic crosslinking as a result of insufficient hydrazone bonds. As the Ca²⁺-free nLinker concentration increased up to 2.25 wt%, LivGels were formed. At any constant nLinker concentration ≤2 wt%, increasing Ca²⁺ concentration initially yielded LivGels, followed by non-strain-stiffening gels. At 2.25 wt% of nLinker, Ca²⁺ rendered the NCH non-strain-stiffening. Accordingly, the NCH are converted to LivGels at a specific concentration range of nLinker and Ca²⁺.

3.4. Quantitative analysis of strain-stiffening behavior in LivGels

To quantify stiffening, differential shear modulus, $K' = \partial \sigma / \partial \gamma$, where σ and γ are shear stress and strain, respectively, was calculated for the NCH, containing varying nLinker and Ca²⁺ concentrations. Fig. 4a shows K' versus shear stress at a constant frequency for NCH, containing varying Ca2+-free nLinker concentrations. A linear region at low stress, followed by a nonlinear region at higher stress were observed at any nLinker concentration, except when nLinker concentration = 1.25 wt% at which no NCH was formed. In the lower stress region, K' was equal to G' in the LVR (G_0) , whereas in the high stress region, K' had an exponential correlation with shear stress $(K' \propto \sigma^m)$. The stiffening index (m) and critical stress (σ_c) are indicators of stiffening intensity and the onset of nonlinear region, respectively. When m is close to 0.5, the dominant stiffening mechanism is filament stretching; however as m approaches 1.5, the network fluctuation mainly regulates the stiffening responses.44 By conducting oscillatory amplitude sweep tests and extracting the exponents, we investigated how the LivGel components, specifically nLinker or Ca2+-loaded nLinker, and the corresponding networks affect the transition between the stiffening mechanisms (i.e., network fluctuation versus single filament stretching) and control the dominant power-law scaling behavior.

Based on the K' plots (Fig. 4a), the m and σ_c values for LivGels containing varying concentrations of Ca²⁺-free nLinkers were obtained, which are presented in Fig. 4b and c, respectively. The m values for LivGels containing 1.75, 2, and 2.25 wt% of nLinker were 0.85 \pm 0.10, 0.72 \pm 0.07, and 0.30 \pm 0.11, respectively, indicating a decrease in the stiffening extent by increasing the nLinker concentration. The m values imply that the dominant stiffening mechanism is a combination of filament stretching and network fluctuations at low nLinker concentrations, which shifts to more filament stretching by increasing the nLinker concentration (≥ 2 wt%). The σ_c values of LivGels, containing 1.75, 2, or 2.25 wt% nLinker were 7 \pm 3, 100 \pm 15, or 95 \pm 11 Pa, respectively, showing a significant increase in σ_c , *i.e.*, less nonlinear responsiveness, by increasing the nLinker concentration. The decrease in m and increase in $\sigma_{\rm c}$ may be associated with an increase in the stiffness (G_0) by increasing the crosslinking density via the formation of hydrazone bonds and/or the reinforcing contribution of nLinker,

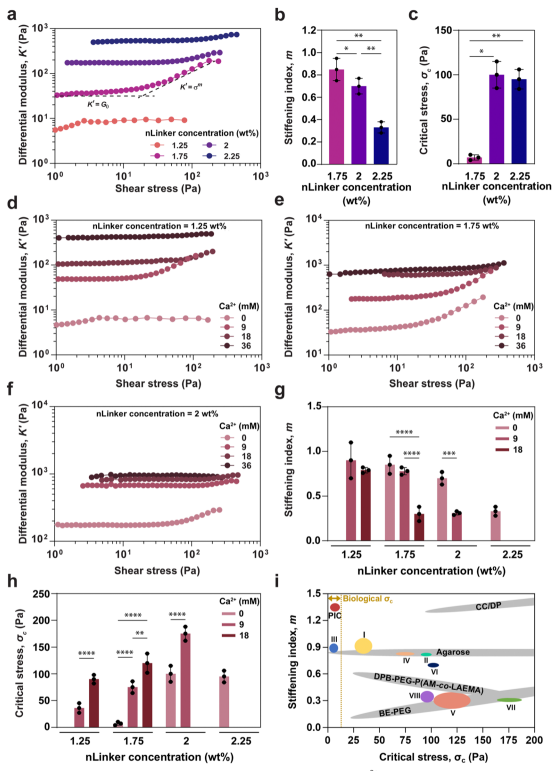


Fig. 4 Quantitative analysis of strain-stiffening behavior in LivGels, containing varying nLinker and Ca²⁺ concentrations. (a) Differential modulus (K') versus shear stress at a constant frequency of 1 rad s⁻¹, (b) stiffening index (m), and (c) critical stress (σ_c) of LivGels, containing varying Ca²⁺-free nLinker concentrations. K' versus oscillatory shear stress at a constant frequency of 1 rad s⁻¹, measured for the LivGels, containing varying Ca^{2+} concentrations and a constant nLinker concentration of (d) 1.25 wt%, (e) 1.75 wt%, or (f) 2 wt%. (g) m and (h) σ_c of the LivGels at varying Ca²⁺ (0–18 mM)-loaded nLinker concentrations (1.25–2.25 wt%). The LivGels containing 36 mM of Ca^{2+} did not have strain-stiffening behavior (n=3, *p<0.05, **p<0.01, ****p<0.001, ****p<0.001. (i) $m-\sigma_c$ profiles of biological and synthetic gels, compared with the LivGels, containing 1.25 wt% of nLinker with (I) 9 mM of Ca²⁺ or (II) 18 mM of Ca²⁺, LiveGels containing 1.75 wt% of nLinker with (III) 0 mM of Ca²⁺, (IV) 9 mM of Ca²⁺, or (V) 18 mM of Ca²⁺, LiveGels containing 2 wt% nLinker with (VI) 0 mM of Ca²⁺ or (VII) 9 mM of Ca²⁺, or (VIII) LiveGels containing 2.25 wt% of nLinker with 0 mM of Ca²⁺. PIC, DPB-PEG-P(AM-co-LAEMA), CC/DP, and BE-PEG stand for synthetic polyisocyanides, diphenylboronic acid-terminated telechelic poly(ethylene glycol)-poly(acrylamide-co-2-lactobionamidoethyl methacrylamide), O-carboxymethyl chitosandibenzaldehyde-terminated telechelic-ploy(ethylene glycol), and boronate ester-crosslinked 4-arm poly(ethylene glycol), respectively. Biological σ_c range $\sim 0.1-10 \text{ Pa.}^{1,55}$

rendering the LivGels less dynamically responsive to the applied strain. These results show that by changing the nLinker concentration, the nonlinear mechanical response of LivGels is readily tailored at a constant matrix biopolymer (DH-ALG) concentration (2 wt%).

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To quantify the effect of Ca²⁺ on the strain-stiffening behavior of NCH, the K' of LivGels, containing 2 wt% of DH-ALG and varying Ca2+ concentrations, was measured as a function of oscillatory shear stress at a constant nLinker concentration of 1.25, 1.75, 2, or 2.25 wt%, as presented in Fig. 4d-f, or Fig. S5 (ESI†), respectively. For the LivGels containing 1.25 wt% or 1.75 wt% of nLinker loaded with 9 or 18 mM Ca²⁺, or 1.75 wt% or 2 wt% of Ca²⁺-free nLinker, as well as the LivGel containing 2 wt% of nLinker loaded with 9 mM Ca²⁺, a linear region at low stress and a nonlinear region at higher stress were observed, which were consistent with the oscillatory strain sweep results (Fig. 3). For all the LivGels, G_0 at low stress increased by increasing the Ca²⁺ concentration, confirming an increase in the ionic crosslinking density. The m and σ_c values of LivGels containing varying Ca²⁺ concentrations are shown in Fig. 4g and h, respectively. The m values for LivGels, containing 1.25 wt% nLinker loaded with 9 or 18 mM Ca $^{2+}$ were \sim 0.87 \pm $0.18 \text{ or } \sim 0.73 \pm 0.03$, respectively. For these gels, by increasing the Ca^{2+} concentration, the σ_c value shifted significantly from 28 \pm 11 Pa at 9 mM of Ca²⁺ to 90 \pm 8 Pa at 18 mM of Ca²⁺. Similarly, in the LivGels containing 1.75 wt% or 2 wt% nLinker, by increasing the Ca^{2+} concentrations, m decreased and σ_c increased. This reduction in the mechanical responsiveness of LivGels may be attributed to an increase in stiffness as a result of dual dynamic hydrazone/ionic crosslinking and the increased network connectivity beyond the isostatic threshold, rendering the semi-flexible networks stiff. Accordingly, when the LivGels, containing 2.25 wt% of nLinker were loaded with Ca²⁺ (Fig. S5, ESI†), they did not have any strain-stiffening behavior. Additionally, the m values of LivGels, containing 1.75 wt% of nLinker, imply that the stiffening mechanism in the absence of Ca^{2+} (m = 0.85 \pm 0.10) may be a result of filament stretching and network fluctuations, whereas by increasing the Ca^{2+} concentrations (e.g., $m = 0.30 \pm 0.08$ at 18 mM), the stiffening may be regulated by filament stretching. A reason for this transition may be a high density of ionic crosslinking by which the local reorientation (bending) of nLinkers along the strain direction is restricted, and therefore the stiffening is governed via filament stretching.

Fig. 4i summarizes the stiffening parameters (m and σ_c) of biological and synthetic gels, as well as the LivGels. The LivGels, containing Ca²⁺-free nLinker (1.75 wt%) were engineered to resemble the strain-stiffening characteristics of biological gels (e.g., 0.5 Pa $< \sigma_{\rm c} < 10~{\rm Pa}^{1,55}$). In the stressstiffening region, m has been predicted to be 1.5 for a single semiflexible polymer, as observed in fibrin fibers⁵⁵ and actin filaments.⁵⁶ Among the developed hydrogels, synthetic polyisocyanide (PIC) gels have comparable strain-stiffening parameters with biological analogs. Synthetic boronate estercrosslinked 4-arm poly(ethylene glycol) (BE-PEG) hydrogels have strain-stiffening properties with σ_c ranging from 20 Pa

to 1000 Pa and m ranging from 0.1 to 0.5, regulated by the polymer concentration, temperature, and pH.57 Agarose hydrogels are a class of ECM-like biomaterials with strain-stiffening behavior because of their semi-flexible nature. The LivGels offer a broad range of m, spanning from ~ 0.87 (9 mM Ca²⁺-loaded nLinker concentration = 1.25 wt%) to ~ 0.25 (Ca²⁺-free nLinker concentration = 2.25 wt%), with σ_c ranging from ~28 Pa to ~ 157 Pa, adjusted by the crosslinking density or modes (i.e., hydrazone and ionic bonding). Diphenylboronic acidterminated telechelic poly(ethylene glycol)-poly(acrylamide-co-2-lactobionamidoethyl methacrylamide) (DPB-PEG-P(AMco-LAEMA))58 and O-carboxymethyl chitosan-dibenzaldehydeterminated telechelic-ploy(ethylene glycol) (CC/DP)⁵⁹ are other hydrogels with stiffening responses that are mechanically mimicked by the LivGels. Note that DPB-PEG-P(AM-co-LAEMA) and CC/DP gels do not recapitulate the stiffening properties, specifically σ_c , of biological gels. Overall, LivGels provide a novel bio-based platform that mimic the dynamic mechanical properties of ECM.

3.5. Qualitative and quantitative analyses of LivGel self-healing

The dynamic covalent hydrazone bonds and physical ionic interactions have been adopted extensively to develop selfhealing gels.⁶⁰ Accordingly, our LivGels were engineered using an aldehyde- and carboxylate-bearing nLinker to include reversible bonds that would enable self-healing behavior when mixed with DH-ALG. The self-healing capability of LivGels was assessed by strain amplitude measurements in which repeating cycles of low (1%) and high (500%) strains were used. Fig. 5a-d present the recovery tests in which the dynamic moduli of LivGels, containing varying nLinker concentrations loaded with 9 mM of Ca2+, are registered over time. The recovery tests of other NCH, containing varying nLinker and Ca²⁺ concentrations are shown in Fig. S6 (ESI[†]). These experiments were conducted to investigate how rapidly the LivGels recover their solid-like structure after the elimination of high strain. By applying strain of 500%, the LivGels had a fluid-like (viscous) behavior, reflected in G' < G''. Right after each cycle of high strain, the LivGel recovery was monitored at 1% strain and 1 rad s⁻¹ angular frequency. All strain-stiffening LivGels, containing either Ca2+-free nLinker (formed via hydrazone bonds) or Ca²⁺-loaded nLinker (formed via both ionic interactions and hydrazone bonds), had excellent recovery within seconds. At an extremely large strain, a fraction of hairy junctions/crosslink points between nLinker/nLinker or nLinker/DH-ALG may break. These broken junction points, made of reversible bonds, reform upon strain reduction, healing the structure of LivGels. Fig. 5e shows a qualitative self-healing assessment of LivGels, containing Ca²⁺-free nLinker (2 wt%) or 9 mM Ca2+-loaded nLinker (2 wt%). The LivGels were prepared in star-shape molds, cut into two pieces, and healed in 10 min after juxtaposing the cut interfaces. The healed LivGels, containing Ca2+-free nLinker or Ca2+-loaded nLinker, were mechanically robust and preserved their integrity upon hanging.

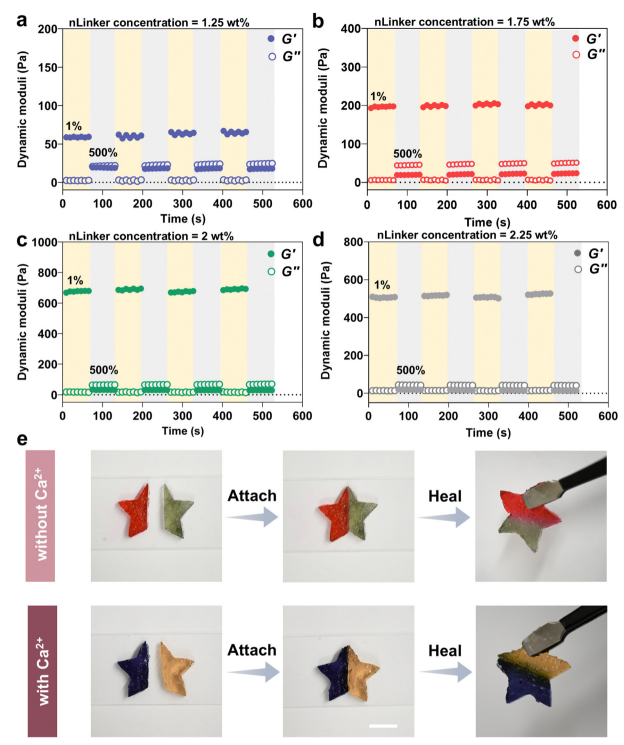


Fig. 5 Self-healing of LivGels. Oscillatory shear strain amplitude tests at alternating low (1%)-high (500%) strain cycles, conducted on the LivGels, containing varying nLinker concentrations of (a) 1.25 wt%, (b) 1.75 wt%, or (c) 2 wt%, all loaded with 9 mM Ca²⁺, or (d) 2.25 wt% without Ca²⁺ (f) Representative photos, showing the self-healing behavior of LivGels, crosslinked with the nLinker (2 wt%) covalently or with Ca²⁺(9 mM)-loaded nLinker ionically and covalently (scale bar = 1 cm).

4. Conclusions

Biopolymeric acellular nanocomposite living hydrogels, referred to as LivGels, were developed with nonlinear mechanical behavior and self-healing properties. The LivGel design was inspired by the shear-stiffening semi-flexible polymeric networks of natural ECM and cell-imparted healing after tissue damage. Via rationally tailoring the crosslinking of a network-forming biopolymer and anisotropic cellulose-based hairy nanoparticles (nLinker), conventional static gels were converted to dynamically responsive analogs using ionic and dynamic covalent hydrazone bonds. The anisotropic bifunctional nLinker, bearing aldehyde and carboxylate groups, enabled tailoring of the m, σ_c , and G' of LivGels within the range of 0.30 to 0.87, 7 to 175 Pa, and 30 to 591 Pa, respectively, at a constant matrix biopolymer (DH-ALG) concentration. The strain-stiffening LivGels underwent rapid recovery under cyclic low (1%)-high (500%) strains. This work may pave the way for developing bio-based acellular living materials with complex dynamic attributes for a broad range of applications, including in vitro ECM models, tissue engineering, additive manufacturing, and soft robotics.

Data availability

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Data pertaining to this work are presented in the main manuscript and the ESI.† Further relevant data are available from the corresponding author upon reasonable request.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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