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Hyperexpandable, self-healing macromolecular crystals with integrated polymer networks

Ling Zhang¹, Jake B. Bailey¹, Rohit H. Subramanian¹, Alexander Groisman², and F. Akif Tezcan^{1,3,*}

¹Department of Chemistry and Biochemistry, University of California, San Diego, La Jolla, CA, USA.

²Department of Physics, University of California, San Diego, La Jolla, CA, USA.

³Materials Science and Engineering, University of California, San Diego, La Jolla, CA, USA.

Abstract

The formation of condensed matter typically involves a tradeoff between structural order and flexibility. As the extent and directionality of interactions between atomic or molecular components increase, materials generally become more ordered but less compliant, and vice versa. Nevertheless, high levels of structural order and flexibility are not necessarily mutually exclusive; there are many biological (such as microtubules^{1,2}, flagella³, viruses^{4,5}) and synthetic assemblies (for example, dynamic molecular crystals $^{6-9}$ and frameworks $^{10-13}$) that can undergo considerable structural transformations without losing their crystalline order and that have remarkable mechanical properties^{8,14,15} that are useful in diverse applications, such as selective sorption¹⁶, separation¹⁷, sensing¹⁸ and mechanoactuation¹⁹. However, the extent of structural changes and the elasticity of such flexible crystals are constrained by the necessity to maintain a continuous network of bonding interactions between the constituents of the lattice. Consequently, even the most dynamic porous materials tend to be brittle and isolated as microcrystalline powders¹⁴, whereas flexible organic or inorganic molecular crystals cannot expand without fracturing. Owing to their rigidity, crystalline materials rarely display self-healing behaviour²⁰. Here we report that macromolecular ferritin crystals with integrated hydrogel polymers can isotropically expand to 180 per cent of their original dimensions and more than 500 per cent of their original volume while retaining periodic order and faceted Wulff morphologies. Even after the separation of

^{*} tezcan@ucsd.edu. Correspondence and requests for materials should be addressed to F.A.T. Author contributions

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L.Z., J.B.B. and F.A.T conceived the work and designed the experiments. L.Z. conducted the majority of the experiments and performed light microscopy analysis. L.Z. and J.B.B. performed XRD and SAXS analyses. L.Z. and R.H.S. performed confocal microscopy analysis. A.G. designed the microfluidic cells. F.A.T wrote the manuscript with contributions from all authors.

Online content

Any Methods, including any statements of data availability and Nature Research reporting summaries, along with any additional references and Source Data files, are available in the online version of the paper at https://doi.org/10.1038/s41586-018-0057-7. Competing interests

L.Z., J.B.B. and F.A.T. have submitted a patent application based on the work described here.

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neighbouring ferritin molecules by 50 ångströms upon lattice expansion, specific molecular contacts between them can be reformed upon lattice contraction, resulting in the recovery of atomic-level periodicity and the highest-resolution ferritin structure reported so far. Dynamic bonding interactions between the hydrogel network and the ferritin molecules endow the crystals with the ability to resist fragmentation and self-heal efficiently, whereas the chemical tailorability of the ferritin molecules enables the creation of chemically and mechanically differentiated domains within single crystals.

Hydrogel polymers present a stark contrast to molecular crystals in that they lack structural order, but are highly elastic and adaptive, can expand considerably and self-heal when equipped with dynamic bonding functionalities^{21,22}. Previously, the isotropic swelling -contraction behaviour of hydrogels has been used to modulate the lattice spacing of colloidal nanoparticle arrays²³, and recently, to expand biological tissue samples and thus facilitate high-resolution fluorescence imaging²⁴. In this study, we examine whether the mechanical properties of hydrogels could be endowed upon molecular crystals. That is, can crystal lattices that are formed by discrete molecules that are connected via specific bonding interactions be mechanically modulated through the integration of polymeric hydrogels? To create hydrogel-expandable molecular crystals, we surmised that the following design parameter conditions should be met: (1) lattices should be mesoporous to enable the hydrogel network to penetrate efficiently and uniformly into the crystals; (2) intermolecular interactions between the constituents of the lattices should be reversible and chemically specific (that is, contain directional and dynamic bonds), such that they disengage with ease during expansion and re-engage with high fidelity upon contraction; (3) interactions between the constituents of the lattice and the hydrogel network should be extensive to maintain the integrity of the crystal-polymer hybrid at all times and sufficiently dynamic to minimize the build-up of local strain and to enable self-healing.

With these parameters in mind, we arrived at hybrid materials composed of ferritin crystals integrated with the superabsorbent poly(acrylate-acrylamide), or p(Ac-Am), copolymer hydrogels, whose swelling-contraction behaviour can be modulated by the ionic strength and pH²⁵. Ferritin is a 24-meric, quasi-spherical protein with 432 symmetry, an outer diameter of 12 nm, an inner diameter of 8 nm, and a molecular weight²⁶ of more than 500,000 Da. Human heavy-chain ferritin forms highly ordered, face-centred cubic (fcc) crystals that routinely grow to more than 200 µm in size and diffract to less than 2.0 Å. The fcc lattice (Fig. 1a) is characterized by a mesoporous network consisting of cube-shaped, 6-nm-wide chambers (Fig. 1b) that are interconnected by smaller, octahedron-shaped cavities that taper to a pore size of about 2 nm at their narrowest (Fig. 1c), thus fulfilling condition (1). The lattice is formed through highly specific, metal-mediated contacts between neighbouring ferritin molecules (Fig. 1d), which are promoted through the K86Q surface mutation to enable metal coordination²⁷. The absence of any other interprotein contacts means that the entire lattice bonding framework of ferritin molecules can be formed or broken via binding or removal of metal ions (such as Ca^{2+}), satisfying condition (2). Finally, ferritin bears a small negative charge, with a zeta potential ranging from -5.5 mV at pH 6.0 to -7.3 mV at pH 7.5 (Extended Data Fig. 1a, b). The exterior surface of ferritin presents a diffuse distribution of both negatively and positively charged residues (Extended Data Fig. 1c),

which should enable uniform association with the p(Ac-Am) network through a combination of ionic and H-bonding interactions, thus fulfilling condition (3) (Fig. 1e).

We first examined the efficiency of molecular diffusion and polymerization within ferritin crystals. Diffusion into single ferritin crystals was assessed using the fluorescent tracer rhodamine B by confocal fluorescence microscopy experiments. These experiments showed that a typical crystal (edge length, l_{edge} = 50–250 µm) was completely infiltrated by rhodamine B (Extended Data Fig. 2a), which is considerably larger (479 g mol⁻¹) than the Ac and Am molecules (both 71 g mol⁻¹), within 15 min. In a typical preparation of crystalhydrogel hybrids, ferritin crystals were incubated with polymer precursors (8.625% (w/v) sodium acrylate, 2.5% acrylamide and 0.2% N,N'-methylenebis(acrylamide)) for at least 10 h to ensure their uniform distribution in the lattice interstices. This treatment caused no apparent damage to the crystals (see Supplementary Information for quantification of polymer precursor concentrations inside the crystals). Crystals were then transferred into a solution containing 1% (w/v) ammonium persulfate (APS) and 1% (v/v) tetramethylethylenediamine (TEMED) to initiate free-radical polymerization within the lattice, as well as 4 M sodium chloride (NaCl) to limit swelling during polymerization (Fig. 1e). To assess the kinetics of polymerization inside the crystals, we added 0.3% (w/v) 8hydroxypyrene-1,3,6-trisulfonic acid (pyranine) to the aforementioned co-monomer mixture. Pyranine has been reported to become covalently incorporated into the polymer backbone upon radical-mediated crosslinking and undergo a shift²⁸ in its emission maximum from 512 to 420 nm. Thus, the extent of in crystallo polymerization could be monitored through the decrease of green fluorescence intensity (emission wavelength $\lambda_{emission} = 500-550$ nm, excitation wavelength $\lambda_{excitation} = 488$ nm), indicating that hydrogel formation was complete in less than 2 min for a crystal with $l_{edge} = 70 \,\mu\text{m}$ (Extended Data Fig. 2b, c and Supplementary Video 1; see Extended Data Fig. 3 for polymer quantification via ¹⁹F nuclear magnetic resonance, NMR). Polymerization was promptly followed by intrusion of the aqueous NaCl solution into the crystal-hydrogel matrix, which was clearly visualized owing to the difference between the refractive indices of the salt solution ($n_D = 1.3676$) and the matrix ($n_D \approx 1.34$) (Extended Data Fig. 2b and Supplementary Videos 1, 2). The solvent permeation process typically finished within 10 min and was accompanied by a small but noticeable enlargement of the crystals (5% increase in edge length) (Extended Data Fig. 2c).

Full expansion of hydrogel-infused ferritin crystals was initiated by placing them in deionized water. As observed using light microscopy, the expansion of the crystals was highly isotropic and their sharply faceted, polyhedral morphologies were preserved even after they grew to 210% of their original dimensions (Fig. 2a and Supplementary Video 2; see Extended Data Fig. 4a and b for additional examples), often without the appearance of any defects. The expansion kinetics was biphasic, with time constants $\tau_{fast} < 100$ s and $\tau_s \iota_{ow} \gg 300$ s (Fig. 2a). Isotropic growth continued indefinitely, until the edges of the materials were not discernible, but we typically stopped the process after <10 min, when considerable expansion had already occurred. No substantial release of ferritin molecules from the lattices was evident during the first 50 min of expansion (Extended Data Fig. 4c). Addition of a concentrated monovalent salt solution (NaCl or KCl) led to rapid dehydration

and isotropic contraction of the expanded crystals to nearly their original size (Fig. 2a and Extended Data Fig. 5a). Recovery of the original crystal dimensions could be achieved by further addition of $CaCl_2$, owing to the ability of Ca^{2+} to both screen the negatively charged polymer backbone more effectively and to re-engage specific interactions between ferritin molecules. The same effect was observed with other divalent metal-ion salts (Extended Data Fig. 5b). The expansion-contraction cycle could be repeated at least eight times without apparent loss in amplitude and change in crystal morphology when a monovalent metal-salt solution was used to induce contraction (Extended Data Fig. 6). We observed that crystals contracted with CaCl₂ displayed considerably smaller expansion owing to the enhanced strength of the polymer network and protein-protein interactions. In control experiments, we examined other hydrogel formulations, including hydrogels that only contained polar but non-charged (pAm or poly-tris(hydroxymethyl)methyl(acrylamide)) or non-polar (poly-Nisopropylacrylamide) side-chains (Extended Data Fig. 7). All of these polymers led to either dissolution or disintegration of crystals after initiation of in crystallo polymerization, suggesting a lack of substantial interactions between the functional groups on these polymers and on the ferritin surface.

Interestingly, pAc hydrogels promoted isotropic expansion of the crystals in the absence of Am co-monomers (Extended Data Fig. 7), indicating that carboxylate side-chains are the primary mediators of interactions with ferritin molecules. By contrast, treatment of ferritin crystals with pre-formed pAc polymers, which cannot diffuse into the lattice, led to crystal dissolution upon transfer into water (Extended Data Fig. 7c). Together, these observations confirm that (i) there are extensive non-covalent interactions between ferritin molecules and the p(Ac-Am) hydrogel matrix that preserve the structural integrity of even highly expanded crystals, and (ii) the hydrogel matrix continuously and uniformly pervades the entire lattice, thus promoting cooperative transmission of any lattice deformations to enable isotropic expansion–contraction.

We investigated the expansion-related changes in the lattice arrangement of ferritin molecules using small-angle X-ray scattering (SAXS). Initial experiments entailed bulk measurements of a large number of ferritin crystals suspended in a capillary tube. Figure 2b shows the evolution of the 'powder' SAXS pattern of more than 100 single p(Ac-Am)infused crystals upon the initiation of polymerization though the addition of APS/TEMED in a solution that contains no salt; thus, polymerization is immediately followed by expansion (see Methods for experimental details). The spectrum of the unexpanded crystals is indicative of an fcc lattice with a unit-cell parameter of a = 182.40 Å. The isotropic growth of the unit cell is evident from the correlated shifts of the Bragg peaks to lower angles. The decay of the higher-angle peaks is considerably more rapid and is accompanied by the emergence of the ferritin form factor. This is consistent with the picture that as the crystal expands, the hydrogel matrix becomes less dense around the ferritin molecules, leading to their increased mobility. However, the (111) reflection is still evident after 20 min of expansion, which means that some long-range periodic order is still present when the unit cell has grown to a = 325 Å (Fig. 2c) and the volume of the material has increased to 570% of its original value.

To probe the reversibility of lattice expansion, we set up a micro-fluidic flow cell for singlecrystal SAXS experiments (Extended Data Fig. 8), which circumvent the inherent issues associated with bulk measurements in a small capillary tube (such as sample heterogeneity and inefficient solvent diffusion). The SAXS data in Fig. 2d indicate that a single-crystal lattice that has expanded by 27%—corresponding to a separation of 35 Å between neighbouring ferritin molecules—can return to its original dimensions upon NaCl/CaCl₂induced contraction. To examine whether this recovery also occurs at the level of atomic periodicity, we conducted high-angle, single-crystal X-ray diffraction (XRD) experiments at room temperature (Fig. 3a-c). These experiments showed that crystals that expanded by up to 40% could fully regain their native diffraction pattern upon contraction with divalent metal-ion salts (Extended Data Fig. 5b). With such expanded and Ca-contracted crystals, we consistently obtained datasets with resolutions <1.15 Å at a synchrotron source at 100 K (Extended Data Table 1). Interestingly, the resulting crystal structures revealed two different conformational states of the Ca^{2+} -bridged ferritin-ferritin interfaces (Fig. 3d): about 60% of these interfaces were found in the native configuration (as shown in Fig. 1d, but with a well resolved Ca-coordinated water molecule), whereas the remaining 40% presented an alternative coordination mode for Ca²⁺, probably stemming from lattice rearrangements during contraction. Notably, the 1.06-Å-resolution crystal structure (R-factors, R_{work} = 9.10%; $\mathbf{R}_{\text{free}} = 10.26\%$; estimated coordinate error (dispersion precision indicator), 0.011 Å) is the highest-resolution ferritin structure reported until now. Our findings suggest that hydrogel infusion and the expansion-contraction process do not diminish XRD data quality and may actually improve it.

Any local anisotropy developed during the expansion or contraction of the hydrogel matrix would be expected to cause dislocations in the embedded ferritin lattice. Indeed, exposure of hydrogel-infused crystals to rapid changes or temporary spatial gradients in NaCl or CaCl₂ concentrations frequently led to fracturing (see, for example, Fig. 3b inset). However, these materials showed a remarkable ability to self-heal, whereby the cracks were spontaneously and, in some cases, scarlessly sealed (Fig. 4a, b and Supplementary Video 3), owing to the reversible bonding interactions of the hydrogel network with the protein molecules (Fig. 4c). It is important to note that covalently crosslinked hydrogels like p(Ac-Am) do not typically self-heal unless they are modified with dynamic bonding functionalities^{22,29}. In the case of our materials, the role of such functional groups is fulfilled by the ferritin molecules, which act as interaction hubs for polymer chains. During expansion-contraction cycles, cracks tended to reoccur in the same loci in a given crystal (Supplementary Video 3). This observation suggests that the healed interfaces had not fully regained the original hydrogel crosslinking density of the bulk material, at least in the time scale (several minutes) of the experiments. Hydrogel integration substantially mitigated the brittleness of native ferritin crystals (Extended Data Fig. 9a). We observed no fragmentation, even in cases of substantial fracturing that propagated throughout the crystals, and fissures as wide as 20 µm could be closed to recover near-native crystal morphology (Fig. 4b). The ferritin crystal-hydrogel hybrids had a reduced modulus of about 1 GPa, which is similar to that of ferritin crystals (Extended Data Fig. 9b), but several orders of magnitude higher than those of hydrogels³⁰. The hybrids are also highly thermostable, maintaining their crystalline order at 80 °C (Extended Data Fig. 9c).

Owing to the inherent chemical tailorability of ferritin molecules, the crystal-hydrogel hybrids could be functionalized in different ways. They could be constructed from ferritin molecules with mineralized ferrihydrite in their interior cavity (Extended Data Fig. 9d and Supplementary Video 4), thus exploiting ferritin's native function as a ferroxidase, or with fluorescent tags covalently attached to their exterior (Fig. 4d, e and Supplementary Video 4). Additionally, spatially differentiated, core-shell crystals were created using a layer-by-layer growth method (Fig. 4d, e). When infused with p(Ac-Am), such nanoparticle- or fluorophore-functionalized lattices displayed the same isotropic expansion-contraction behaviour as non-functionalized ones. The layer-by-layer growth process was further modified whereby the core lattice domain (labelled with rhodamine groups) was first covalently fixed through the chemical crosslinking of ferritin molecules with glutaraldehyde, followed by the growth of an uncrosslinked, unlabelled shell layer and the incorporation of the p(Ac-Am) polymer into the composite lattice. Hydration of such 'fixed core/expandable shell' crystals led to complete fragmentation of the shell layer due to the strain generated at the mechanically mismatched core-shell interface, exposing the morphologically unaltered core layer (Fig. 4e). These examples highlight the facility with which chemical and mechanical patterning are achieved in protein crystal-hydrogel hybrids.

We have reported here a new form of materials that integrate macro-molecular protein crystals with synthetic polymer networks. These hybrids seamlessly combine the structural order and periodicity of crystals, the adaptiveness and tunable mechanical properties of polymeric networks and the chemical versatility of protein building blocks. Additionally, the ability to reversibly expand-contract crystal lattices and mobilize their protein components may provide a new means to improve XRD quality and explore otherwise inaccessible protein structural states using three-dimensional protein crystallography. Protein crystals are often highly porous, sometimes containing up to 90% solvent, and are usually assembled through weak, non-covalent packing interactions; therefore, our approach should be applicable to other protein lattices. Their potential for generalizability, coupled with the chemical tailorability of synthetic polymers and the genetic mutability of proteins, should make protein crystal-hydrogel hybrids a rich medium for materials science.

METHODS

Protein expression, purification and characterization.

The plasmid for the C* variant of human heavy-chain ferritin (HuHF), devoid of all native cysteine residues (C90E, C102A and C130A), was obtained via site-directed mutagenesis as previously described³¹. Expression and purification of C* was performed according to the previously published protocol³².

Determination of zeta potentials.

Purified ferritin was concentrated to about 200 μ M and exchanged into a buffer solution containing 50 mM 2,2-Bis(hydroxymethyl)-2,2[/],2^{//}-nitrilotriethanol (Bis-Tris) (pH 6.0), 50 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) (pH 7.0) or 50 mM HEPES (pH 7.5) using an Amicon Ultra centrifugal filter unit (10 kDa cutoff). The zeta potentials of ferritin in the three different buffers were determined using a Zetasizer Nano

ZS90 (Malvern Instruments). Experimental runs were performed to collect 12 datasets with a He-Ne laser at 633 nm.

Formation of crystal-hydrogel hybrids.

Polymer precursor solution. 25 mM (HEPES; pH 7.0), 30 mM CaCl₂, 917 mM (8.625% w/v) sodium acrylate, 352 mM (2.5% w/v) acrylamide and 13 mM (0.2% w/v) N,N[;]- methylenebis(acrylamide). *Polymerization solution.* 4 M NaCl, 1% (w/v) APS and 1% (v/v) TEMED. Octahedron-shaped ferritin crystals formed over 1–2 days in a buffered solution containing 25 mM HEPES (pH 7.0), 3–14.5 μ M protein (per 24meric ferritin cage) and 4.5–7.5 mM CaCl₂. Once the ferritin crystals matured, the crystallization solution was replaced with the polymer precursor solution. Crystals were soaked for more than 10 h to ensure full infusion of the monomers into the ferritin crystals, and were then individually transferred with a mounted CryoLoop (Hampton) to the polymerization solution for 5 min, initiating in crystallo polymerization. Alternatively, the crystallization solution was replaced with the polymerization for the bulk polymerization of many crystals at once.

Measurement of the rate of diffusion into ferritin crystals.

A large ferritin crystal was transferred with a mounted CryoLoop onto a glass slide, and 20 μ l of a solution containing 20 μ M (0.01 mg ml⁻¹) rhodamine B, 30 mM CaCl₂ and 25 mM HEPES (pH 7.0) was added to the crystal. The rhodamine diffusion process was monitored with a 10 × air objective installed on a spinning-disk confocal Axio Observer inverted microscope (Zeiss) equipped with a pair of Quantum 5125C cameras (Roper), using a filter to collect light at 575–650 nm (red channel). Differential interference contrast (DIC) and fluorescence (564 nm excitation) images were captured at 1-s intervals with a 10-ms exposure. Images were collected in Slidebook 6 (Intelligent Imaging Innovations) and analysed using Fiji (http://fiji.sc/Fiji).

Determination of in crystallo polymerization dynamics.

Ferritin crystals were incubated in a polymer precursor solution supplemented with 5.7 mM (0.3%) pyranine (Sigma-Aldrich). After 12 h, an individual crystal was transferred onto a glass slide and polymerization was initiated by adding 10 µl of the polymerization solution. Hydrogel polymerization throughout the crystal and the corresponding decrease of pyranine fluorescence were monitored with a 20 × air objective on the confocal microscope as described above, using a filter to collect light at 500–550 nm (green channel). DIC and fluorescence (488 nm excitation) images were captured at 1-s intervals with 100-ms (DIC) and 1-s (fluorescence) exposures.

Scanning electron microscopy of ferritin crystals.

Native ferritin crystal and crystal-hydrogel hybrid samples were deposited onto glowdischarged, Formvar/carbon-coated Cu grids (Ted Pella Inc.). Each grid was blotted with filter paper to remove excess liquid. Grids were mounted onto a STEM 12x v2 sample holder and imaged using a Sigma 500 scanning electron microscope (Zeiss) at an accelerating voltage of 1 kV using a 30-µm aperture.

Polymer quantification with ¹⁹F NMR.

Large-scale crystallization of ferritin was carried out in a 24-well culture plate (Costar). 100 µl of 25 µM ferritin in 15 mM Tris (pH 7.4) and 150 mM NaCl) was combined with 100 µl of a buffered solution containing 50 mM HEPES (pH 7.0) and 12 mM CaCl₂. Crystals formed overnight and matured over 72 h. The solution in each well was replaced with 100 µl of a polymer precursor soaking solution containing: 25 mM HEPES (pH 7.0), 30 mM CaCl₂, 179.9 mM 2-(trifluoromethyl)acrylic acid, 744.8 mM sodium acrylate, 350.7 mM acrylamide and 20.4 mM N,N -methylenebis(acrylamide). After soaking overnight, this solution was removed, and the crystals were washed with a buffered solution (25 mM HEPES, pH 7.0; 30 mM CaCl₂) to remove unincorporated monomers. Polymerization was initiated by replacing the washing solution with 100 µl of the polymerization solution. After 10 min, the crystals were transferred into an Eppendorf tube and centrifuged at 2,000g for 60 s. The supernatant was decanted, and the crystals were resuspended in 1 ml D_2O . Concentrated HCl was added until the pH of the solution was approximately 4.0 to facilitate crystal decomposition. 705 µl of this solution was transferred into an NMR tube and supplemented with 4.6 mM trifluoroacetic acid. The ¹⁹F-NMR spectrum was collected using a 300M Bruker AVA spectrometer with a ¹⁹F probe (Extended Data Fig. 3). The peak at -64.94 p.p.m. corresponds to free 2-(trifluoromethyl)acrylic acid, the cluster of peaks near -67.07 p.p.m. to 2-(trifluoromethyl)acrylic acid that has been incorporated into the polymer, and the peak at -75.51 p.p.m. to the trifluoroacetic acid standard. From the integration of these peaks it was deduced that (a) the total concentration of 2-(trifluoromethyl)acrylic acid in the solution was 8.2 mM and (b) 74.7% of the monomeric precursor was incorporated into

the polymer matrix inside the crystals. The protein concentration was determined to be 60.0 μ M using the Bradford assay³³, and the molar ratio of 2-(trifluoromethyl) acrylic acid to ferritin was calculated as 137:1. Given this ratio and the fact that each unit cell of the ferritin crystals contains four ferritin cages and has a volume of about 5832 nm³ (*a* = 18 nm), the concentration of 2-(trifluoromethyl)acrylic acid in the crystal lattice was calculated as 155.6 mM, which is very similar to its concentration (179.9 mM) in the soaking solution.

Monitoring crystal expansion-contraction using light microscopy.

Single crystals were transferred with a mounted CryoLoop onto a glass slide with a microscopic ruler (OMAX). All images and videos were obtained on an SZX7 (Olympus) microscope equipped with an Infinity 1 charge-coupled device (CCD; Lumenera). For crystals that had not been polymerized, 10 μ l of the polymerization solution was carefully added to minimize crystal movement. This solution was removed before water addition. For previously polymerized crystals, water (Milli-Q 30 μ l) was added and crystal expansion was observed over 5–20 min. To initiate crystal contraction, water was replaced with a solution containing either 4 M NaCl or 1 M CaCl₂. This expansion-contraction cycle could be repeated at least eight times for a crystal if NaCl was used to induce crystal contraction. Crystal size was determined by measuring the edge length of a facet relative to the microscopic ruler using the Fiji image processing package.

Monitoring crystal expansion using confocal microscopy.

Crystals containing polymer precursors were prepared as described above. One of these crystals was transferred onto a glass slide and imaged on a confocal microscope. After capturing an initial image, the crystal was polymerized in 10 μ l of the polymerization solution, and its expansion in 30 μ l water was monitored. DIC images were captured at different time intervals with a 100-ms exposure until the crystal was no longer visible.

Quantification of protein release during expansion.

Large-scale crystallization of ferritin was carried out as described above. Once crystals fully matured, the well solution was replaced with 100 μ l of the polymer precursor solution. After 12 h, the crystals were all combined into a single Eppendorf tube and 500 μ l of the polymerization solution was added. Crystals were expanded by replacing the polymerization solution with 1 ml water. During this experiment, aliquots (100 μ l) of the protein solution were removed and replaced with 100 μ l of water, and each aliquot was used to determine the protein concentration using the Bradford assay.

Multi-crystal expansion monitored using SAXS.

Crystals for multi-crystal small-angle X-ray scattering were prepared as described above and transferred into the polymer precursor solution. A large number (n > 100) of crystals were transferred to an Eppendorf tube. After the crystals had settled at the bottom, they were transferred, along with 50 µl of solution, into a 1.5-mm quartz capillary (Hampton). Crystals in capillaries were analysed at beamline 5-ID-D of the Advanced Photon Source (Argonne National Laboratory). Data were collected using collimated X-ray radiation (0.7293 Å, 17 keV) calibrated with both a glassy carbon standard and a silicon diffraction grating. After the sample was mounted on the instrument, a thin tube (with a diameter of 0.51 mm) was inserted into the capillary to facilitate the addition of 50 μ l of solution with a syringe injector during X-ray exposure. The injected solution contained a more concentrated polymerization solution without NaCl (2% APS and 2% TEMED) in water. After the first exposure, the solution was injected, and an image with a 1-s X-ray exposure was collected every 30 s. Peaks corresponding to the original lattice were visible throughout the process, indicating that some of the crystals in the bulk sample did not expand. This is probably due to limited solvent diffusion or incomplete polymerization within the capillary tubes used for the SAXS experiments. It is important to note that in this procedure, 'polymerized' crystals immediately began expanding upon the commencement of data collection. The reason for this experimental strategy (instead of polymerization in a high-ionic-strength solution, followed by the initiation of expansion through lowering the ionic strength) is that it was not possible to sufficiently dilute the high-ionic-strength polymerization solution in the thin capillary tubes used for SAXS (which cannot accommodate addition of large volumes of solution) to enable expansion.

Scattered radiation was detected using a CCD area detector and onedimensional scattering data were obtained through the azimuthal averaging of the two-dimensional data to produce plots of the scattering intensity as a function of the scattering vector length, $q = 4\pi \sin(\theta/\lambda)$, where θ is one-half of the scattering angle and λ is the wavelength of the X-rays used.

Analysis of the one-dimensional data was performed using the powder diffraction processing software JADE (MDI) or Origin (OriginLab).

Multi-crystal SAXS at elevated temperatures.

Large-scale crystallization of ferritin was performed as described above. The crystallization solution was removed, and ferritin crystals were resuspended in either the polymer precursor solution or a buffered solution containing 25 mM HEPES (pH 7.0) and 30 mM CaCl₂. After 72 h, the polymer precursor soaking solution was replaced with the polymerization solution. After 10 min, this was also replaced with a buffered solution containing 25 mM HEPES (pH 7.0), 4 M NaCl and 30 mM CaCl₂. Both samples, containing either native ferritin crystals or the crystal-hydrogel hybrids, were transferred into 1.5-mm quartz capillaries (Hampton). Data were collected at beamline 4–2 of SSRL using collimated X-ray radiation (1.1271 Å, 11 keV) calibrated with a silver behenate standard. The samples were heated using a custom-built thermal stage operating at 1 °C min⁻¹, and images with a 1-s X-ray exposure were collected every minute. Scattered radiation was detected using a Pilatus3 X 1M detector (Detectris) and processed as described above.

Single-crystal SAXS.

Crystals containing the polymer precursors were prepared and polymerized using the polymerization solution as described above, and were analysed at SSRL (beamline 4–2). Single crystals were harvested with a mounted CryoLoop and transferred into a 2 M NaCl solution in the 400-µm-diameter central well of a custom-made microfluidic chip (Extended Data Fig. 8a, b). The microfluidic chip was sealed with a coverslip, attached to a syringe injector and mounted on beamline 4–2 at SSRL for data collection. Data were collected using collimated X-ray radiation (1.127 Å, 11 keV) calibrated with a silver behenate standard. Water was injected into the microfluidic chip at 1 µl s⁻¹ to initiate expansion, and 0.5-s-exposure images were taken every about 2.5 s for 4 min. After the data acquisition for crystal expansion was complete, the process was repeated—in the order 4 M NaCl, water, 1 M CaCl₂, water—to monitor repeated contraction and expansion processes. Data were collected using a Pilatus3 X 1M detector (Detectris). The unit-cell parameters were determined by calculating the radial distance of individual reflections, after fitting the spot intensity to a two-dimensional Gaussian surface.

Single-crystal XRD at room temperature.

Crystals containing polymer precursors were prepared and imaged using light microscopy as described above. A single crystal was transferred onto a MicroMount precision tool (MiTeGen) with a 100-µm aperture and sealed with a MicroRT capillary (MiTeGen). Data were acquired on an APEX II CCD diffractometer (Bruker) using Cu K α radiation (1.5418 Å) at 295 K. Three images (60-s exposure) were collected at rotation angles $\varphi = 0^{\circ}$, 60° and 120°. The crystal was removed from the instrument and soaked in 10 µl of the polymerization solution for 2 min. The crystal was transferred onto a microscopic ruler and 30 µl of water was added. Crystal expansion was measured over 3 min. This crystal was returned to the MicroMount with the MicroRT capillary and an identical three-image dataset was collected. This process was repeated using 30 µl of a 1 M CaCl2 solution. After the

crystal had contracted (1 min), another three-image dataset was collected. Images were analysed with the Apex III software (Bruker).

Single-crystal XRD at 100 K.

Crystal-hydrogel hybrids were prepared and imaged using light microscopy as described above. Two crystals were harvested, 30 µl of water was added, and crystal expansion was monitored over 5 min for both crystals. After 5 min, the water was removed and 30 µl of either a solution containing 1 M CaCl₂ (crystal A) or 4 M NaCl (crystal B) was added. Crystal B was re-expanded in 30 µl water. After 5 min, the water was replaced with 30 µl of a 1 M CaCl₂ solution to contract crystal B. After contraction, both crystals were cryoprotected in perfluoropolyether (Hampton) and frozen in liquid N2. Single-crystal XRD data for the contracted ferritin crystals were collected at 100 K at beamline 9-2 of SSRL using 0.98-Å radiation. The data were integrated using iMosflm³⁴ and scaled with Aimless³⁵ (Extended Data Table 1). The structures for crystal A and crystal B were determined at resolutions of 1.06 Å and 1.13 Å, respectively. Molecular replacement was performed with Phaser³⁶ using the HuHF structure (PDB ID, 5CMQ) as a search model. Rigid-body, positional, anisotropic thermal and atom-occupancy refinements were carried out using Phenix³⁷. Coot³⁸ was used for iterative manual model building. The interstitial solvent content was calculated by subtracting the solvent volume of each crystal from the volume of the inner cavity of ferritin (calculated using VOIDOO)³⁹. All figures were produced with Pvmol⁴⁰.

Nanoindentation measurements of crystals.

The mechanical properties of the native ferritin crystals and the crystal-hydrogel hybrids were determined using a Hysitron TI 950 Triboindenter test instrument (Bruker). All crystals were dried before the indentation experiments. A Berkovich probe (TI-0039, 142.3°, 100 nm tip radius) was used to determine the hardness and reduced modulus of the native crystals and crystal-hydrogel hybrids. Experiments were conducted in displacement control mode using a displacement of 1,000 nm.

Preparation of iron-loaded ferritin.

Iron-loaded ferritin was prepared by adding 10.8 ml of 10 mM $(NH_4)_2Fe(SO_4)_2$ over 2 h to 144.8 ml of a vigorously stirring solution containing 1 μ M ferritin, 15 mM Tris (pH 7.4) and 150 mM NaCl. Subsequently, the solution was stirred for an additional hour before being concentrated to about 3 ml using a 10-kDa Amicon membrane. A 10DG column (Bio-Rad) was used to remove any unbound iron. The iron content was assessed using a 2–2 '- bipyridine-based colorimetric assay⁴¹ and the protein concentration was determined using the Bradford assay. Each ferritin cage contained about 800 Fe atoms.

Formation of core-shell ferritin crystals.

Expandable core/expandable shell crystals. Mature ferritin crystals were transferred to a buffered solution containing 25 mM HEPES (pH 7.0), 30 mM CaCl₂ and 1.9 mM (1 mg ml ⁻¹) 5-(and 6)-carboxytetramethyl rhodamine succinimidyl ester (NHS-rhodamine; Thermo Fisher Scientific). After soaking for 12 h, an individual crystal was removed and washed

three times in a buffered solution containing 25 mM HEPES (pH 7.0) and 30 mM CaCl₂ to remove unbound NHS-rhodamine. The crystal was transferred to a well containing 10 μ l of 12.5 μ M ferritin, 25 mM HEPES (pH 7.0) and 6 mM CaCl₂. A transparent layer of ferritin formed around the rhodamine-labelled ferritin crystal over 12 h (creating a red core and a transparent shell). This crystal was soaked in a polymer precursor solution and polymerized as described above to yield an expandable core/expandable shell crystal.

Fixed core/expandable shell crystals. Fixed core/expandable shell ferritin crystals were prepared similarly to the expandable core/expandable shell crystals described above. The only difference was that after the rhodamine labelling step, the crystal was transferred into a solution containing 2.5% (v/v) glutaraldehyde, 25 mM HEPES (pH 7.0) and 30 mM CaCl₂. After 12 h, the crystal was washed five times with water to remove unbound glutaraldehyde, followed by the epitaxial growth of the transparent layer of ferritin crystals on top of the core layer in a fresh crystallization solution containing 12.5 μ M ferritin. This crystal was then soaked in a polymer precursor solution and polymerized as described above to yield a fixed core/expandable shell crystal.

Data availability.

Crystal structures have been deposited in the Research Collaboratory for Structural Bioinformatics Protein Data Bank under accession codes 6B8F (ferritin-polymer hybrid crystal 1; https://www.rcsb.org/struc-ture/6b8f) and 6B8G (ferritin-polymer hybrid crystal 2; https://www.rcsb.org/structure/6b8g). Raw data are available from F.A.T.

Extended Data



Extended Data Fig. 1 \mid . Distribution of electrostatic charge on the surface of ferritin and size distribution of ferritin in solution.

a, pH-dependent zeta potentials of ferritin, determined by dynamic-light-scattering measurements. **b**, Dynamic-light-scattering profile of ferritin (200 μ M) in a solution of 50 mM HEPES (pH 7.0). **c**, Representation of the electrostatic charge distribution on the ferritin surface, as viewed along the two-, three- and four-fold symmetry axes. Positive (+5k_BT/e) and negative (-5k_BT/e) charges are shown in blue and red, respectively. k_B, Boltzmann constant; *e*, electron charge.

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Extended Data Fig. 2 |. Molecular diffusion and polymerization in ferritin crystals, monitored using confocal microscopy.

a, Diffusion of rhodamine B into a ferritin crystal over 15 min. **b**, **c**, In crystallo polymerization of the hydrogel network, monitored through the decrease of integrated pyranine fluorescence (green fluorescence channel). The corresponding bright-field (DIC) images show the diffusion of the aqueous NaCl solution into the crystal. The ring-shaped diffusion front becomes evident at time t = 108 s and disappears by t = 216 s. The crystal expands by approximately 5% (edge length) during polymerization. Scale bars in a and b correspond to 100 µm. **d**, Scanning electron microscopy images of native ferritin crystals (top) and crystal-hydrogel hybrids (bottom).



Extended Data Fig. 3 |. Quantification of an acrylic acid analogue using 19FNMR.

a, ¹⁹F-NMR spectrum, showing peak assignments for the trifluoroacetic acid standard, free 2-(trifluoromethyl)acrylic acid, and 2-(trifluoromethyl)acrylic acid incorporated into the polymer. **b**, Diagram illustrating the experimental protocol for the quantification of 2-(trifluoromethyl)acrylic acid uptake into the crystal lattice. The concentration of the 2-(trifluoromethyl)acrylic acid in the crystal lattice (155.6 mM) is approximately the same as its concentration in the soaking solution (see Methods for details).



Extended Data Fig. 4 |. Isotropic hyperexpansion of ferritin crystal-hydrogel hybrids.

a, **b**, Continuous expansion of two different crystal-hydrogel hybrids in deionized water, monitored using confocal microscopy. Crystal facets are still discernible after expansion for more than 2 h. Scale bars correspond to 100 μ m. **c**, Ferritin release into the solution from expanding crystal-hydrogel hybrids (*n* > 10,000) over about 4 h. Negligible ferritin release is observed until about 1 h. Protein concentrations were determined using the Bradford assay. **d**, Confocal microscopy images of highly expanded crystal-hydrogel hybrids, showing the structural deterioration of the facets and the edges.



 $\mbox{Extended Data Fig. 5 |. Expansion and contraction behaviour of crystal-hydrogel hybrids in the presence of different metal ions. } \label{eq:extended_extended_extended}$

a, Light micrographs of the crystal-hydrogel hybrids at different stages of expansion and contraction in response to different metal ions. **b**, XRD patterns (T = 273 K) of expanded crystal-hydrogel hybrids, acquired upon contraction with different metal ions. Contraction with divalent cations (Ca, Mg, Cd, Zn, Ni and Co) reproducibly leads to the recovery of the full atomic-level order, whereas contraction with monovalent cations (Li, Na and K) only reinstates low-order diffraction peaks.



Extended Data Fig. 6 |. Successive expansion-contraction cycles for a single ferritin crystalhydrogel hybrid.

Light micrographs of a hybrid crystal at pre- and post-expansion stages in each cycle are shown on the left, and the corresponding changes in edge length upon expansion-contraction are shown on the right. The separation between the major ticks of the ruler is 100 μ m. The crystal expands to a lesser extent during the first expansion cycle, which we ascribe to residual CaCl₂ (which forms strong polymer-polymer and protein-protein interactions) remaining in the solution that is transferred on the loop along with the crystal. The subsequent variability in the rate and extent of expansion is attributed to the different amounts of residual NaCl transferred in each cycle.

а			post-	
	polymer precursors	chemical structure	polymerization	expanded
	-sodium acrylate -MBAm -acrylamide	$ \underbrace{\overset{O}{\longleftarrow}}_{NH_2} \underbrace{\overset{O}{\longleftarrow}}_{H} \underbrace{\overset{O}{\longleftarrow}}_{H} \underbrace{\overset{O}{\longleftarrow}}_{H} \underbrace{\overset{O}{\longleftarrow}}_{H} \underbrace{\overset{O}{\longleftarrow}}_{O^-Na^+} \underbrace{\overset{O}{\longleftarrow}}_{Na^+} \underbrace{\overset{O}{\longleftarrow}_{Na^+} \underbrace{\overset{O}{\longleftarrow}}_{Na^+} \underbrace{\overset{O}{\longleftarrow}_{Na^+} \underbrace{\overset{O}{\longleftarrow}}_{Na^+} \underbrace{\overset{O}{\longleftarrow}_{Na^+} \underbrace{\overset{O}{\longleftarrow} \underbrace{\overset{O}{\leftarrow}_{Na^+}$		
	-sodium acrylate -MBAm	$\mathbf{A}_{O^{-}Na^{+}}^{O}$		
	-sodium acrylate -acrylamide	NH ₂ O ⁻ Na ⁺	0	0

b	polymer precursors	chemical structure	monomer infused	post- polymerization
	-acrylamide -MBAm	NH ₂ NH ₂		
	-N-isopropylacrylamide -MBAm		0	
	-TRIS-acrylamide -MBAm	OH OH OH OH OH	C	0
c	polymer	chemical structure	polymer soaked	hydration
	-sodium polyacrylate M _w = ca. 2100 Da	° Na ⁺		200 1

Extended Data Fig. 7 |. Alternative hydrogel formulations.

a, Alternative monomer combinations that yield successful in crystallo polymerization and crystal expansion. **b**, Monomer combinations that lead to crystal dissolution during polymerization. **c**, A crystal soaked in a solution containing polyacrylate (molecular weight, $M_W = 2,100$ Da) dissolves upon being transferred into water. The separation between the major ticks of the ruler is 100 µm. MBAm, N,N '-methylenebis(acrylamide).

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Extended Data Fig. 8 |. **SAXS imaging of a single crystal-hydrogel hybrid in a microfluidic chip. a**, Schematic diagram of the microfluidic chip. **b**, Side-view representations of the microfluidic chip. **c**, Photograph of the microfluidic chip, mounted on beamline 4–2 at SSRL. **d**, Single-crystal SAXS diffraction patterns observed at different stages of crystal expansion and contraction. The Miller indices of each visible spot are indicated. Reflections with the highest signal-to-noise ratio $(I/\sigma I)$ are circled in red. **e**, Spot profiles of the highest- $I/\sigma I$ reflections indicated in d.

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Extended Data Fig. 9 |. **Mechanical and thermal properties of ferritin crystal-hydrogel hybrids. a**, Light-microscopy images showing the fragmentation of a native ferritin crystal and of a crystal-hydrogel hybrid upon application of external force with a needle at the location indicated with the arrow. The separation between the major ticks of the ruler is 100 μm. **b**, Temperature dependence of the SAXS profiles of native ferritin crystals and crystal-hydrogel hybrids. The small-angle reflections (that is, periodic order) in both samples are maintained at 80 °C (the maximal temperature experimentally attainable). c, Determination of the hardness and reduced modulus of native ferritin crystals and crystal-hydrogel hybrids using atomic force microscopy nanoindentation measurements. **d**, Light-microscopy images showing the expansion and contraction of a crystal-hydrogel hybrid containing Fe-loaded ferritin molecules. The separation between the major ticks of the ruler is 100 μm.

Extended Data Table 1 |

X-ray data collection and refinement statistics

Crystal A (6B8F) Crystal B (6B8G)

Data collection

	Crystal A (6B8F)	Crystal B (6B8G)	
Space group	F432	F432	
Cell dimensions			
<i>a</i> , <i>b</i> , <i>c</i> (Å)	180.40	179.95	
$\alpha,\beta,\gamma~(^{\circ})$	90	90	
Resolution (Å)	63.65-1.06	63.62–1.13	
Number of unique reflections	111189	92912	
Multiplicity	14.0 (2.4)	32.2 (13.8)	
CC1/2	0.999 (0.897)	0.999 (0.628)	
Rmerge	0.060 (0.222)	0.131 (1.131)	
<1/σ1>	24.8 (3.2)	19.1 (2.3)	
Completeness (%)	99.0 (86.4)	100 (100)	
Average mosaicity (°)	0.17	0.31	
Total solvent content (%)	57.42	57.36	
Interstitial solvent content (%)	39.72	39.62	
Refinement			
$R_{ m work}/R_{ m free}$	0.0910/0.1026	0.1029/0.1213	
Number of atoms			
Protein	1687	1699	
Ligand/ion	13	13	
Water	340	372	
B-factors (Å ²)			
Protein	8.51	9.81	
Ligand/ion	10.19	11.96	
Water	21.83	23.75	
Root-mean-square deviations			
Bond lengths (Å)	0.013	0.011	
Bond angles (°)	1.317	1.241	
MolProbity ⁴⁴ score	1.20	1.38	
Clashscore	4.12	6.99	
Ramachandran plot (%)			
Favoured	98.82	98.82	
Outliers	0.00	0.00	
Rotamers (%)			
Favoured	97.35	96.88	

Numbers in parentheses correspond to the highest resolution shell. Rmerge and CC 1/2 are measurements used to determine an appropriate high-resolution limit for XRD data.

0.52

0.015

Supplementary Material

Poor

 $DPI~(\text{\AA})^{45}$

Refer to Web version on PubMed Central for supplementary material.

0.53

0.011

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Fig. 1 |.

Packing arrangement in ferritin crystals and their expansion-contraction mediated by the infused hydrogel network. a-c, The fcc packing arrangement of ferritin crystals (Protein Data Bank identifier, PDB ID, 6B8F). The unit cell, the 200 plane and the 111 plane are outlined in black, green and red, respectively. d, Ca-mediated intermolecular interactions between ferritin molecules in the lattice. Ca^2 + ions (blue) are coordinated by two pairs of D84 and Q86 side-chains (magenta). e, Schematic representation of the formation, expansion and contraction of ferritin crystal-hydrogel hybrids.



Fig. 2 |.

Characterization of the expansion and contraction behaviour of ferritin crystal-hydrogel hybrids. a, Structural evolution of a ferritin crystal-hydrogel hybrid during the polymerization-expansion-contraction process. Black arrows indicate the addition of different solutions or water to the crystal. The numbered images (i-vi) in the top panels correspond to the selected time points shown as red circles in the bottom panel. The separation between the major ticks of the ruler is 100 μ m. b, SAXS profiles of hydrogel-infused ferritin crystals during lattice expansion, plotted against the scattering vector length *q*. The progression of scattering peaks to lower angles is indicated with grey dashed lines. Peaks corresponding to the original lattice parameters (designated with red asterisks) are visible throughout the process. c, Changes in the unit-cell parameter *a* during lattice expansion, calculated from the SAXS profiles shown in b. The schematics correspond to the red circles and are drawn to scale. The error bars were determined from the full-widths at halfmaximum of the scattering peaks. d, Expansion and contraction of a single crystal, monitored using SAXS.



Fig. 3 |.

Atomic-level structural characterization of ferritin crystal-hydrogel hybrids by XRD. a-c, XRD patterns (at temperature T = 293 K) of a ferritin crystal infused with polymer precursors (a), after polymerization and expansion (b) and after contraction with CaCl₂ (c). Light micrographs of the crystal are shown in the insets; the separation between the major ticks of the ruler is 100 µm. d, 1.06-Å-resolution structure (T = 100 K; PDB ID, 6B8F) of the contracted ferritin crystal-hydrogel hybrid, showing the electron density surrounding the Ca-mediated ferritin-ferritin interfaces and highlighting the two observed Ca coordination conformations. The electron density ($2F_0$ - F_c) map (grey) is contoured at 1.5 σ . Water molecules and Ca ions are shown as red and blue spheres, respectively.



Fig. 4 |.

Self-healing behaviour and functionalization of ferritin crystal-hydrogel hybrids. a, Light microscopy images of crystal-hydrogel hybrids, showing the self-healing of cracks that appear during Ca-induced contraction. b, Extensive cracks can also appear during crystal expansion or during the initial stages of NaCl-induced contraction, but eventually self-heal. The arrows point to the termini of the major crack extending through the crystal. c, Schematic of crack formation and self-healing through the interactions between polymer strands and ferritin molecules. d, Isotropic expansion and contraction of a crystal-hydrogel hybrid with an expandable core and an expandable shell. e, Swelling-induced fragmentation of the shell of a crystal-hydrogel hybrid with a fixed core and expandable shell. All scale bars, as well as the separation between the major ticks of the ruler in e, correspond to a length of 100 µm. See Supplementary Videos 3 and 4 that accompany this figure.