Recent advances in bio-orthogonal and dynamic crosslinking of biomimetic hydrogels

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In recent years, dynamic, ‘click’ hydrogels have been applied in numerous biomedical applications. Owing to the mild, cytocompatible, and highly specific reaction kinetics, a multitude of orthogonal handles have been developed for fabricating dynamic hydrogels to facilitate ‘4D’ cell culture. The high degree of tunability in crosslinking reactions of orthogonal ‘click’ chemistry has enabled a bottom-up approach to install specific biomimicry in an artificial extracellular matrix. In addition to click chemistry, highly specific enzymatic reactions are also increasingly used for network crosslinking and for spatiotemporal control of hydrogel properties. On the other hand, covalent adaptable chemistry has been used to recapitulate the viscoelastic component of biological tissues and for formulating self-healing and shear-thinning hydrogels. The common feature of these three classes of chemistry (i.e., orthogonal click chemistry, enzymatic reactions, and covalent adaptable chemistry) is that they can be carried out under ambient and aqueous conditions, a prerequisite for maintaining cell viability for in situ cell encapsulation and post-gelation modification of network properties. Due to their orthogonality, different chemistries can also be applied sequentially to provide additional biochemical and mechanical control to guide cell behavior. Herein, we review recent advances in the use of orthogonal click chemistry, enzymatic reactions, and covalent adaptable chemistry for the development of dynamically tunable and biomimetic hydrogels.

1. Introduction

Since late-2000’s, hydrogels synthesized by click chemistry have become a unique class of biomaterials suitable for a variety of biomedical applications.1–3 In contrast to radically initiated chain-growth polymerization of (meth)acrylated macromers, which yield random crosslinks and stochastically distributed
mesh size, orthogonal click chemistries are milder, more cytocompatible, and produce hydrogel networks with less structural defects. Click hydrogels also have significantly improved mechanical properties. The selective reactivity of click chemistry handles (e.g., azide/alkyne, azide/cyclooctyne, thiol/norbornene, thiol/maleimide, etc.) not only facilitate hydrogel crosslinking, but also allows for specific tailoring of gel physicochemical properties via a secondary reaction. Click chemistry affords the crosslinked network with a wide range of bond stability, such as purely covalent bonds, cleavable linkages, and covalent adaptable crosslinks. This level of tailorable is instrumental in modulating cell fate and tissue morphogenesis, as the stability of the crosslinks often dictates how a cell respond to its extracellular microenvironment. Click chemistry also provide an elegant means of tuning hydrogel physicochemical properties in a spatiotemporally controllable manner.

An expansive ‘toolbox’ of click reagents are now commercially available for a multitude of bioconjugation and polymerization applications. In general, click gel crosslinking starts from conjugating mutually reactive click reagents to multifunctional macromers, followed by reacting the modified macromers via simple mixing. By adjusting the concentration, molecular weight, and stoichiometric ratio/functionality of the macromers, hydrogels can be fabricated with different network connectivity or crosslinking density. This in turn affects mechanical properties of the hydrogels that may guide cell fate processes. Fabrication of click hydrogels can also be facilitated by photochemistry or enzymatic reactions, which provide an additional gelation control without sacrificing the orthogonality of robust click chemistry. Click chemistry has also been widely adapted for post-gelation modification of an otherwise inert and static hydrogel network. This is commonly achieved by a secondary photochemistry, by swelling in additional click-based macromers, or by removing molecular cage capping the otherwise reactive motifs pre-conjugated in the hydrogels. The integration of multiple click chemistries in a single hydrogel results in the formation of a smart and highly responsive hydrogel network. In this review, we highlight the recent advances in major bio-orthogonal reactions used to form cell-laden hydrogels with highly tunable physicochemical properties. In particular, we focus on the use of three types of bio-orthogonal chemistries: Irreversible click chemistry, enzymatic crosslinking, and covalent adaptable network (Fig. 1). The common criterion for these three bio-orthogonal chemistries is that the crosslinking reactions between the macromers should have high specificity. Hence, we do not include hydrogels formed with random chain-growth polymerizations, supramolecular/host–guest interactions, and enzymatic reactions with low substrate specificity (e.g., acid-amine coupling).

### 2. Chemistry for crosslinking of biomimetic hydrogels

#### 2.1. Orthogonal irreversible click chemistry

**2.1.1. Thiol–ene/alloc/vinyl photopolymerization of hydrogels.** Bio-orthogonal thiol–norbornene click chemistry has emerged as an effective and versatile tool for crosslinking of biomimetic cell-laden hydrogels (Fig. 2A). Thiol–norbornene photoclick reactions are free-radical initiated but only require a low concentration of photoactivatable initiator to generate the radicals. Furthermore, thiol–norbornene crosslinking strategies do not consume oxygen and can be used to prepare hydrogels as microparticles or bulk hydrogels for cell culture. Uniquely, this chemistry can also be used to spatially pattern extracellular matrix (ECM) components to study cell behaviors. Typical initiation strategies include using UV light or visible light polymerization. Mechanistically, UV light irradiation cleaves a type I initiator (e.g., lithium aryl phosphinate, LAP)
into free radicals, which abstract protons from sulfhydryls to create thiol radicals. The thiol radicals react with nearby norbornene (NB) while regenerating additional thiol radicals for propagation of the crosslinking reactions. Alge and colleagues utilized UV light to crosslink 4-arm PEG-NB with bis-thiol containing dithiothreitol (DTT) in the presence of LAP to generate hydrogel microgels that could be further modified with other chemistries.\(^8\) In another example highlighting the modularity of this strategy, Ki and colleagues generated schizophyllan, a water soluble β glucan, hydrogels using NB modified schizophyllan macromers, DTT, and LAP.\(^{10}\) In contrast to using UV-light, visible light-initiated crosslinking often involves the use of non-cleavage type (type II) photoinitiators (e.g., eosin-Y, rose bengal), which can be excited into a triplet state for extracting protons and initiating crosslinking. Our lab has utilized eosin-Y as a visible light photoinitiator,\(^{11}\) and others have shown the use of additional crosslinking.\(^{12}\) Eosin-Y photopolymerization has recently been used in conjunction with other click chemistry (e.g., tetrazine–norbornene reaction) to assemble microporous annealed particle (MAP) hydrogels.\(^{13}\) Other materials crosslinked using thiol–ene chemistry include alginate,\(^{14}\) pectin,\(^{15}\) gelatin,\(^{9}\) and hyaluronic acid.\(^{16}\)

In addition to thiol–norbornene photopolymerization, thiol-allyloxycarbonyl (alloc) is often utilized to generate hydrogels and tether additional molecules to a network. For example, Sawicki and Kloxin developed a protocol for encapsulation of cells using alloc-functionalized peptides and multi-arm PEGSH.\(^{17}\) Similar to thiol–norbornene reaction, this approach was shown to also enable photopatterning of hydrogels and achieve biocompatible crosslinking for 3D cell culture. Kloxin and colleagues also used thiol–alloc reaction to tether collagen mimetic peptides to PEG-based hydrogels for regulating mesenchymal stem cell (MSC) fate processes.\(^{18}\) Besides fabricating hydrogels, the alloc moiety has been shown to ‘click’ to acrylate groups for modifying the crosslink network structure of PEG-acrylate hydrogels.\(^{19}\) Increasing the concentration of a mono-alloc-functionalized peptide significantly reduced the modulus of the hydrogels, resulting in variation of growth factor release kinetics.

Thiol–vinyl polymerization is also a commonly used reaction for generating a wide range of hydrogels requiring the use of natural materials such as chitosan or synthetic molecules like PEG.\(^{20,21}\) As with other thiol-based photopolymerizations, these hydrogels exhibit a high degree of tunability and modularity. For example, Hughes and coworkers developed an injectable, photocurable gelatin-based hydrogel by modifying gelatin with thiol and acrylate moieties.\(^{22}\) The injectability and cytocompatibility of the gelatin-based hydrogel was demonstrated using rabbit corneal wound model. Our lab has also used thiol–vinyl crosslinking to fabricate PEG-acrylate-based hydrogels for investigating the HIPPO pathway activation in hepatocellular carcinoma cells.\(^{23}\)

2.1.2. Michael-type addition reactions and hydrogel crosslinking. Widely used for forming cell-laden hydrogels, Michael-type additions are typically base-catalyzed reaction between a Michael donor (e.g., thiols, amines) and electrophilic Michael acceptors (e.g., maleimides, vinyl sulfones, acrylates) (Fig. 2B). The gelation kinetics of Michael-type hydrogels are highly dependent on the concentration or molecular structure of the functional groups. García and coworkers designed PEG-based thiol-maleimide hydrogels to encapsulate pancreatic islets and intestinal organoids.\(^{24–26}\) Segura and colleagues demonstrated the importance of controlling Michael-type reaction kinetics to achieve a crosslinked hydrogel with low degree of network defects.\(^{27}\) They showed that faster crosslinking kinetics (i.e., 2–5 seconds) led to heterogeneous crosslinking and peptide labeling of PEG-based hydrogels, whereas a slower gelation rate (a minute or longer) yielded hydrogels with appropriate mixing time and a more homogeneously crosslinked gel.\(^{27}\) On the other hand, Del Campo and coworkers demonstrated that multi-arm PEG-methysulfone (PEG-MS), which reacted with PEG-SH with a moderate reactivity, was particularly advantageous for hydrogel crosslinking owing to a better mixing of pre-polymer solutions.\(^{34}\)

2.1.3. Inverse electron-demand Diels–Alder (iEDDA) cross-linking of hydrogels. iEDDA reactions between diene tetratetrazine (Tz) and dienophiles (e.g., NB, trans-cyclooctetene (TCO)) are fast

![Fig. 2](image-url)

**Fig. 2** Reaction schemes of (A) thiol–norbornene photopolymerization (P.I.: photoinitiator), (B) Thiol-MAL Michael-type addition reaction, (C) iEDDA-mediated crosslinking (by Tz-NB click reaction), (D) SPAAC reaction (by azide-DBCO click reaction), and (E) Diels–Alder reactions (by furan-MAL click reaction).
and irreversible click reactions that have increasingly been used for formulating hydrogels in 3D cell culture applications (Fig. 2C). Conjugating the diene and dienophile moieties to macromers for iEDDA-based gel crosslinking is synthetically simple as numerous amine-, carboxyl-, sulphydryl-reactive derivatives are commercially available. Mooney and coworkers used carbodiimide chemistry to generate Tz- and NB-alginate macromers for facile \textit{in situ} cell encapsulation.\textsuperscript{29,30} Tz-NB crosslinking has been combined with other click chemistry to generate double network hydrogels with high mechanical strengths.\textsuperscript{31} Controlling Tz-NB reactivity is critical for providing appropriate mixing and pipetting time, which is essential for homogeneous cell encapsulation in bulk hydrogels. In an effort to control the initiation of Tz-NB crosslinking, Forsythe and colleagues synthesized dihydrogen-Tz (dHTz) that requires oxidation prior to iEDDA reaction to precisely control Tz-NB gelation. The reaction was initiated by low concentration of Horseradish peroxidase (HRP) or red-light induced oxidation in the presence of methylene blue.\textsuperscript{32,33} Recent reports also show that tetrazine can be replaced with a more stable derivative methylphenyltetrazine to slow the gelation rate and to prevent reagent degradation.\textsuperscript{34,35} Nonetheless, macromer formulations must be carefully optimized for achieving homogeneous mixing without significantly delaying gelation speed and matrix stiffness. Another potential consideration for this reaction is the generation of nitrogen gas by-product. While nitrogen gas is inert and does not pose significant toxicity, a slowly crosslinked network typically generates larger nitrogen gas bubbles, which may cause local heterogeneity in mechanical properties of the hydrogel network.

Aside from Tz-NB crosslinking, Tz-TCO can be used for rapid generation and post-fabrication of hydrogels. Tz-TCO reaction is significantly faster than Tz-NB reaction and therefore has application in injection, interfacial polymerization, and bio-printing. Jia and colleagues utilized Tz-TCO for fibrous scaffold fabrication. This was achieved by interfacial polymerization with a trifunctional TCO monomer and a PEG-based bifunctional Tz monomer, and the scaffold was demonstrated to promote fibroblast viability and migration.\textsuperscript{36} Pei and coworkers also used Tz-TCO reaction for \textit{in situ} hydrogel crosslinking, encapsulation, and release of bone morphogenetic protein-2 (BMP2).\textsuperscript{37}

2.1.4. Strain-promoted alkyne-azide cycloaddition crosslinking (SPAAC). Copper-catalyzed azide–alkyne cycloaddition (CuAAC) was one of the earliest utilized click chemistries in biomaterials-based applications.\textsuperscript{38} However, its use in formulating cell-laden hydrogel was limited due to the toxicity of copper. Copper-free SPAAC (Fig. 2D), initially established by Bertozzi and colleagues,\textsuperscript{39} has proven to maintain high specificity and reaction rates while exhibiting limited cytotoxicity, thus making it ideal for cell encapsulation.\textsuperscript{40–42} As with other click chemistries, numerous functional precursors have been developed to tune the gelation kinetics and reactivities of SPAAC. For example, Heilshorn \textit{et al.} developed elastin-like proteins (ELPs) decorated with either azide or bicyclo[6.1.0]nonyne (BCN) moieties for SPAAC-induces gelation.\textsuperscript{43} In another example, an injectable hyaluronic acid (HA) based hydrogel was fabricated using azide-conjugated PEG and cyclooctyne-conjugated HA.\textsuperscript{44} Adronov and coworkers utilized aza-dibenzocyclooctyne (DIBAC or DBCO), which has higher reactivity than other cyclooctynes, to generate PEG-based hydrogels with tunable mechanics.\textsuperscript{45,46} Other uses of DBCO include the development of microgel scaffolds for hMSC secretome analysis as described by Anseth and colleagues.\textsuperscript{47} Kloxin and coworkers also utilized DBCO to generate PEG-based SPAAC hydrogels decorated with thrombin-cleavable fluorescent proteins.\textsuperscript{48}

2.1.5. Diels–Alder reactions. Diels–Alder (DA) reactions permit spontaneous crosslinking of hydrogels at physiological pH and temperature (Fig. 2E). However, DA gelation is typically slower than other click chemistries, particularly for the commonly used diene maleimide and dienophile furan pair.\textsuperscript{49,50} Increasing the reaction rate requires an acidic environment (pH = 5.5), but this is an unfavorable condition for \textit{in situ} cell encapsulation. In this regard, Schoichet and colleagues used electron-rich methylfuran to accelerate crosslinking of HA-methylfuran and PEG-MAL hydrogel at pH 7.4.\textsuperscript{51} In another example, Heilshorn and Madl achieved a 30 second gelation rate using 8-arm PEG-fulvene and 8-arm PEG-MAL.\textsuperscript{52} DA reaction has been used in conjunction with other chemistry such as acylhydrazone bond formation or metal coordination bonding to generate double network hydrogels with self-healing properties.\textsuperscript{53–55} Other dually crosslinked networks involving DA reaction include using a rapid, thermo-responsive gelation approach, followed by slow furan-MAL crosslinking to provide the gel with enhanced mechanical stability.\textsuperscript{56}

2.2. Enzymatic crosslinking

2.2.1. Horseradish peroxidase (HRP). Enzymatic reactions are popular strategies for \textit{in situ} crosslinking of hydrogels due to their highly predictable reaction kinetics and enhanced substrate specificity. Of particular interest is the use of HRP for \textit{in situ} crosslinking of phenolic polymers (\textit{e.g.} hydroxyphenylacetic acid (HPA)), tyramine, or tyrosine-modified polymers. Upon activation by H\textsubscript{2}O\textsubscript{2} (Fig. 3A), HRP abstracts a hydrogen from the phenolic groups to generate radicals, which terminally join to form dityrosine crosslinks.\textsuperscript{57} In a recent example, HRP was used to polymerize tyramine-substituted silk (SF-TA) or gelatin (G-TA) to improve crosslinking and mechanical properties of silk-based hydrogel.\textsuperscript{58} Despite the use of H\textsubscript{2}O\textsubscript{2} at 0.01% (ca. 3.3 mM), the hydrogels were reported to sustain both 2D and 3D spreading of hMSCs.\textsuperscript{58} In an effort to mitigate the cytotoxicity and enzyme inactivation effects of exogenously added H\textsubscript{2}O\textsubscript{2}, many have exploited glucose oxidase (GOX) and glucose to supply H\textsubscript{2}O\textsubscript{2} indirectly. In a recent example, Kim and coworkers employed the GOX/HRP dual enzyme system to generate a gelatin-based hydrogel for 3D encapsulation of human dermal fibroblast.\textsuperscript{59} Gantumur \textit{et al.} reported that H\textsubscript{2}O\textsubscript{2} could be generated from self-oxidation of cysteine residues within HRP.\textsuperscript{60} To achieve self-oxidation, high concentration of HRP (300 U ml\textsuperscript{-1}) was used along with glucose as a reducing agent. Despite the use of high HRP concentration, the hydrogel system showed no cytotoxicity to the encapsulated 10T1/2 fibroblasts.\textsuperscript{60}
Beside phenolic functional groups, HRP has also been shown to generate radicals from thiols to initiate disulfide bond crosslinking. Gelatin-based hydrogel formation via this mechanism exhibited enhanced adhesiveness for long-term culture of human dermal fibroblasts (hDF). Via targeting thiol residues, HRP could also initiate thiol–norbornene gelation. In our own recent work, we showed that HRP could be used to crosslink 8-arm PEG-NB with bis-cysteine peptides for in situ cell encapsulation. Furthermore, the incorporation of tyrosine residues within a peptide crosslinker significantly increased the kinetics of HRP-mediated thiol–norbornene gelation. It was hypothesized that tyrosyl radicals were generated on the tyrosine residues that rapidly transferred to nearby cysteines, yielding thyl radicals for initiating gelation. Furthermore, these tyrosine residues remained available for mushroom tyrosinase (MT) mediated dynamic gel stiffening.

### 2.2.2. Transglutaminase

Transglutaminase (TG) is another useful enzyme that is popular for in situ hydrogel crosslinking. TG catalyzes the formation of isopeptide bonds between primary amine and carboxyamide group of glutamines (Fig. 3B). Blood coagulation factor XIII was employed to crosslink high molecular weight HA conjugated with TG substrates to form hydrogels that enabled in situ encapsulation of rat neurons. The encapsulated neurons displayed impressive neurite outgrowth, as well as axonal and dendritic speciation. TG crosslinking can be used in combination with other reactions to generate double network hydrogels. For example, Chen et al. used TG to crosslink gelatin in combination with alginate/Ca²⁺ to generate a cyctocompatible interpenetrating network (IPN) hydrogel for 3D cell culture and bioprinting. In another example, microbial TG was utilized to crosslink gelatin microgels to produce a biofunctional, injectable, and macroporous hydrogel. The porous hydrogel supported higher proliferation and migration of hDFs than the nonporous hydrogel over 2 weeks. A notable recent advance of TG-based crosslinking was the use of ultrasound to liberate liposome-encapsulated Ca²⁺, which activated TG to achieve tunable fibrin gelation.

### 2.2.3. Other enzymes

Many enzymes routinely used in cellular and molecular labeling are increasingly explored for hydrogel applications. For example, sortase A (SrtA) catalyzes...
the transpeptidation between LPXTG (where X is any amino acid except proline) and G\textsubscript{n} (n > 1) sequences to yield LPXTG\textsubscript{n} and a leaving G (Fig. 3C). The high specific SrtA transpeptidation has been used for post-gelation modification,\textsuperscript{69} controlled degradation,\textsuperscript{70} and crosslinking of PEG-based hydrogels.\textsuperscript{71} These examples have demonstrated the cytocompatibility of SrtA-mediated reaction for \textit{in situ} cell encapsulation. Uniquely, dynamic controls of hydrogel properties (e.g., on-demand stiffening or softening) could be achieved \textit{via} designing peptide crosslinkers with orthogonal specificity to different enzymes (e.g., tyrosine residues for tyrosinase, protease-labile sequence).\textsuperscript{71}

Another orthogonal enzyme ligation strategy for hydrogel crosslinking is the SpyTag–Spy Catcher system, which were engineered from the CnaB2 domain of the fibronectin adhesion protein FbaB. SpyTag is a peptide sequence (AHIVMV-DAYKPTK) containing reactive D amino acid and SpyCatcher is the protein domain containing the reactive K residue.\textsuperscript{72} Mixing of SpyTag and SpyCatcher results in the formation of isopeptide bond between the reactive D and K amino acids (Fig. 3D). This reaction was applied to assemble hydrogels that can be further functionalized with adhesion proteins to modulate human mammary epithelial cells behaviors in 3D.\textsuperscript{73} In another example, a protein-based hydrogel network was created using ELPs conjugated with SpyTag and SpyCatcher.\textsuperscript{74} In addition to the SpyTag–Spy Catcher covalent crosslinks, the gels were designed to entrap engineered p53 domains, which can dimerize to produce physically entangled interchain interactions. Hence, the resulting hydrogels could be created with various dissipation modes and would be useful for investigating cell–matrix interactions.\textsuperscript{74}

Thrombin, which catalyzes the cleavage of fibrinogen into fibrin, is also frequently used in hydrogels crosslinking. By cleaving fibrinogen at the specific Arg–Gly residues, fibrin polymerization domains are exposed and fibrin monomers can self-associate into insoluble fibrin.\textsuperscript{75} Taira \textit{et al.} exploited this highly specific reaction for the development of an electrochemically-induced fibrin gelation for encapsulation of human lung fibroblasts.\textsuperscript{76} In another recent example, Zhao \textit{et al.} used thrombin to prepare aptamer-functionalized fibrinogen injectable hydrogel that allowed for \textit{in vivo} co-delivery of multiple growth factors to promote angiogenesis.\textsuperscript{77}

Another promising enzyme is tyrosinase, which specifically oxidizes tyrosine residues into catechols, dihydroxyphenylalanine (DOPA)-quinones and eventually to DOPA-dimer linkages (Fig. 3E). Recently, a novel tyrosinase, derived from \textit{Streptomyces avermitilis}, was employed to crosslink a tissue adhesive hydrogel from tyramine-conjugated HA and gelatin.\textsuperscript{78} Comparing to tyrosinase extracted from \textit{Agaricus bisporus} or \textit{Bacillus megaterium}, this new tyrosinase exhibited a superior reactivity that significantly reduced gelation rate to less than 50 s.\textsuperscript{78}

2.3. \textbf{Covalent adaptable networks}

The effects of viscoelasticity on cell fate have received increasing attention in recent literature.\textsuperscript{79–84} Several covalent adaptable networks (CANs) have been developed to recapitulate tissue viscoelasticity. CANs are networks containing reversible bonds that undergo changes between bound and free states over time or in response to environment stimuli.\textsuperscript{85} CANs are a great toolkit for adjusting hydrogel viscoelasticity since they offer dynamically adaptable network connectivity. Hydrogels containing CANs provide a robust yet dynamic framework, from which to study how matrix viscoelasticity affects cell fate processes.

\textbf{2.3.1. Boronate ester.} Boronic acid (BA) complexes with 1,2- or 1,3-diols to produce reversible boronate ester-diol bonding (Fig. 4A). The equilibrium of the complex is sensitive to pK\textsubscript{a} of the boronic acid groups. Thus, to form stable hydrogels, the pH of the buffer solution usually has to be higher than the pK\textsubscript{a} of BA to prevent dissociation.\textsuperscript{86–88} Boronate ester bond is widely employed in cell encapsulation due to its selective and non-toxic nature. The Anseth group has recently reported a fast stress-relaxing hydrogel containing both stable azide–alkyne bonds and dynamic covalent boronate ester bonds.\textsuperscript{89}

![Fig. 4](image-url) Reaction schemes of (A) boronate–ester bond, (B) imine formation, (C) oxime ligation, (D) hydrazone bond, (E) thioester exchange, (F) Fe–catechol coordination complex.
Despite the absence of chemically and hydrolytically degradable bond, this hydrogel system could promote hMSCs spreading through dynamic linkages that were sensitive to cell-induced forces. By varying the phenylboronic acid derivatives with different p$K_a$ values, the stress relaxation rate can be systematically controlled over a broad range of timescales to capture the dynamic mechanical properties of ECM in various tissues.$^{89}$ In another work, PEG was functionalized with benzaldehydes and phenylboronic acid at each end of the chain to obtain multifunctional PEG (MF-PEG).$^{90}$ This MF-PEG simultaneously cross-linked with poly(vinyl alcohol) (PVA) through the boronate ester bonds and glycol-chitosan via imine formation to generate an adhesive and self-healing hydrogel with a unique double dynamic network (DDN) structure under mild conditions (pH $\approx 7, 25^\circ C$).$^{90}$

When incorporated within hydrogel constructs, the dynamic nature of boronate ester bonds not only gives rise to viscoelasticity, but also provides hydrogels with self-healing ability, which is uniquely advantageous for generating co-culture systems. For example, polymer 2-acylamidophenyl-boronate acid was crosslinked with PVA to form self-healing hydrogels.$^{91}$ Two hydrogels separately encapsulated fibroblasts or breast cancer cells were fused together upon close contact due to the formation of boronic acid-diol complex at the gel interfaces. The result hydrogel was used to probe directional cell migration and invasion of the co-cultured cells.$^{91}$

2.3.2. Imine/enamine. Imines/enamines (Schiff bases) are generated from the condensation reaction between amine and aldehyde (or ketone) (Fig. 4B). In general, Schiff base bonds are more stable in neutral pH and have also been used to generate hydrogels under physiological conditions.$^{92}$ For example, Zhang et al. utilized the reaction between the amino groups along the dendric oligoethylene glycol-functionalized chitosan and dialdehyde crosslinkers to generate a hydrogel at pH 7.4.$^{93}$ The hydrogels were cytocompatible for 3D stem cell culture despite the use of aldehydes.$^{93}$

Although promising for designing adaptable hydrogels, the rapid coupling and uncoupling of Schiff base linkages usually causes poor structural stability.$^{94}$ To overcome this challenge, Wei et al. developed a dual network hydrogel formed via two chemical reactions: Schiff base and Michael-type addition.$^{95}$

The reaction between the amino groups of N-carboxyethyl chitosan and the aldehyde groups on oxidized and acylated HA (OHA) constituted the dynamic hydrogel network. On the other hand, the Michael addition reaction of acrylates moieties on OHA chain with the dithiol of matrix metalloproteinase-sensitive crosslinkers yielded a network with both biofunctionality and enhanced stability. Owing to the self-healing effect of the dynamic chemistry, multiple hydrogel pieces can be fabricated with different RGD concentrations then assembled together to create a hydrogel string with RGD gradient to investigate cellular response.$^{95}$

2.3.3. Oxime. A special type of imine, oxime, is produced by reacting aldehyde or ketone with hydroxyl amine functional groups (Fig. 4C).$^{96}$ Oxime bonds are susceptible to hydrolysis, which occurs faster under acidic conditions (pH $\leq 5$).$^{97}$ However, oxime bonds still possessed higher stability than regular imines.$^{98}$ Therefore, comparing with other Schiff base gels, oxime-mediated hydrogels exhibited better mechanical properties, but still maintain the dynamic nature. It was reported that hydrogels created from alkoxyamine- and aldehyde-modified alginates permitted tunable viscoelasticity,$^{99}$ which was adjusted by simply varying the molar ratios between oxime and aldehyde. This material was used to probe the growth and migration of encapsulated B-cells.$^{99}$ In another case, to offer more control over the spontaneous oxime reaction, the DeForest group created photo-triggered oxime ligation by protecting alkoxyamine with 2-(2-nitrophenyl) propyloxycarbonyl (NPPOC) photocage.$^{100}$ Upon UV light expose, the photocage was removed, revealing the oximoyl on the PEG chain for immediate crosslinking with benzaldehyde-modified PEG and generating hydrogel within minutes.$^{100}$

2.3.4. Hydrazone. Another dynamic covalent bond that is widely used to crosslinking biomimetic hydrogel crosslinking is hydrazone, which is also a special type of imine. Hydrazone results from the condensation reaction between hydrazine and aldehyde (Fig. 4D). Similar to oxime bond, the exchange between free and complexed state of the hydrazone bonds occurs through hydrolysis and recondensation, which constitutes the dynamic property of this chemistry. In an example, the polymerization of HA-hydrazine, HA-aldehyde/HA-benzaldehyde and collagen I rendered an IPN hydrogel system that recapitulated the viscoelasticity and fibrillarity of ECM in tissues.$^{101}$ The stress relaxation in the IPN hydrogels was tuned via the two modes of stress relaxation present within the network: one from collagen and the other from the hydrazone dynamic bond. It was revealed that faster relaxation promotes MSCs spreading, fiber remodeling, and focal adhesion (FA) formation.$^{101}$ Light can also be employed to initiate hydrazone gelation. In this case, 4-arm PEG-hydrazide and PEG-nitrobenzyl was used to fabricate hydrogels.$^{102}$ Under UV-light exposure, pendant photoreactive 2-nitrobenzyl alcohol was converted into 2-nitrobenzaldehyde and reacted with PEG-hydrazine to form a hydrogel.$^{102}$

To further extend the mechanical properties as well as the applicability of this chemistry, a double-network hydrogel was crosslinked via vinyl and hydrazone bonds.$^{103}$ Since hydrazone is subjected to decomposition under acidic conditions, the hydrogel stiffness can be adjusted by incubating the gels in pH 7.4 or pH 5 buffers. This dynamic double network hydrogel was used to demonstrate the differential migratory behaviors of endothelial cells (ECs) and MSCs against gels with different stiffness. Specifically, ECs migrated much deeper into the softer hydrogel that were pre-treated with pH 5 buffer, whereas MSCs migrated more easily into the stiffer hydrogel.$^{103}$

2.3.5 Other CANs. Additional dynamic covalent chemistries for generating stress relaxing hydrogels include disulfide bond, catechol–Fe complexes, and Diels–Alder (DA) cycloaddition. To generate stress-relaxing hydrogels via thioester exchange, gels were initially crosslinked by thiol–ene photo-crosslinking of 8-arm PEG thiol macromers with the thioester d(vinyl ether) (TEDVE) crosslinker with an off-stoichiometry ratio.$^{104}$ The excess thiols reacted with the backbone thioester in a thioester
exchange reaction to afford tunable viscoelasticity (Fig. 4E).\textsuperscript{104} Using an alternative chemistry, four-arm PEG modified with dopamine was reacted with Fe$^{3+}$ to form reversible bonds (Fig. 4F).\textsuperscript{105} In addition to the dynamic bond formation, the spontaneous oxidation of catechol resulted in formation of irreversible covalent bonds causing the hydrogel to have both self-healing and shape memory properties.\textsuperscript{105} Lastly, the orthogonal DA bonds mentioned in Section 2.1 also possess dynamic nature and could potentially be used to generate self-healing and adaptable hydrogels. As mentioned above, HA modified with methylfuran, a more electron-rich diene than furan, could couple with maleimide under a physiological pH to form more stable hydrogels.\textsuperscript{51} The improvement in stability suggests slower forward and reverse DA reactions, which might affect the time-dependent mechanical properties and the self-healing effect of the hydrogels.

Even though dynamic covalent bonds provide a promising means for cell encapsulation, solely relying on this chemistry in the network structure does not provide a stable network, especially for dynamic bonds with fast dissociation constants. Furthermore, gel degradability and bio-adhesivity should also be incorporated in the network to support cell fate processes. In one example, DeForest and colleagues utilized gels with oxime bonds to encapsulate 3T3 fibroblasts.\textsuperscript{108} They noticed that the lack of cell adhesion ligands and biodegradable network resulted in cells with rounded morphologies. Similarly, by encapsulating cells using Schiff base chemistry, the Gerecht group found that cells within regions of high RGD and MMP-sensitive crosslinker concentration were significantly more elongated than regions without any bioactive peptides.\textsuperscript{95} In addition, the reversibility of dynamic covalent bonds will also affect how fast cells can spread. In particular, the Anseth group reported hMSCs encapsulated within hydrazone hydrogels demonstrated no extensive spreading due to low hydrolysis rate of aromatic hydrazone crosslinking, which restricted cell mobility.\textsuperscript{106} It was proposed that the use of more dissociative bonds would be beneficial in decreasing physical confinement and promoting cell expansion and migration. Finally, a study by Lee and coworkers reported that gels with high viscosity would not always promote cell spreading.\textsuperscript{107} Chondrocytes encapsulated within high viscosity gel secreted more ECM and spread less than those in gels with a lower viscosity. The effect of viscosity on cells behavior is still relatively unexplored and not yet fully understood, further development of hydrogels with tunable viscoelasticity would certainly further decipher the physiological significance of matrix viscoelasticity.

3. Methods for dynamic tuning of hydrogel crosslinking

Tuning the crosslinking density of cell-laden hydrogels post-gelation is of paramount importance for studying dynamic biological processes, including tissue fibrosis, tumor progression, or stem cell differentiation. Dynamic stiffening (or softening) of ECM is a result of excess deposition and crosslinking (or breakdown) of matrix proteins [e.g., collagen fibers].\textsuperscript{108} Strategies to control dynamic hydrogel crosslinking post-gelation include photopolymerizations, orthogonal click chemistries, and enzymatic reactions.

3.1. Dynamic stiffening of hydrogels

Irreversible stiffening of hydrogels is useful to mimic tissue fibrosis and tumor stiffening. This is often achieved by performing secondary crosslinking in a cell-laden hydrogel in the presence of additional macromers and initiators. Anseth and colleagues developed dynamic PEG-based thiol–norbornene hydrogels by diffusing additional 8-arm PEG-NB, 8-arm PEG-SH, and LAP into the gels, followed by UV-light induced secondary crosslinking.\textsuperscript{109} Also utilizing light-based secondary crosslinking in the presence of additional photoinitiator LAP, Burdick and coworkers achieved dynamic stiffening in thiol-methacrylate Michael-type hydrogels crosslinked with excess methacrylates.\textsuperscript{110} Alternatively, Engler and colleagues implemented a two stage photopolymerization to stiffen HA-methacrylate hydrogels by simply varying the UV-light irradiation time.\textsuperscript{111} In a recent study, the Anseth group exploited the ability of DBCO group to undergo radical-induced photocrosslinking for preparing stiffening hydrogels to probe myoblast mechanotransduction.\textsuperscript{104} This approach eliminated the need to diffuse additional macromers as long as the initial network contained excess cyclooctyne (relative to azide). Rosales and coworkers used o-nitrobenzyl crosslinkers and free methacrylate groups within a hydrogel fabricated by Michael-type addition to achieve UV-mediated photocleavage and visible-light and LAP-mediated stiffening.\textsuperscript{112} In another example, Bryant and colleagues demonstrated that through sequential hydrogel deposition and thiol–norbornene photopolymerization in a cylindrical mold, hydrogels with multiple, discrete regions of stiffness can be created.\textsuperscript{113} Alg and coworkers used microgel packing in conjunction with thiol-norbornene photopolymerization to generate microporous annealed particle (MAP) hydrogels with an $E$ gradient between 9.8 to 29.2 kPa.\textsuperscript{114} Using the MAP gels, the researchers revealed the influence of gradient stiffness on hMSC proliferation and cell volume. Anthracene photodimerization can also be used to stiffen PEG-based hydrogels.\textsuperscript{115} Specifically, PEG-anthracene hydrogels were formed by UV light-mediated [4+4] photodimerization of anthracene functional groups, and stiffening was achieved by additional dimerization through further UV light treatment.

In addition to UV-light based photo-stiffening, our lab has developed a visible light induced tyrosine dimerization approach for achieving secondary crosslinking of PEG–peptide hydrogels.\textsuperscript{116} Dynamic stiffening was initiated by diffusing Flavin mononucleotide (FMN) into the PEG–peptide hydrogels containing tyrosine residues, followed by visible light exposure. Additional methods to achieve spatial stiffening of hydrogels include the use of protecting groups in conjunction with thiol-VS crosslinking.\textsuperscript{117} The otherwise reactive thiol groups were isolated using a protecting group. Rosales and coworkers used 8-arm PEG-NB and 8-arm PEG-N3 as the initial network contained excess cyclooctyne (relative to azide). Rogers and coworkers used thiol–norbornene and thiol–methacrylate Michael-type hydrogels crosslinked with excess methacrylates.

Our lab has also utilized visible light induced tyrosine dimerization to achieve spatial stiffening of hydrogels. We have developed dynamic PEG-based thiol–norbornene hydrogels by implementing a two stage photopolymerization to stiffen hydrogels by simply varying the UV-light irradiation time. This approach eliminated the need to diffuse additional macromers as long as the initial network contained excess cyclooctyne (relative to azide). We have also utilized visible light induced tyrosine dimerization to achieve spatial stiffening of hydrogels. We have developed dynamic PEG-based thiol–norbornene hydrogels by implementing a two stage photopolymerization to stiffen hydrogels by simply varying the UV-light irradiation time. This approach eliminated the need to diffuse additional macromers as long as the initial network contained excess cyclooctyne (relative to azide). We have also utilized visible light induced tyrosine dimerization to achieve spatial stiffening of hydrogels. We have developed dynamic PEG-based thiol–norbornene hydrogels by implementing a two stage photopolymerization to stiffen hydrogels by simply varying the UV-light irradiation time. This approach eliminated the need to diffuse additional macromers as long as the initial network contained excess cyclooctyne (relative to azide).
free thiols that could react with pendant VS groups in the network. Enzymes can also be employed to stiffen hydrogels. In contrast to photopolymerization which requires a light source and radical species for initiating secondary crosslinking, enzymatic reactions rely only on diffusion-reaction kinetics. Enzymatic stiffening of hydrogels is a cytocompatible strategy that can be modeled using Michaelis–Menten kinetics. Our lab has reported tyrosinase-induced dynamic hydrogel stiffening. Tyrosinase converts mono-phenolic motifs (e.g., tyrosine, hydroxyphenylacetic acid or HPA) into DOPA dimers, resulting in dynamic stiffening of PEG–peptide and gelatin/HA hybrid hydrogels. The latter study revealed a synergistic effect of hydrogel stiffening and presence of HA on epithelial-to-mesenchymal transition (EMT) of pancreatic cancer cells (Fig. 5A). In another example, Jia and colleagues explored Tz-TCO reactions to stiffen hydrogels. TCO-modified HA was diffused into a hydrogel network crosslinked by thiol-acrylate reaction. Notably, the gels were prepared with excess Tz moieties, which permitted secondary crosslinking in the presence of additional TCO-modified HA. Interestingly, while the same strategy was effective in dynamic labeling of cell adhesive ligands (e.g., TCO-RGDS), stiffening with TCO-HA was not as significant.

3.2. Dynamic softening of click crosslinked hydrogels

Irreversible softening of hydrogels may be achieved by hydrolysis, enzymatic cleavage, or photo-lysis of the crosslinkers. In addition to hydrolysis and enzymatic linker cleavage, photo-labile o-nitrobenzyl groups are increasingly used for controlled hydrogel softening. For example, Forsythe and coworkers synthesized o-nitrobenzyl ester-containing PEG-methacrylate for crosslinking with gelatin-methacrylate into photodegradable hydrogels. The o-nitrobenzyl groups were cleaved upon exposure to 365 nm light, causing a reduction of crosslinking density and softening of the hydrogels. In an effort to tune the photodegradation rate of hydrogels, nitrobenzyl–ester, –amide, –carbonate, and –carbamate were synthesized and incorporated into hydrogels. It was determined that carbamate linkers had the fastest photodegradation rate, which was useful in temporal control of protein release (Fig. 5B). Anseth and colleagues used addition–fragmentation chain transfer to degrade PEG-based hydrogels crosslinked by allyl sulfide-containing linker. Photodegradation was controlled by LAP and mono-functional PEG-SH induced thiol–allyl sulfide exchange.

Similar to stiffening strategies, softening of hydrogels can be attained using visible light irradiation. In particular, ruthenium polypyridyl complexes were used as a photolabile linker in CuAAC-crosslinked PEG hydrogels. The linker was cleaved upon exposure with visible light (400–500 nm), resulting in hydrogel degradation. Perylene, which is cleaved at 520 nm light, is another example of a visible light photosensitizer used for hydrogel degradation. Forsythe et al. used perylene in conjunction with o-nitrobenzyl groups to induced wavelength-dependent photodegradation of SPAAC-crosslinked hydrogels.

3.3. Reversible chemistry for tuning hydrogel crosslinking

Many tissue morphogenesis events involve both matrix stiffening and softening at different timescales and magnitudes. A few reversible conjugation chemistries are exploited for recapitulating...
these dynamic events, including light-mediated azobenzene cis-trans isomerization, and reversible enzymatic reactions.

Photoswitchable chemistry has been used for reversible stiffening of cell-laden hydrogels. For example, Anseth and colleagues formulated Michael-type addition hydrogels with PEG-VS and thiol-containing peptide-azobenzene crosslinkers. By exposing the hydrogel to 365 nm light, the azobenzene-containing linkers underwent isomerization into the cis configuration, resulting in hydrogel softening. On the other hand, when exposed to heat or visible light, azobenzene was converted to the trans-configuration, thus stiffening the hydrogel. Other uses of azobenzene isomerization to reversibly stiffen and soften hydrogels showed the effect of stiffness on hMSC spreading on poly(acrylamide) hydrogels (Fig. 6A).

Switchable proteins (i.e., proteins that can be reversibly converted between monomeric and oligomeric states) are also exploited to reversibly tune hydrogel stiffness. A photoswitchable protein known as Dronpa145N, which can be converted to a tetrameric state under 400 nm light and back to a monomeric state under 500 nm light, was tethered to 4-arm PEG molecules. By alternating the light exposure, the hydrogel crosslinking density could be significantly modulated to approximately 5-fold of the initial modulus. Another photoswitchable pair used for modulating the hydrogel crosslinking density includes LOV2 and J helices. The molecules were expressed in bacteria as azide-LOV2-J-azide and crosslinked into a hydrogel network using SPAAC (Fig. 6B). Treating the hydrogels with UV-light created a conformational change between the LOV2-J pair, lengthening the molecule on the order of tens of Angstroms. The lengthening of the engineered proteins softened the bulk hydrogel mechanics. The LOV2-J domain was returned to its initial length upon removal of light exposure, leading to recovery of the initial hydrogel stiffness.

Supramolecular interactions, particularly guest–host interactions, lend themselves to dynamic modulation of hydrogel properties. Our lab and others have used supramolecular interactions to reversibly stiffen and soften hydrogels. For example, thiol–norbornene hydrogels immobilized with β-cyclodextrin (β-CD) could be stiffened by diffusing multi-functional PEG-adamantane. The additional supramolecular crosslinks were stripped by subsequent diffusion of soluble β-CD, which competed with immobilized β-CD for the adamantane crosslinkers. Azobenzene also interacts with CD functionalities at differing binding affinities based on the state of azobenzene isomerization. Cheng and colleagues exploited this phenomenon for stiffening and softening hydrogels consisting of α-CD, azobenzene, and N-isopropylacrylamide (NIPAAm). Inducing trans-azobenzene to cis-azobenzene by UV-light treatment softened the gel whereas infrared radiation (IR) thermally induced stiffening of the hydrogel by converting NIPAAm to a hydrophobic state.

Certain enzymes exhibit reversible conjugation/cleavage of the peptide substrates. For example, the transpeptidase SrtA reversibly ligates LPXTG and G6 sequences. Our lab has reported PEG-based hydrogels with unique peptide designs that rendered the hydrogels susceptible to reversible SrtA-mediated stiffening and softening (Fig. 6C). Specifically, the hydrogels were crosslinked by multi-arm PEG-NB and bis-cysteine-containing peptide linkers flanked with the two SrtA substrates (i.e., GGG-C-GGG-C-LPRTG). Following thiol–norbornene gelation, the two SrtA substrates became pendant peptides that were accessible for SrtA-induced ligation, which increased the cross-linking density of the hydrogel. Subsequent incubation with SrtA and soluble glycine peptides or glycynamide resulted in cleavage of the additional LPRTGGG crosslinks, thus softening the hydrogels.

3.4. Methods and examples to tune stress-relaxation

It has been established that cells actively respond to matrix viscoelasticity; however, the mechanisms and biological pathways responsible for these behaviors are still not well understood. Engineered hydrogels with tunable relaxation timescales would greatly facilitate the study of mechano-responsive molecular pathways. Typically, hydrogel stress relaxation can be controlled by using a network containing dynamic reversible bonds. In one example, PEG hydrogel networks were formed from mixtures of two phenylboronic acid (PBA) derivatives with different pKₐ, 4-carboxyphenylboronic acid, and o-aminomethylphenylboronic acid. Since the stability of the networks depended heavily on the pKₐ of the PBA groups, the relaxation time and the mechanical response to dynamic shear stress were exquisitely controlled by the concentrations of the PBA derivatives. Similarly, a hydrazone hydrogel was formed with two hydrazone-derivatives of different binding constants: alkyl-hydrazone (aHz) and benzyl-hydrazone (bHz). The average stress relaxation times, ranging from hours ($4.01 \times 10^3$ s) to months ($2.78 \times 10^6$ s), were modulated by varying the percentage of aHz and bHz within the same hydrogel network. Chondrocytes encapsulated in hydrogels exhibiting faster stress relaxation and deposited more ECM (e.g., collagen and glycosaminoglycans) as compared with gels with slower relaxation time.

Photochemistry can also be exploited to control hydrogel viscoelasticity. Tamate et al. described a cytocompatible and dynamic hydrogel comprised of ABA triblock copolymers, where A blocks contained NIPAAm and coumarin acrylate and B block contained PEG (Fig. 7A). The hydrogel was initially formed via physical crosslinking through thermo-responsive NIPAAm. Under UV light irradiation, the coumarin moieties dimerized, causing a significant increase in elasticity to the stress-relaxing hydrogels. Likewise, Carberry et al. established a technique to turn a stress-relaxing hydrogel into an elastic gel. In this contribution, viscoelastic hydrogels were prepared with off-stoichiometric ratio of 8-arm PEG thiol and 8-arm PEG-thioester-NB via photoinitiated thiol–ene polymerization. The excess thiols served as a trigger to induce radical-mediated dynamic thioester exchange. This was achieved by swelling the viscoelastic thioester gels in LAP and 5-norbornene-2-carboxylic acid, followed by UV-light exposure to consume the remaining thiols in the gels. Consequently, the degree of thiol–thioester exchange was reduced, so was the hydrogel viscoelasticity, in the presence of cells. The utility of this unique dynamic hydrogel was demonstrated by examining viscoelasticity-induced YAP/TAZ signaling.
nuclear translocation in fibroblasts.\textsuperscript{138} The usage of photochemistry has, therefore, further expanded the toolkits for tunable viscoelastic hydrogels where user-defined spatial and temporal control can be incorporated. An alternative approach that could potentially be used for dynamically tuning viscoelasticity is reported by Accardo et al., where hydrogel mechanics were tuned by light-induced dynamic covalent diol-boronic acid crosslinking (Fig. 7B).\textsuperscript{139} Under light exposure, the BA conjugated with azobenzene can switch between $E$ and $Z$ isomers, resulting in changes in $pK$ of boronic acid groups. Using this method, the dynamic covalent bonds can break and reform irreversibly on demand, which would be beneficial in isolating the cell response specifically to changes in viscoelasticity.\textsuperscript{139}
Viscoelasticity can also be dynamically introduced in a hydrogel matrix. The Caliari group devised a method where viscoelasticity and stiffening of a hydrogel was introduced simultaneously via the formation of βCD–adamantane host–guest bonding and additional thiol–ene crosslinking (Fig. 8A). To achieve this, an elastic hydrogel was first generated with dual-functional HA-βCD-norbornene via photo-mediated thiol–ene chemistry. To trigger stiffening and increase viscoelasticity, hydrogels were incubated with LAP and adamantane conjugated–thiol peptide (Fig. 8B). Following UV light exposure, adamantane–thiol peptide reacted with left-over norbornene within the hydrogels. Since stiffening was achieved with the introduction of additional βCD–adamantane binding, both the storage and loss moduli of the hydrogel were increased simultaneously.

4. Methods to control presentation of bioactive ligands

As with modulation of crosslinking density, designing materials to recapitulate dynamic presentation of biomolecules would prove invaluable for modeling disease progression and for promoting tissue morphogenesis. Methods to tune ligand presentation are similar to that of controlling the crosslinking density of hydrogels. Instead of introducing additional crosslinks, a chemically reactive moiety can be attached to the biomolecule of interest for its tethering into the hydrogel network. On the other hand, a chemically labile linker can be used to permit stimuli-responsive cleavage of immobilized biomolecules. Reversible chemistry can also be employed to achieve cyclic addition and removal of biomolecules.

![Fig. 7](URL) Strategies for tuning hydrogel viscoelasticity. (A) Chemical structure of the ABA triblock polymer and the modulation of viscoelasticity through photo-induced dimerization of coumarin. Reprinted with permission from ref. 137. Copyright 2016, American Chemical Society. (B) Photo-switch control of a dynamic covalent crosslink stability using equilibrium constants for esterification of an o-azobenzene boronic acid. Reprinted with permission from ref. 139. Copyright 2018, Royal Society of Chemistry.

![Fig. 8](URL) (A) Elastic hydrogel system formed by covalent thiol–norbornene crosslinks vs. viscoelastic hydrogel system formed by thiol–ene photo-chemistry and supramolecular interactions between CD-HA and Ad-thiolated peptide. (B) Secondary introduction of covalent and supramolecular crosslinks to modulate stiffness and viscoelastic properties. Reprinted with permission from ref. 140. Copyright 2019, American Chemical Society.
4.1. Photopatterning and cleavage

Photopatterning and/or cleavage of proteins and small molecules are increasingly used for dynamic cell culture applications. Compared with other approaches, photopolymerization affords spatial control of ligand immobilization by using diffusion of initiators and a photomask. While patterning cysteine-bearing peptides is well established, whole-protein conjugation is more challenging. Dai and colleagues demonstrated spatial-temporal patterning of whole protein by conjugating thiolated vascular endothelial growth factor (VEGF) in thiol–NB hydrogels with excess NB groups. Reversible photochemistries provide further control of ligand presentation. For example, Tirell and DeForest used a photo-sensitive oxime-ligation regime along with o-nitrobenzyl ester functional groups to tether and remove bovine serum albumin (BSA) into a SPAAC-crosslinked hydrogel network. Photoaddition of the protein occurred by incorporating caged-alkoxyamines that were removed by light exposure, which permitted reaction with the aldehyde-functionalized proteins. As the proteins are also functionalized with o-nitrobenzyl moieties, they were readily cleaved with additional light exposure. In another example, Anseth and colleagues showed that thiol–ene chemistry can be used to conjugate thiol-radical-conjugated protein (i.e., transferrin) to allyl sulfide containing hydrogels. Release of protein was achieved by a second thiol–ene reaction with monofunctionalized PEG-SH. Incorporating bromohydroxy-coumarin, a photocage molecule, into a hydrogel network also allows controlled protein release. Upon treatment with...
UV-light and near-IR, the photocage was cleaved, resulting in tunable payload release. Deforest and colleagues used photochemistry to tether numerous proteins, enzymes, and growth factors site-specifically to hydrogels (Fig. 9A). As opposed to random modification, the proteins were engineered with site-specific reactive handle. Under visible light exposure, the photocleavable protein (PhoCl) underwent irreversible β-elimination and peptide backbone cleavage, leading to site-specific protein patterning.147

4.2. Enzymatic addition and removal of ligands
Enzymatic reactions can be used to immobilize and remove whole proteins that have been tagged with relatively simple substrates. Particular examples include SrtA-mediated transpeptidation, SpyTag/Spycatcher, and SnoopTag/SnoopTCatcher interactions. As alluded to earlier, enzymatic reactions require biological substrates; therefore, sequences can be translationally added to proteins of interest for site-specific labeling. Deforest and coworkers utilized sortase-mediated transpeptidation to install a variety of orthogonal click chemistry reactants.148 For example, green fluorescent protein (GFP) was tethered to a hydrogel and later removed using the above mentioned combined photodeprotection-oxime and α-nitrobenzyl regimes. The sortase-sensitive sequence conferred site-specific labeling to reduce any potential detrimental effects of non-specific protein labeling that may occur. Sortase A has also been used for patterning nerve growth factor (NGF) into hydrogels containing photocaged oligoglycine substrate.149 Conducting two-photon uncaging at precise locations within the hydrogel enabled the reaction between LPXTG moieties conjugated to Avidin/Biotin-NGF and available GGG moieties in the hydrogel. SpyTag–SpyCatcher reaction also enables the tethering of bioactive molecules in hydrogels. For example, West and colleagues generated hydrogels that could be tagged with RGDS molecules by incorporating SpyTag molecules in the network. Dynamic ligand tethering occurred when additional SpyCatcher-RGDS was diffused into the hydrogel (Fig. 9B).150 In another example, SnoopTag/SnoopCatcher was used in conjunction with SpyTag/SpyCatcher for crosslinking and decorating HA-based hydrogels with EpCAM and E-cadherin.73 Through genetic engineering, Griffith and colleagues exploited the reversible SrtA transpeptidation for addition and removal of network-immobilized epidermal growth factor (EGF), which was tagged with SrtA substrate GGG at the protein N-terminus (i.e., NH₂–GGG–EGF).69 By incubating the gel with additional glycine and sortase, GGG–EGF was cleaved and removed from the hydrogels.

Invented by the Hahn lab, LOVTRAP is a pair of light-sensitive proteins that consist of LOV2 domain that binds to a small protein Zdark (Zdk) only in the dark.151 Upon irradiation with blue light wavelength (400–500 nm), Zdk readily dissociates from LOV2 and would release any protein that is conjugated to this domain. Utilizing this light-sensing protein system, the West lab developed a PEG-based hydrogel system immobilized with the LOV domain, which provides light-sensitive presentation of proteins within the PEG hydrogel.152 In particular, the PEG–LOV hydrogel network were able to reversibly bind and release mCherry-Zdk in response to blue light irradiation. When the light was removed, LOV2 underwent thermal relaxation and allowed additional Zdk-mCherry to rebind to the network. Since the LOVTRAP protein pair is highly orthogonal and can be controlled within the visible light range, it can be used in the presence of cells to modulate biochemical changes within the hydrogels.

4.3. iEDDA-mediated addition of ligands
Tetrazine–norbornene reaction is a synthetically accessible strategy for immobilization of bioactive moieties within a hydrogel. This can be achieved through simple diffusion of a tetrazine-modified protein into a hydrogel with excess norbornene functional groups. Jia and colleagues utilized Tz-TCO reaction to tether RGD dynamically into a hydrogel crosslinked with excessed Tz motifs. The infiltration of TCO-RGD molecules led to biochemical modification of the otherwise inert hydrogel (Fig. 9C).121 Additionally, Alge and colleagues showed that Tz-NB crosslinking could be used in conjunction with thiol–ene photochemistry to generate and label microgels fabricated with excess NB functionalities.8

5. Conclusion and future outlook
The needs for creating biomimetic microenvironments for cell-laden hydrogels are rapidly evolving. Up until the early 2000s, research efforts were focused on synthesizing hydrogels with pre-determined properties (e.g., crosslinking density and hydrolytic degradability).153–155 The next decade (2000 to 2010) had witness the rapid development of cell-responsive hydrogels where the materials properties could be modulated by cellular activity (e.g., protease-labile network and cell traction force induced matrix deformation).156–160 In the past 10 years, a diverse array of chemistries has been exploited for creating cell-laden hydrogels with on-demand tunable properties (e.g., spatiotemporal patterning of matrix mechanics and presentation of bioactive signals).161–164 This new class of dynamic and biomimetic hydrogels are able to recapitulate and/or direct multiple aspects of cellular fate processes. These materials have been widely used for disease treatment and modeling, tissue regeneration, and even diagnostic applications.

The ability to tune the physicochemical properties of cell-laden hydrogels at will stems from the elegant integration of multiple orthogonal chemistries. A recent report from the Anseth group exemplifies this strategy, where a photoinitiated thiol–yne reaction, an azide–alkyne cycloaddition, and a methyltetrazine-TCO iEDDA reaction were integrated in a hydrogel network to independently control the presentation of several bioactive proteins to valvular interstitial cells (VICS).165 Going forward, careful considerations must be taken when and removal of residual, potentially immunogenic initiators and unreacted monomers prior to using the hydrogel as an injectable or implant. Dynamic tuning viscoelasticity strategies can be applied to build tissue models that experience loss of elasticity, or elastin-related disease (for example, valvular aortic stenosis
that caused by elastin mutations leading to reduced levels of functional elastin). Double network (DN) crosslinking and interpenetrating hydrogels using both reversible and irreversible chemistry to help increased stability, mechanical properties, and allow for integrating different biopolymers to better mimic the hierarchical structures and heterogeneity of ECM. For example, one potential is the use the first network of the DN hydrogels to afford tunable elastic moduli while using the second network to afford controllability in matrix stress-relaxation. Another integrated strategy is to engineer therapeutically relevant growth factors or cytokines with clickable handles for their spatiotemporal presentation in the cell-laden hydrogels.\(^{166}\) This will increase the biomimicry of the hydrogels as whole proteins contain information that may be lost in short biomimetic peptides. Other dynamic properties such as self-healing, shear-thinning, or shear-thickening can also be integrated within the orthogonally crosslinked hydrogels to support multi-tissue co-culture, bioprinting, and tissue regeneration. Reversible host–guest interaction or metal ion chelation may be integrated with any of the bio-orthogonal click chemistries reviewed above to create dynamic matrices with sustained release of therapeutically relevant or immunoregulatory molecules.\(^ {167-169}\) Host–guest interactions may also be designed within a click hydrogel to generate hydrogels with tunable mechanics and bioactive ligands (e.g., RGD peptide).\(^ {170,171}\)

To broaden the applications of dynamic and biomimetic cell-laden hydrogels in biomedicine, there is a need to simplify the chemical synthesis process using non-toxic reagents. In order to fully capitalize on the increasing synthetic ability to modular hydrogel properties, there is also a need to utilize non-destructive imaging modality that permits longitudinal monitoring of cell fate processes. Methods to increase image resolution like machine learning and deep neural network can be utilized to analyze and extract more information from images. While traditional molecular biology techniques were adequate for hypothesis-driven studies, the benefit of dynamic hydrogel platform can only be realized with high content images. While traditional molecular biology techniques were adequate for hypothesis-driven studies, the benefit of dynamic hydrogel platform can only be realized with high content analysis of cell fate. This can be accomplished via integrating the dynamic hydrogels with high throughput screening, 3D printing, and other automation tools. Finally, in the age of a global pandemic, there exist an opportunity for using click hydrogels for targeted DNA or RNA delivery, as biosensors for pathogen detection and treatment (e.g., CRISPR controlled release and diagnostic devices).

**Conflicts of interest**

There are no conflicts to declare.

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