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Controlling Mechanical Properties of Cell-Laden Hydrogels by Covalent **Incorporation of Graphene Oxide**

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Abstract

Graphene-based materials are useful reinforcing agents to modify the mechanical properties of hydrogels. Here, we present an approach to coval ntly incorporate graphene oxide $\langle GO \rangle$ into hydrogels via radical copolymerization to enhance the dispersion and conjugation of GO sheets within the hydrogels. GO is chemically modified to present surface-grafted methacrylate groups Pathistical in small edical form about 12.100, 514-323 deviate 1420, 1502-small 201302182.

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Supporting Information

Supporting Information is available online from the Wiley Online Lives y or from the author.

(MeGO). In comparison to GO, higher concentrations of MeGO can be stably dispersed in a pregel solution containing methacrylated ζ_{relum} , $\text{del}M_A$) without aggregation or significant increase in viscosi⁺, 11 ad lition, the resulting MeGO-GAMA hydrogels demonstrate a significant increase in fracture strength with increasing MeGO concentration. Interestingly, the rigidity of the hydrogels is not significantly affected μ y the covalently incorporated GO. Therefore, our approach can be used to enhance the structural integrative and resistance to fracture of the hydrogels without in advertently ϵ' recting their rigidity, which is known to affect the behavior of encapsulated cells. The biocompatibility of MeGO-Gelma hydrogels is confirmed by measuring the viability and prolif, ration of the encapsulated fibroblasts. Overall, this stray highlights the advantage of covalently incorporating GO into a hydrogel system, and improves the quality of cell-laden h drogels. gel soletion consuming mobilized signature collear works on whom the presention of viscosity, it and if the the consumer in the soletic stress of the consumer stress and properties and interest in the theorem is the stres

Keywords

methacrylated graphene oxide (MeGO); methacrylated gelatin (GelMA); hydrogel; toughness; cell encapsulation

1. Introduction

Hydrogels are widely used as extracellular matrix (ECN_f -n imicking materials to provide suitable cellular microenvironments in various biomedical a_P plications, because the elastic polymeric network of $\frac{1}{2}$ an successfully mimic certain traits of the natural ECM structure.^[1, 2] Lydroge¹ can be designed to exhibit various chemical and physical factors to optimize cell survival and induce specific cell \mathbb{E} - \mathbb{E} -aviors.^[2] \mathbb{E} or example, hydrogels are often modified with cell recognition domains, such as A_{g} -Gly-Asp ('RGD peptide') to promote cell adhesion and \mathcal{L}_{μ} vival, and matrix metalloproteinase recognition domains to allow enzymatic degradation of the hydrogel.^[3] Recently, extensive result of efforts have been focused on studying the effect of rigidity of hydrogel on the cells, as the mechanical signals imparted by the ECM influence a diverse array of cell phenotypes as well as the differentiation fate of s em cells.^[4] **Pack Considers concentring** s of NeGO can be stably dispersed in a preschanging the control of the stable scheme interesting (MeGO can be stable along the stable along the stable scheme interesting (MeGO can called a sign

Hydrogel rigidity is most commonly modulated by controlling the candidated by density of the polymer network *via e* qustments of monotour concentration and the ratio of monomer to crosslinker.^[5] However, varying the crosslinking density in advertently affects the hydrogel toughness, *i.e.* the ability to w^{it} istand applied mechanical energy, without fracture, due to the correlation between rigidity and toughness of polymeric networks. Increasing the crosslinking density to enhance rigidity often results in brittleness, while decreasing the crosslinking density to reduce the rigidity leads to structural weal ness.^[6] $\frac{1}{4}$ nus, $\frac{1}{4}$ is challenging to improve the toughness of hydrogel while maintaining rigidity.

It has been previously shown that incorporating nanostructures with characteristic thysical properties into a hydrogel plays a significant role in determining the mechanical properties of the overall hydrogel structure.^[7,8] Graphene is a highly robust yet the vibishmacromolecular nanomaterial, composed of $s\sigma^2$ carbon atoms in a single two-dimension layer.^[9] Owing to its favorable physical properties (*e.g.* electrical and optical propeties, high mechanical strength, and biocompatibility), graph \mathcal{L} i.e-based m/ \mathcal{L} erials are increasingly used

in biomedical applications.^[10] Graphene oxide (GO), readily prepared from the oxidation of graph te, has abundant hydrophilic functional groups on the graphene layer, which allows for dispersion in aqueous media and chemical modifications, and thus has been commonly used in t iological applications over pure graphene.^[9, 11] Recent research efforts on engin ering GO-composite hydrogels with improved mechanical strength have been : ported^[12] It is suggested that incorporating GO into hydrogels would significantly enhance the toughness of nydrogels. However, the solubility of GO in biological buffers and ρ re-ge' solutions is rather limited, which impedes the homogeneous incorporation of GO within the polymeric network especially at high concentrations.

Here, we present an approach to chemically modify $G()$ to introduce methacrylate groups on the GO surface, termed methacrylated graphene oxide (MeGO), for the covalent incorporation of GO into a hydrogel system via radical copolymerization. Mechanical properties and t_2 bio degradation rates of the resulting MeGO-linked hydrogels were compared with those made with unmodified σ O to evaluate the effects of covalent con ugation. In addition, spectroscopic and microscopic methods were employed to analyze the dispersion of MeGO within the pre-gel solution and hydrogel network. Finally, the μ ioc μ mpatibility of MeCC-linked hydrogels was evaluated by measuring the viability and pro $\frac{1}{2}$: cration of $\frac{1}{2}$ icapsulated fibroblas. s.

2. Results and Discussion

2.1. Synthesis and Characterization of Methacrylated Graphene Civide (MeGO)

Methacrylate groups were conjugated onto GO by reaction with 3-(trimethoxysilyl)propyl methacryl ite (TMSPMA) to prepare methacry to graphene oxide (MeGO) (**Figure 1a**). A large number of hydroxy₁ functional groups on GO were converted to methacrylic groups *via* silanization, as evidenced by the FT-IR spectroscopy of McCO; the presence of characteristic vibrational spectral peaks corresponding to siloxyl, sily and methacrylate groups of TMS_V via (1108 cm⁻¹ (vSi-O), 1300 cm⁻¹ (vSi-C), 17¹⁰ cm⁻¹ (vC=O)), and the decrease in hydroxyl peak (3419 π m⁻¹ (θ O-H)) due to the reaction between hydroxyl groups and TMSPMA (Figure 1b). The atomic force microscopic (AFM) images of GO and MeGO showed that the chemical reaction did not all the sheet structure of GO and induce aggregation of multi le GC sheets (Figure 1c). **EXAMPLE A has also heliot** the process model with the state and the state of the state of the state of the state and the state and the state of the state and the state and the state of the state of the state of the state

2.2. Dispersion of MeGO in GelMA Solution

Inducing proper dispersion of nanoparticles within a polymer system is critical for imparting reinforcing effects of nanoparticles to the composite material.^[13] Therefore, we find examined the dispersion of MeGO in \mathcal{L} e-gel solutions to evaluate the effect of \mathcal{L} and methacrylic groups on the dispersibility of GO sheets. Here, methacrylated genann (GelMA) was chosen as a model photocrosslinkable polymer system.^[14, 15] First, varying amounts of unmodified GO or MeGO up to 3 mg mL^{-1} were added to pre-gel solutions consisting of 8 wt% GelMA and sonicated to induce designation. GO dispersed readily up to 0.8 mg mL⁻¹. However, large aggregations of GO began to appear in the pre-gel solution above 1 m_g mL⁻¹, which only disappeared after high-temperature treatment (80 °C for ¹ hour). Above 1.6 mg mL⁻¹, the pre-gel solution became viscous with highly diminished fluid mobility and **Example 12**
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 $U\checkmark$ -vis spectroscopy was used to further analyze $\mathcal L$ e dispersion of MeGO in GelMA solution. GO displays a characteristic absorption n^2 k at 231 nm, which corresponds to $\pi \rightarrow$ π^* transition and therefore identifies the dispersion of GO layers (denoted as I_1 , Figure 2h) $[17]$ McCC showed similar characteristic absorption spectra as GO, which demonstrated that dispersibility of GO layers were not affected by the presence of methacrylic groups (Figure $2c$). When CO or MeGO was incorporated within GelMA solution, the characteristic peak was red-shifted to 254 nm (denoted as I₂, Figure $2d$ and 2e), which is associated with the interaction between GO and polymers.^[18] The ratio of I_2 to I_1 (I_2/I_1), which measures the change in GO dispersion, significantly decreased (by 60 %) when the concentration of G_o was increased to 1.6 mg mL⁻¹, sug esting there was significant aggregation of GO (inset in Figure 2d). However, there was caly a small decrease in I_2/I_1 values (by 15 %), when the concentration of MeGO was increased to 1.6 mg mL⁻¹, demonstrating that MeGO ren aine a effectively dispersed in GelMA solution at a higher concentration than GO (inset in Figure 2e). It should be noted that UV-vis $\sin \theta$ and $\sin \theta$ or MeGO in GelMA at 3 mg mL^{-1} could not be obtained because high concentration of graphene oxide layers absorbed much \overline{c} . UV-vis irradiation. These results demonstrated that the presence of methacrylate groups on GO counter-the effectively prevent aggregation between GO layers, and induce better dispersion within polymer s^1 ... **EVERTS** 25). Previous studies and is a signific simulation for the same properties and the observation of the context of the same properties are the same properties and CO alternation of the same of the same of the same Page and the control of the disasonciated by high-temperature treatment
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2.3. Mechanical Properties of MeCO-GelMA Hydrogels

GelMA hydrogels incorporated with varying amounts of GO (GO-CelMA hydrogels') or MeGO ('MeGO-GelMA hydrogels') were fabricated by photoinitiated radical copolymerization (Figure 3). The hydrogels became darker with increasing amount of GO or MeGO (Figure S1a in Supporting Information). Microscopic observation of the hydrogels showed that micron-sized agglomerates began to appear in GO-GelMA hydrogels with GO concentration above 1 m , m L⁻¹, whereas 1.0 such agglomerates were observed in MeGO-GelMA hydrogels (Figure S1b in Supporting Information).

Mechanical properties of the MeGO-GelMA hydrogels were evaluated by uniaxial compression (Figure 3b and 3c). Elastic modulus, determined by the slope of the e^i astic region of the stress-strain curves, *i.e.* t_{tot} initial linear portion of the curves, increased 2.7fold when the concentration of MeGO was increased up to 3 mg mL⁻¹ (Figure 3d and 3f). On the other hand, the presence of MeGG had a more profound effect on the toughness of the hydrogels, as the stress values began to increase significantly at strains above 50%. There was an 11-fold increase in the u'umate stress of the MeGO-GelMA h / crogels when MeGO was increased to 3 mg mL^{-1} (Figure 3e, 3g, $\sin 44$.).

Elastic moduli and ultimate stress values of GO-GelMA hydrogel were similar to mose of MeGO-GelMA hydrogels up to 1.6 mg mL⁻¹. However, the elastic modulus and ultimate

stress of GO-G^{-1MA} hydrogel at 3 mg m₁⁻¹ dramatically decreased (Figure 3d, 3e, 3g, and 4b). This result is in line with the righly limited dispersibility of GO in GelMA solution at 3 mE^{-1} as presented above, which suggests that a large amount of agglomerates prevented proper hydrogel formation. As a result, these agglomerates within the hydrogels acted as structural defects, and led to structural deterioration even at lower strain.

These result; also demonstrated that in corporating GO, regardless of the mode of incorporation, had greater influence on toughness than rigidity of the hydrogels. These findings are in contrast with previous studies incomparating other types of carbon-based nanostructures, such as carbon nanotubes (C_N) and nanodiamonds (NDs), to reinforce hydrogels where both rigidity and toughness were significantly influenced. For example, Shin *et al.* demonstrated that incorporating CNT_s in to GelMA hydrogel system resulted in a significant increase in modulus $(3-f_0)^2$ increase at 0.5 mg mL⁻¹ GO), while minimally affecting the ultimate stress of the hydrogels.^[19] Furthermore, Yildirim *et al.* showed increases in both elastic modulus and tensile strength of CNT-alginate composite hyd oget.^[20] in both studies, however, the **b**rittleness of the hydrogel was also increased wⁱ in CNT, as evidenced by the decrease in ultimate strain. Behler *et el.* created NDpolyacrylonitrile composite film which showed 4-fold increase in modulus and 2-fold increase in scratch hardness when the concentration of NDs was increased up to 20 wt%.^[21] In other words, CNT or ND-incorporated hydrogels behave like a typical composite system, in which stiffer composites are generally more brittle. In comparison, GO-GelMA and MeGO-GelMA hydrogels deviate from this typical behavior with a significant increase in ultimate stress (11-fold) and a less pronounced increase in star ess (2.7-fold). It is therefore suggested that characteristic material properties of G O played a critical role in determining the mechanical properties of t_{tot} overall hydrogel structure. The highly flexible macromolecular sheet structure of GO could effectively dissipate energy applied to the hydrogel through highly aynamic conformational changes, and therefore had a more profound effect on the hydrogel toughness, whereas CNTs and NDs that do not possess such conformational flexibility also had a significant effect on the rigidity of the hydrogel. Therefore, incorporating MeGO into hydrogels could by nighly use ul to improving their mechanical toughness, vithout significantly affecting their rigidity which is a known regulator of cellular behavior **EVALUAT THE ASSOCIATE CONSULTERATION** (FOR CONSULTERATION CONSULTERATION) (FOR THE ASSOCIATE THE ASSOCIATE CONSULTERATION (FOR THE ASSOCIATE THE ASSOCIATE THE SCALE CONSULTERATION (FOR THE ASSOCIATE THE ASSOCIATE THE ASSO **AHFormatter**

2.4. Morphological Evaluation G MeGO-GelMA Hydrogels

The stark difference in mechanical properties between GO-Gella hydrogel and MeGO-GelMA hydrogels at high GO \sim MeGO content (3 mg mL⁻¹) as shown in Figures 3 and 4, suggest that the presence of methacrylic groups on $\mathcal{C}O$ sheets facilitation their integration into hydrogels even at high concentrations. To gain further insight into the effect of covalently incorporating GO into $\frac{G}{\epsilon}$, $\frac{G}{\epsilon}$ and $\frac{G}{\epsilon}$ scanning electron microscopy (SEM) was used to visualize the detailed structural features of GelMA hydrogels in corporated with high concentrations of GO or MeG λ , at λ mg mL⁻¹. The GO-GelMA hydrogel ^{4:} played highly irregular porous structure, with significant portions of the wall structure being fractured (Figure 5a). In addition, GO vas not well distributed within the nydrogel detwork, as evidenced by the uneven distribution of nighly writing and rough surface, which is caused by the presence of GO (inset in Figure 5a). Such structural irregularities were not

observed with GO-GelMA at low GO concentration (0.8 mg mL⁻¹, Figure S2a in supporting Information). It has been shown that the presence of GO can distort the p o. yme ric matrices, resulting in wrin¹ ded structures.^[22] Here, high GO content in localized areas of the network with out proper dispersion likely weakened the strength of the polyn eric network, $a \cdot d$ led $\cdot b$ fracture during the lyophilization process for sample \therefore reparation. These findings support the significant decrease in mechanical properties of GelMA hy² ogels incorporated with high concentration of GO as shown in Figure 3.

On the other hand, MeGO-GelMA hydrogets, regar lless of the concentration of MeGO, showed highly ordered porous structure, without any fractured areas (Figure 5b, Figure S2b in Supporting Information). In α^2 information, the entire surface of the hydrogel network was evenly wrinkled, which indicates that GO was well distributed throughout the hydrogel (inset in Figure 5b). These observations suggested that the covalent conjugation of GO effectively prevented aggregations, and allowed stable dispersion of the GO sheets within the hydrogels e^{γ} at high concentrations. It is well known that there is enhanced entropydriven depletion attraction between nanoparticles during polymeric network formation, because it is energetically unfavorable for the polymers to form networks surrounding the i and particles $[1^2, 2^3]$ This, coupled with the attractive interaction between GO sheets, makes GO more susceptible for aggregation or phase separation within the polymeric network. Ho vever, the covalent linkage between GQ and polymer deting the polymerization reaction likel \prime stabilized the dispersion and incorporation of GO within the hydrogel network. Furthermore, flexible shoet structures are known to increase the fracture resistance of the composite materials by reducing their Poisson ratio.^[8, 24] T_{left} fore, the significant increase in toughness of McGO-GelMA hydrogel could also be attributed to the presence of MeGO within the polymeric network allowing the material ω expand in response to external force, thus effectively dissipating the applied energy without weakening the structure. Supporting intersections (Fig. 2). The size of the the presence of CO specific and the properties of the network with survey in the size of the strength and the solicity repeated in Fig. 1, we provide the solicity repeate ²⁰ C² and the worst correlation (0.8 mg m1⁻¹, Figure S2a in each
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2.5. Biodegradation of MeGO-GelMA Hydrogels

GelMA hydrogels have been shown to undergo enzymatic degradation, as gelatin contains functional sequences recognized by collagenase.^[14, 2,] Thus, we explored the effect of covalent conjugation of GO to the GelMA hydrogels on the enzymatic degradation. MeGO-GelMA hydrogels were treated with type II collagenase, and the weight of the remaining hydrogel at various time points were measured over time. Degradation of CO-GelMA hydrogels was also evaluated as a control.

Figure 6a & 6b show the plots of $(M_t/M_0)^{1/2}$ is \therefore , where M_t/M_0 r_{-p}resents the fractional weight of the hydrogel at ume, *t*, for GO-GelMA ny trogels and MeGO-GelMA hydrogels, respectively. The plots were then fitted with eq. $\binom{1}{k}$ to obtain the degradation rates (*k_D*) of the hydrogels. k_D values for C_J -GelMA hydrogels did not change revardless of the amount of GO, which indicates physical association of GO with the GelMA network had little effect on the enzymatic cleavage of the gelatin backbone (Figure 6a & 6c). However, there was a significant decrease in k_D values with increasing amount of MeGO in the MeGO-G ϵ MA hydrogels (Figure 6b & 6c). GO sheets covalently linked to GelMA molecules were likely able to bridge the cleaved GelMA chains, and delayed the hydrogel decomposition. This

result further confirms that MeGO was at le to covalently incorporate into the hydrogel network.

2.6. Coll Encaps ulation in MeGO-GelMA Hydrogels

To assess the biocompatibility of MeGO-linked hydrogel, NIH-3T3 fibroblasts were encapsu¹ at e1 within MeGO-Gell and hydrogets (0.8 mg mL⁻¹ MeGO) and their viability and proliferation were evaluated. As a control, cells encapsulated in pure GelMA hydrogels and $GO-C_{et}MA$ hydrogels (0.8 mg mL⁻¹ GO) were evaluated. The initial viability of encapsulated cells, measured one hour after encapsulation, showed that the cell viability in MeGO-GelMA l ydrogels ($\frac{2}{2} \pm 2\%$) and GO-GelMA hydrogel (94 \pm 5%) was higher than utat in GelMA hydrogel (84 $\pm \frac{4}{3}$) (Figure S3 in Supporting Information). This suggests that the presence of GO, regardless of mode of incorporation within GelMA hydrogels, protected the cells from harmfu¹ environment during the crosslinking reaction. Shin *et al.* \ln ve t cently reported a similar finding in which cells cultured on CNT-reinforced scaffold we e protected against induced oxidative stress.^[19] The decrease in the initial viability of cells encarsulated within radically polymerized hydrogels is often attributed to the free r_{adica} and reactive oxidative species affecting the cells. It is suggested that the GO within the nydrogel may have acted as a scavenger that removes unreacted free radicals and prevented \sim death, since GO is well known to readily react with free radicals due to its electron-rich surface.^[26]

The viability of encapsulated cells was continuously monitored over the period of 7 days (Figure $7a$, Figure S4 in Supporting Information). In all conditions, the cell viability remained high throughout the experiment and the cells were able to spread and proliferate over time, den onstrating that the presence of $CO \subset$. MeGO in the hydrogels did not have any adverse effect on the long term viability of the energy sulated cells (Figure 7a & 7b). Interestingly, however, the proliferation rate was significantly higher in GO-GelMA hydrogels and MeGO-GelMA hydrogels as compared with pure GelMA hydrogels (Figure 7c). The cells became more elongated in GO-GelMA nydrogels as compared to those in MeGO-GelMA hydrogels, likely due \overline{t} the increased crosslinking density by covalent incorporation of MeGO more constrained the cells in MeGO hydrogels. However, no significant difference \ddot{r} , proliferation rate between GO-Gell \ddot{A} hydrogels and MeGO-GelMA hydrogels w.s observed, indicating the presence of GO layers within the hydrogels, not the mode of linkage to the hydrogel, was responsible for the effect on the cells, several previous studies have also reported the enhanced cell behavior on graphene-based materials.^[27] Khang *et al.* proposed that the presence of carbon nanomaterials within the polymeric matrix increased protein adsorption, que to increased surface roughness in the nano-scale, which was also shown in MeGO-C $\sin A$ (Fig. 5).^[28] Although the exact mechanism of biological responses have not been fully elucidated to Late, these results further demonstrate the feasibility of validing GO-incorporated hydrogels demonstrating high mechanical strength for tissue engineering applications. Example 12.1
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3. Conclusion

Taken together, we have chemically modified GC to introduce methacrylate functional groups onto GO and generated methacrylic graphene oxide (MeGO) in order to covalently

conjugate GO into hydrogel systems via radical copolymerization. Photocrosslinkable gelatin (GelMA) hydrogels with varying amounts of MeGO were fabricated, and the \cos ilting hydrogels displayed improved mechanical toughness with increased concentrations of N eGO, whereas hydrogels incorporated with GO showed mechanical failure at lower GO concentration than M ϵ GO. Morphologies: study of the hydrogels showed that covalently incorporating GO by using MeCO allowed stable dispersion and interfacial bonding bety een GC and polymeric network. Interestingly, the effect of MeGO on hydrogel mechanics was more pronounced on toughness than rigidity, which could be attributed to the c_c aformational flexibility of GO layer effectively assipated the energy accumulated within the polymeric network, but had smaller the rigidity. Thus, incorporating GO into $\frac{1}{4}$ hydrogel can be used to enhance the fracture strength v hile minimizing the change in rigidity which is known to influence cell behavior. Furthermore, the biocompatibility of M_{ν} GO-Genvi A hydrogels was confirmed by evaluating the viability and proliferation of encapsulated for order strategy of covalently incorporating GC presented in this study can be successfully utilized to significantly improve the structural integrity and resistance to fracture in a wide range of cell-encapsulating hydrogels v thout inadvertently affecting their rigidity. polarity (citable), the signal control is equiliblent in the signal control is the signal control of Net(k) whereas hydrogen is more and the Mongole Sink of Net(k) whereas hydrogens is an of the Mongole Sink of Net(k) whe **Example 19** (synsoms via Tidual copolymerization, Photocrosslinkable
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4. Experimental Section

Synthesis of MeGO

Graphene oxide (GO) was first prepared using modified Hummer's method.^[29] Dried GO flakes were suspended in ctuanol (1 mg mL⁻¹) and sonicated for 20 minutes, which resulted in stable homogeneous dispersion. 3-(trimethoxysily1)propyl methacrylate (Sigma Aldrich) was slowl *i* ad led to GO suspension (50 µL per each in g of GO) with sonication, and continuously stirred for 12 hours at 50 °C. The mixture was dialyzed against ethanol, and then dried under vacuum to obtain the product. MeGO was dispersed in deionized (DI) water at 4 mg m L^{-1} as a stock solution.

Synthesis of GelMA

5 g of gelatin and 0.5 g of 4-(dimethylamino)pyridine (Sigma Aldrich) were dissolved in 50 mL of dimethyl sulfox de at 50° C. Then, 2 mL of glycidyl methappy at (Sigma Aldrich) was slowly added to the mixture. The mixture vas continuously stirred for \sim heaves at 50 °C under dry N₂ gas, and then dialyzed γ anst DI water to remove byproducts. The powdered product, GelMA, was obtained by lyonhinization.

Spectroscopic analyses of MeGO

For Fourier transform infrared $(F^T\mathbf{I}\mathbf{x})$ spectroscopic analysis, dried CO or MeGO sample was first mechanically ground and pressed into a pellet with KBr powder. FT-IR transmittance spectra in a wavelength region between 500 and 4000 cm⁻¹ were acquired using a FT-IR spectrometer (Spectrum P \overline{A} , Perkin Elner).

For atomic force microscopic (AFM) analysis, GO \angle MeGO dispersed in cannol (0.05 mg) mL^{-1}) was spin coated onto a circular silicon substrate (8 mm diameter). Then, AFM images

were taken in tapping mode using a silicon-SPM tip (POINTPROBE®, NanoWorld), with a scan r te of 1.5 Hz (Digital Instruments Dimension 3000).

UV vis spectroscopy was used to analyze the dispersion of GO or MeGO within GelMA solution. Varying concentrations of GO or MeGO was dissolved in 8 wt% GelMA solution in phosphate buffered saline (PBS, pH 7.4), and sonicated for 30 minutes. Then, absorbance between 200 and 600 nm was measured using a spectrophotometer (ND-1000, Thermo Fisher).

Hydrogel fabrication

Pre-gel solution was prepared by mixing $8 \times \sqrt{6}$ GelMA with varying concentrations of GO or MeGO in PBS. 0.2 wt% of Irgacure® 2959 (Ciba) was also added to each solution as a photoinitiator. Each pre-gel solution was then placed in a custom-made cylinderical mold, and then irradic ded with UV for 2 minutes (output power of 850 mW, OmniCure® S2000) to form a hydrogel disk (8 mm diameter, 2 mm thickness). The hydrogels were then incubated in P_{3S} at 37 °C for 24 hours before characterization. **EXAMPLE 12** (**EVALUATION**) we used to calibrate the priorities of GO of 1.19 (**FV**) and the section of GO of Λ ² (**EVALUATION**) with the section of GO of Λ ² (**EVALUATION**) and Λ (**EVALUATION**) and Λ (**EV EVALUATION** (Fig. 2) the state of the properties (Fig. 2) the $\frac{1}{2}$ of $\$

 NEM was used ω analyze the morphological features of hydrogels. Hydrogels were first washed with DI water and lyophilized. Then, the dried hydrogel samples were sputter-coated with go^l α (2 nm thickness, IBS/TM200S, VCR Gr_{oup}, Inc.), then visualized with SEM (Quanta 200 FEG, FEI™) under high vacuum.

Evaluation of hydrogel mechanical properties

The hydrogel dis¹. were compressed at 1 mm min⁻¹ until they fractured using a mechanical testing system (Model 5943, *I*_{nstron} ®) equipped with a computer-based control/analysis system (B) $\text{lehi} \otimes 3$. Elastic modulus was calculated from the slope of a stress-strain curve at the \tilde{r} st 10 % strain where the curve was unear. Ultimate stress was determined as the maximum stress before the hydrogal fractured.

Evaluation of hydrogel degradation rate

The hydrogel disks were incubate in 1 U mL⁻¹ of collagenase (type II, Worthington Biochemical Co.) at 37 ° C. A ℓ various tine points, a hydrogel sample was taken out and its dried weight was measured. The results were reported as $(\sqrt{(t/T/M_0)})^{1/2}$ vs. *t*, where M_0 is the original dry weight of the hydrogel $x \cdot d$ *M*_t is the dried weight at time, f in the degradation rates (k_D) were obtained by fitting the linear region of the plots (first 15 hours) with the following equation,[31]

$$
\left(\frac{M_t}{M_0}\right)^{1/2} = 1 - k_D \cdot t \quad (1)
$$

Cell studies

NIH-3T3 fibroblasts were suspended in a pre-gel solution $(1 \times 10^6 \text{ cells} \text{ mL}^{-1})$, and then crosslinked to fabricate hydrogels via photocrosslinking, as mentioned above. The cellencapsulated hydrogels were incubated in the culture media (Dulbecco's Modified Eagle

Medium, supplemented with 10 % fetal bovine serum and penicillin/streptomycin, all purchased from Invitragen) at $37 \rightarrow$ with 3% CO₂. To measure the viability of encapsulated $cc₁'s$, the cells were flucrescently labeled with calcein-AM (green, live) and ethidium homodimer-1 (red, dead) using LIVE/DEAD® Viability/Cytotoxicity Assay kit (Invitrogen), and then visualized with a fluorescence microscope (Eclipse Ti, Nikon). The viability was quantified as the percentage of ¹² ve cells from total encapsulated cells. Proliferation rate (k_P) was obtained from the following equation, $[32]$ purchased from metropool 257 - with 5 ¹⁹ CO₂ 1 for a measure the sales the centre of the sales o **AHFormatter**

$$
\frac{w_t}{w_t} - z^k \rho^2 \qquad (2)
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where N_0 is the initial number of live cells and N_t is the number of live cells at time, *t*.

Statistical Analysis

All numerical data obtained in this work were averaged from four independent experiments. The statistical difference between two values were actermined from one-way ANOVA (Tukey's post-hoc method), and p values below 0.05 was considered statistically significant and reported here.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Figure 1.

(a) Surface functionalization of graphene oxide (GO) with methacrylate via silanization to prepare methacrylated graphene oxide (MeGO). (b) FT-IR spectra of GO (black) and MeGO (red). Characteristic peaks \sim noted in numbers. 1: $2\pi 19$ cm⁻¹ ($v_{st}(\text{O-H})$), 2: 2957 cm⁻¹ $(v_s(C-C)),$ 3: 1712 cm⁻¹ (v_s(C=O)), 4: 1300 cm⁻¹ (v_s(Si-C)), 5: 1.08 cm⁻¹ (v_s(Si-O)). (c) AFM images of GO (12h) and MeGO (trht). (Scale bar: $1\text{ }\mu\text{m}$)

Figure 2.

(a) Photographs of 3 mg m^{L-1} of MeGO (left) or C (right) dispersed in GelMA solution. UV-vis absorption spectra of (b) GO, (c) McGO, (d) GO in GelMA, and (e) McGC in GelMA. The concentration of GO or MeGO was varied from 0.15 to 1.6 mg m^{T -1}. The legends in (b) and (c) are the same for (d) and (e), r_{tot} (e), r_{tot} and r_{tot} and r_{tot} (e) represent the ratio of characteristic peaks of GO-GelMA or MeGO-CelMA at 254 nm (I₂) to that of GO or MeGO at 231 nm $(I_1)(\overline{\ast}_{r} \sim 0.05)$

Figure 3.

(a) MeGO-GelMA hydrogel is prepared by photoinitiated radical copolymerization of GelMA and MeGO. Stress-strain curves of GelMA hydrogels with varying amounts of (b) GO or (c) MeGO measured from uniaxia^l compression. (d) Elastic n odulus (*E*) and (e) ultimate stress (*U*) of GO-GelMA hyd ogels and MeG D-GelMA hyd. ogels. f, Normalized elastic modulus (E/E_0) and (g) no malized fracture energy (U/U_0) of General hydrogens incorporated with GO or MeGO. The values are normalized with respect to those of mire GelMA hydrogel (E_0, U_0) . (*p<0.05 at the same concentrations of GO and MeGO)

Fig. vre 4.

GelMA hydrogel incorporated with (a) GO or \oplus) MeGO at $\overline{3}$ mg mL⁻¹ subjected to uniaxial compression. GO-GelMA nydro gel became easily fractured, v hereas MeGO-GelMA hydrogel demonstrated resistance to fracture at high strain (70%) .

Figure 5.

Scanning electron microscopic (SEM) images of f_{m} cross sections of (a) GO-GelMA (3 mg mL⁻¹ GO) hydrogel and (b) MeGO-GelMA (3 mg m^{T-1} MeGO) hydrogel. The images on right show m_{α} ^{rified} views of designated area. (Scale h_{α} : 200 μ m)

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Figure 6.

Biodegradation of (a) GO-GelMA hydrogels and (b) MeGO-GelMA hydrogels, induced by treating the hydrogels with collagenase (¹ U mL⁻¹). The concentration of GO or MeGO was varied from 0 to 3 mg mL⁻¹ (c) The degradetion rates (k_D) of the hydrogels were obtained by fitting the linear region (first 15 hours) of the plots in (a) and (b) vith Equation 1. $(*p<0.05)$

Figure 7.

(a) Fluorescent images of fibroblasts encapsulated in GelMA, GC-GelMA and MeGO-GelMA hydrogels over time. The cells were stained with c^2 cell-AM and ethidium homodimer-1 to visualize live (arc) and dead (red) cells. (Squee bar: 100 µm) (b) Viability of the encapsulated cells at various time points. (c) Proliferation rate (*k_P*) determined from Equation 2.(*p<0.05)