Magnetic Stiffening in 3D Cell Culture Matrices

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ABSTRACT: The mechanical environment of a cell is not constant. This dynamic behavior is exceedingly difficult to capture in (synthetic) in vitro matrices. This paper describes a novel, highly adaptive hybrid hydrogel composed of magnetically sensitive magnetite nanorods and a stress-responsive synthetic matrix. Nanorod rearrangement after application of (small) magnetic fields induces strain in the network, which results in a strong (over 10-fold) stiffening even at minimal (2.5 wt %) nanorod concentrations. Moreover, the stiffening mechanism yields a fast and fully reversible response. In the manuscript, we quantitatively analyze that forces generated by the particles are comparable to cellular forces. We demonstrate the value of magnetic stiffening in a 3D MCF10A epithelial cell experiment, where simply culturing on top of a permanent magnet gives rise to changes in the cell morphology. This work shows that our hydrogels are uniquely suited as 3D cell culture systems with on-demand adaptive mechanical properties.

KEYWORDS: adaptive hydrogels, ferrogels, magnetic iron oxide nanoparticles, extracellular matrices, mammary epithelium

■ INTRODUCTION

The cell and its microenvironment are shaped by their mutual dynamic and reciprocal interactions. In vivo, the local properties of the matrix are not constant but vary in time as a result of cellular protein deposition and remodeling. Recently, new culture systems have been developed, for which the biochemical and biophysical properties can be modulated spatially and temporally, which adds great perspectives for tissue regenerative and disease modeling. Strategies based on external stimuli such as pH, temperature, cross-linking, and light have been introduced to make the culture systems adaptive. These strategies, however, often lack reversibility, only induce small changes, pose biocompatibility challenges, and/or are restricted to 2D cell experiments.

Over the past decades, magnetic fields have been studied as external cues to tune the mechanical properties of soft hydrogels and tissue culture matrices, since they are biocompatible and easily and reversibly applied. Researchers demonstrated how magnetically responsive materials alter cell behavior by applying an external field. Although these reported hydrogel hybrids show potential in adaptive biological applications, they require high loadings of magnetic nanoparticles, which compromise compatibility with 3D cell culture environments. Our research question is as follows: Is it possible to truly benefit from the advantages that magnetic fields offer for cell culture applications, where we consider sensitivity to the field, reversibility, and tailorability as well as biocompatibility?

In this work, we present a magnetically responsive hydrogel (ferrogel) that, even at low iron oxide nanoparticle concentrations (C < 2.5 wt %) shows a reversible and large—up to an 18-fold stiffness change—at low magnetic fields (200 mT). We study the key parameters that determine the magnitude of the magnetic stiffening response, and we provide insight in forces and stresses at the microscopic scale from where macroscopic stiffening originates. The ferrogel is fully biocompatible and does not contract in magnetic fields, which renders it suitable for 3D cell culture experiments. In addition, we demonstrate the potential of magnetic stiffening in a 3D cell culture study where the mammary epithelial cells change their phenotype after culturing simply on top of a permanent magnet.

■ RESULTS AND DISCUSSION

Materials. For our experiments, we use synthetic and highly biomimetic polyisocyanide (PIC)-based hydrogels, which share the fibrous architecture and the associated intricate mechanical properties with biological gels, for instance those based on collagen and fibrin. They are, however, fully...
synthetic and therefore highly tailorable in properties. The thermoreversible PIC gels have developed into a valuable in vitro research tool,27−29 which also shows promising results in in vivo studies.30,31 For the magnetic components, we study the effect of differently shaped ferromagnetic magnetite (Fe3O4) particles (Figure 1A−C, Supporting Information Figure S1): magnetic iron oxide nanorods (MIONRs, diameter 48 nm, length 424 nm), large magnetic iron oxide nanospheres (LMIONSs, diameter 302 nm), and small magnetic iron oxide nanospheres (SMIONSs, diameter 37 nm). Note that the diameters of the nanospheres correspond approximately to the diameter and length of the nanorods. The synthesis of the PIC polymer32,33 and water-dispersible magnetite particles34−36 is described in detail in the Supporting Information.

The ferrogels were prepared by mixing a cold PIC solution with an aqueous precooled well-dispersed nanoparticle solution (Scheme 1A), followed by heating the mixture to 37 °C (Scheme 1B, Supporting Information for details). When the temperature surpasses the PIC gelation temperature (Tgel ≈ 16 °C, Figure S3), bundling of PIC chains is formed, which creates a network wherein the magnetic particles are randomly dispersed (Scheme 1B). In the presence of an external magnetic field (Scheme 1C), the particles align along the direction of the field and interact to induce fiber deformation, which results in stiffening. The gel sample is formed between a plate and TA Instruments’ temperature-controlled magnetorheology accessory with the magnetic field (maximum B = 1 T) directed perpendicular to the shear direction.

Scheme 1. (A−C) Schematic Overview of the Magnetic Stiffening Process and (D) Schematic Representation of a Magnetorheological Setup

“A Precooled PIC polymer (blue) and MIONR (black) solutions are mixed and placed between the plates below Tgel. (B) Heating the hybrids above Tgel induces polymer bundling into a semiflexible fibrous network with the MIONRs randomly dispersed. (C) In the presence of an external magnetic field, the particles align along the direction of the field and interact to induce fiber deformation, which results in stiffening. The gel sample is formed between a plate and TA Instruments’ temperature-controlled magnetorheology accessory with the magnetic field (maximum B = 1 T) directed perpendicular to the shear direction.

Magnetorheology Experiments. The magnetic stiffening experiments were carried out on a stress-controlled rheometer equipped with a magnetic field accessory (Scheme 1D), which can apply fields up to 1 T perpendicular to the shear direction. The gels are formed between (parallel) rheometer plates, and
the mechanical properties, in the absence or presence of an external magnetic field, are probed with a small shear strain. After gel formation at 37 °C, the application of a small magnetic field (100 mT) strongly stiffens the hybrids. At further increasing fields, the stiffening response levels off (Figure 1D). Note that also the corresponding ferrofluids (i.e., the dispersion of the magnetic nanoparticles in absence of the PIC gel) show an increased shear modulus but to a much less extent (Figure S2). Contrarily, PIC gels without magnetite particles do not stiffen (or soften) in the presence of a magnetic field.

We hypothesized that the strong experimentally observed magnetostiffening effect originates from reorientation and interaction of the Fe₃O₄ particles, which is transduced to the strain-sensitive fibrous network of the PIC gel (Scheme 1C). As such, hydrogels based on polyacrylamide (PAAM) that are strain-sensitive in the presence of a 1 T magnetic field (Figure 1D), we observe that at the same particle weight fraction (25 mg mL⁻¹), SMIONSs show the smallest magnetically induced stiffening response Gₘ/R = 5.0; the stiffening response of the LMIONSs (10.7) and of the MIONRs (11.8) is similar. The latter result indicates that magnetic torques have a minor contribution to the stiffening response and that the stiffening mechanism can be mainly attributed to the stress-stiffening character of the PIC matrix, which may allow for extrapolation of our results to other semiflexible fibrous networks doped with magnetic (nano)particles.

The literature indicates that larger particles and particles with larger aspect ratios are less toxic to cells because of the reduced uptake efficacy by avoiding cell wall penetration or membrane damage.⁴⁷,⁴⁸ Considering the expected viability, combined with the fact that the stiffening for both MIONRs and LMIONSs is nearly the same, we focus in the remainder of this manuscript on the application of the Fe₃O₄ nanorods.

Final characteristics of our PIC-transduced magnetic stiffening mechanism are the rate and the reversibility of the process. As the stiffening does not depend on architectural changes inside the network (Figure S5B), we find that the magnetic field (Figures S6, S7 and Movie S3) and do not leak from the hydrogel (Figure S11), highlighting that stiffening is not simply the result of matrix densification and that this strategy may be suitable for 3D cell culture experiments.

We then compare the shape and size of the particles (Figure 1D); we observe that at the same particle weight fraction (25 mg mL⁻¹), SMIONSs show the smallest magnetically induced stiffening response Gₘ/R = 5.0; the stiffening response of the LMIONSs (10.7) and of the MIONRs (11.8) is similar. The latter result indicates that magnetic torques have a minor contribution to the stiffening response and that the stiffening mechanism can be mainly attributed to the stress-stiffening character of the PIC matrix, which may allow for extrapolation of our results to other semiflexible fibrous networks doped with magnetic (nano)particles.

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Final characteristics of our PIC-transduced magnetic stiffening mechanism are the rate and the reversibility of the process. As the stiffening does not depend on architectural changes inside the network (Figure S5B), we find that the
response is very fast (Figure 1E). After a 200 mT field is applied, the stiffness increase is completed within 6 s, and the first data point is recorded, which allows for full temporal control of the mechanical properties. Moreover, the process is fully reversible; as soon as the field is removed, the stiffness returns, albeit somewhat slower. A magnetic field cycle experiment (19 times with a 200 mT field 10 min on and 10 min off) shows full stability for several hours, which gives access to dynamically changing the mechanical properties of the hydrogel.

Effects of Nanoparticle and Polymer Concentration.

The stiffening mechanism discussed so far is attributed to the complementary response of the magnetic-field-responsive MIONPs and stress-responsive PIC. Consequently, the concentration of both components is expected to affect the stiffening. Indeed, stiffening increases with the MIONR concentration, $C_{\text{MIONR}}$ (Figures 2A and S9A). In the absence of a magnetic field, the particle concentration has a negligible effect on the gel stiffness, but at a field of 1 T, we observe that magnetic stiffening increases linearly with the MIONR concentration $G_{\text{T}} \propto C_{\text{MIONR}}^1$ (Figures 2B, S8C,D). The relation even holds for nanorod concentrations as low as 3 mg mL$^{-1}$.

As can be expected, when the PIC concentration, $C_{\text{PIC}}$, is increased, the stiffness of the hybrid gels in the magnetic field also increases (Figure 2C). At the same time, the stiffness of the hybrids in the absence of the field show a stronger increase, $G_{\text{T}} \propto C_{\text{PIC}}^2$, which follows the models for semiflexible polymers. Quantitative analysis shows the linear scaling relation $G_{\text{T}} \propto C_{\text{PIC}}^1$ (Figure 2D), and we find a crossover at $C_{\text{PIC}} > 3$ mg mL$^{-1}$. At this concentration, the MIONRs (at 25 mg mL$^{-1}$) are no longer able to stiffen the network (Figure S8F). Note that the strongest relative stiffening is observed at low PIC concentration: for $C_{\text{PIC}} = 0.25$ mg mL$^{-1}$, we find a stiffening $G_{\text{T}}/G_{\text{T0}} = 18$. At typical cell culture concentrations of $C_{\text{PIC}} = 1$ or 2 mg mL$^{-1}$ (and $C_{\text{MIONR}} = 25$ mg mL$^{-1}$), the stiffening ratio $G_{\text{T}}/G_{\text{T0}} \approx 4$ and 2, respectively.

Earlier work on magnetically induced stiffening of soft materials that studied (spherical) magnetic micro/nanoparticles proposed two distinct steps in particle reorganization; first particles quickly align into chains, which then assemble into aggregates at a slower rate. In our case, both processes are expected to contribute to PIC stiffening, with a major difference that we do not expect full aggregation of the nanoparticles, since they are encapsulated in the PIC network, and the particles are expected to merely interact. Stiffening
Figure 4. Effect of magnetomechanical stimulation on MCF10A cells 3 days after seeding. (A,B) Immunofluorescence imaging of mammary gland fragments cultured in PIC/Matrigel without MIONRs in the absence (A) and in the presence of a $B = 450$ mT magnetic field (B) with nuclei stained blue and E-cadherin stained green. Without MIONRs, the morphology of MCF10A cells is not affected by the field. (C,D) Immunofluorescence imaging of MCF10A clusters in PIC/Matrigel/MIONRs matrices in the absence (C) and in the presence of a $B = 450$ mT magnetic field (D). The morphology of MCF10A cells changed in the presence of both the MIONRs and the magnetic field. Scale bar: 20 μm. (E–G) The quantification of the morphology corresponding to the above four conditions by aspect ratio (E), roundness (F), and solidity (G).

experiments in sweeps where the field is ramped up to 1 T in a 5 min sweeps (Figure S10) show that the response quickly saturated, which indicates that both processes are relatively fast. For quantification, the experimental data was fitted to a two-term exponential decay model (Tables S2 and S3), analogous to earlier work. We find that, indeed, particle alignment is fast (seconds time scale) and independent of MIONR or PIC concentrations. Secondary particle interaction, however, is a slower process with a characteristic time scale that is concentration-dependent: it becomes faster at higher C_{MIONR} and slower at an increased PIC network density.

**Quantification of Magnetic Forces and Forces inside the Network.** We then aimed to study how efficient the magnetic forces generated by the MIONRs after application of the external field are transduced to mechanical forces and how much force is exerted in the network. Since local measurements are very difficult, we then calibrated the stresses generated by the MIONRs with the stress from an external (shear) deformation (Figure 3A). Despite the difference in origin between internal and external stress, the stiffening experiments are very comparable: in the strain-stiffening experiment, a continuous stress (or prestress) is applied that stiffens the material, which is quantified by a small superposed oscillatory stress; and in the magnetic stiffening, the external magnetic field causes the stiffening, which is similarly quantified by a small oscillatory stress. This stress calibration strategy has been used earlier to analyze network stresses in myosin-contracted actin gels and in PEG gels stiffened by the thermoresponsive poly(N-isopropylacrylamide). For the strain-stiffening experiment, we measure the differential modulus $K^\prime = \delta\sigma/\delta\gamma$ (where $\delta\sigma$ and $\delta\gamma$ are the oscillatory stress and strain), which more accurately describes the stiffness of the material under an externally applied prestress. At low external strain or stress, $K^\prime = G'$, but beyond a critical stress, $\sigma_c$, the stiffening regime is entered, where the modulus becomes dependent on the applied external prestress by $K^\prime \propto \sigma^m$ where $m$ is the stiffening index, which, theoretically, is limited to $m \leq 3/2$. The parameters $\sigma_m$ and $m$ that describe the nonlinear mechanics both depend on polymer concentration; hence, strain-stiffening data was recorded for all PIC concentrations discussed in this manuscript.

We now can compare the modulus of the magnetically stiffened hydrogels (Figures 2B and 3A, dotted lines) to the strain-stiffening experiment (Figure 3A) and calibrate the stress that the magnetic particles induce with the stress applied by the rheometer to yield an average internal stress $\sigma_{\text{int}}$. Also, the magnetically induced stresses at different polymer concentrations were similarly calibrated. With all data taken together, we observe that $\sigma_{\text{int}}$ in the magnetically stiffened gels collapses to a single master curve when plotted against the product of PIC and MIONR concentrations (Figure 3B, Table S1).

From the stress $\sigma_{\text{int}}$ and the mathematically determined fiber density, we calculate the internal force on an average PIC fiber, $F_{\text{int}}$ (eqs S2, S3). As can be expected, $F_{\text{int}}$ scales linearly to $C_{\text{MIONR}}$ (Figure 3C) and weakly decreases with increasing $C_{\text{PIC}}$ (Figure 3D). Quantitatively, the piconewton internal forces that are generated by the magnetic field are of a similar order of magnitude as the contractile forces of cells that stiffen a fibrin matrix.
After the experimental analysis, we theoretically calculated whether the magnetic field that acts on the nanoparticles generates enough force to realize the observed effect. From the particle details and magnetic field gradient inside the rheometer setup, we calculated \( F_m \) the maximum magnetic force that the nanorods can apply to an average fiber in the network (eqs S4–S13, details in Supporting Information). We find that \( F_m \) values (Figure 3E,F) are only about an order of magnitude higher than the calculated \( F_m \) levels in the network, which suggest that the magnetic forces from the MIONRs are effectively transduced into a stiffening response. Our quantitative theoretical analysis of force generation provides microscale insight, which serves as valuable input for the next generations of mechanotransductive biomaterials.

**PIC/MIONR Composites as Adaptive Matrix Material.** We then studied if the magnetic stiffening matrix fulfills its promise and can be used to change the mechanical properties in cell culture environments, in particular in a realistic 3D setting. First, cell viability tests were carried out on human foreskin fibroblasts to confirm that the MIONRs are not cytotoxic (Figure S13). Live–dead staining after culturing the cells for 72 h with the MIONRs (0–50 mg/mL) in the medium did not give any indication of toxicity. The fluorescent images show the presence of the nanorods close to the cell but it remains unclear if the nanoparticles have been internalized.

To demonstrate the effect of the in situ stiffening, we selected MCF10A cells, a mammary epithelial cell line that is commonly used in breast cancer models. MCF10A cells form clusters with different morphologies, dependent on the mechanical properties of their environment. The clustering ensures the formation of a stable construct that can be visualized with traditional techniques after the strongly light-absorbing nanorods have been removed from the matrix (see Experimental Section). For cell adherence to the matrix, PIC was functionalized with the cell-binding peptide GRGDS (PIC-RGD). In addition, the PIC matrix was supplemented with basement membrane extract (Matrigel), which is critical for generating functional acini for MCF10A cells. Functionization and the addition of medium and Matrigel changes the linear and nonlinear mechanical properties of the matrix somewhat (decrease in \( G' \) and \( m \)), but the stiffening effect is still present (Figure S16). The cell culture matrix with MIONR (25 mg mL\(^{-1}\)) added stiffens a factor of 2.5 in the presence of a 450 mT magnetic field, and the stiffening response is not relaxing (Figure S17).

The MCF10A cells were seeded in PIC-RGD (2 mg mL\(^{-1}\)) both in the presence or absence of MIONRs (25 mg mL\(^{-1}\)), and the 3D cell cultures were placed in an incubator either on top of a permanent magnet (\( B \approx 450 \) mT, Figure S14) or away from the magnet.

Based on the work by the Chaudhuri lab, we expected the mammary epithelial cells to display a more invasive morphology as a result of an increased matrix stiffness. Indeed, we find that MCF10A display growth-arrested acinar structures in the absence of MIONRs or a magnetic field (Figure 4A–C). Only in the presence of both the nanorods and the external field, their morphology changed to the invasive status (Figure 4D). The morphology change was quantified by determining the aspect ratio (Figure 4E), roundness (Figure 4F), and solidity (Figure 4G) of the cells (details in Supporting Information). The analysis shows significant differences for all three quantities of the experimental group (with MIONRs and an external magnetic field) compared to all controls. Note that we did not observe any significant differences between the control groups, i.e., the application of a magnetic field to the PIC/Matrigel matrix or the addition of the nanorods to the matrix without an external magnetic field has is not influencing the MCF10A cell cluster morphology. The ease of preparation (simply mixing in the particles) and application of the stimulus (placing on a commercially available permanent magnet) makes these magnetoresponsive hydrogels available to any researcher who aims to study (temporal) mechanical effects on cell behavior.

**CONCLUSIONS**

In summary, we reported that PIC hydrogels doped with low concentrations of MIONRs show a strong magnetomechanical effect in the presence of very modest magnetic fields. The effect originates from rearrangement of the ferromagnetic nanorods, which is transduced to the strain-sensitive PIC gel. Our analysis indicates that forces generated inside these soft materials are comparable to contractile forces found in biology. Application of a magnetic field from a simple commercial permanent magnet is sufficient to change morphologies in an MCF10A cell culture experiment, which demonstrates that these hybrids are uniquely suited for 3D cell culture systems with on-demand adaptive mechanical properties. The magnetic field as a stimulus further offers the advantages of ease of use, remote control, strength modulation, and full penetration through the culture system, or even deep penetration in vivo (demonstrated by the development of MRI). Combined with the current attention for synthetic strain-stiffening hydrogels, we foresee broad application of magnetic stiffening in hydrogel platforms in, for instance, organoid development and the formation of complex tissue morphologies.

**ASSOCIATED CONTENT**

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.nanolett.1c00371.

Synthetic procedures and materials characterization, additional mechanical data, relaxation data, magnetic field gradients, and images from cell culture experiments (PDF)

Movie S1. Movie of magnetic guiding of the composite with an external magnet field (MP4)

Movie S2. Movie of magnetic guiding of the composite against gravity (MP4)

Movie S3. Movie showing that the composite does not contract (MP4)

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Author Contributions
W.C. synthesized the polymers and magnetic nanomaterials and did the mechanical characterization. W.C., P.H.J.K., and H.E. analyzed the data. Y.Z. and J.K. did the cell culture. W.C. and P.H.J.K. wrote the manuscript. P.H.J.K. supervised the project.

Notes
The authors declare no competing financial interest.

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