



Supramolecular polymeric biomaterials

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Polymeric chains crosslinked through supramolecular interactions—directional and reversible non-covalent interactions—compose an emerging class of modular and tunable biomaterials. The choice of chemical moiety utilized in the crosslink affords different thermodynamic and kinetic parameters of association, which in turn illustrate the connectivity and dynamics of the system. These parameters, coupled with the choice of polymeric architecture, can then be engineered to control environmental responsiveness, viscoelasticity, and cargo diffusion profiles, yielding advanced biomaterials which demonstrate rapid shear-thinning, self-healing, and extended release. In this review we examine the relationship between supramolecular crosslink chemistry and biomedically relevant macroscopic properties. We then describe how these properties are currently leveraged in the development of materials for drug delivery, immunology, regenerative medicine, and 3D-bioprinting (253 references).

1 Introduction

It is a privilege to live in a time where the global life expectancy continues to increase. Over the past 30 years the fraction of humans aged 65+ has increased over 140%; multiple countries

contain over a fifth of their population in this bracket! However, as the human body ages it becomes more fragile. A rising global life expectancy has been met with increased incidences of cancer and degenerative diseases. We are now left with the challenge of an aging population seeking improved quality of life. Addressing this problem, the health-care community has looked towards smarter and more creative medical solutions through the development of novel therapeutic agents and strategies, or innovative delivery of existing therapeutics. Common to both approaches is an increasing reliance on creative and adaptable biomaterials.

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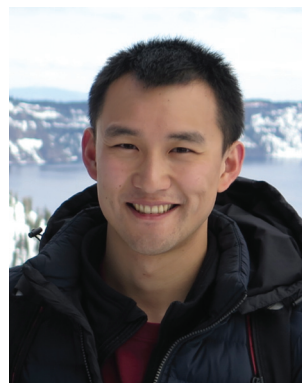
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Joseph L. Mann

Joseph Mann is a PhD student in the Materials Science & Engineering department at Stanford University with a soft spot in his heart for controlled radical polymerization. He is currently funded by a Department of Defense Graduate Fellowship and is interested in the controlled synthesis of topologically diverse polymeric backbones for materials that interface with proteins. More specifically, he is exploring the rational

design of soluble excipients, thermo-responsive artificial chaperones, and injectable supramolecular hydrogels.



Anthony C. Yu

Anthony Yu is a PhD student in the Materials Science & Engineering department at Stanford University and is currently funded by a Kodak Graduate Fellowship. He is interested in developing a fundamental understanding of dynamically cross-linked polymeric materials in order to engineer practical materials solutions to biomedical challenges. In particular, he is exploring the structure–property relations that control

injectability and solute release kinetics of supramolecular hydrogels and how the relations can be leveraged for rational design. Currently, Anthony is designing new injectable materials for controlled drug delivery and sprayable materials for fire retardant encapsulation and retention.

Biomaterials—substances in contact with tissues or biological fluids that are not food or drugs^{1,2}—have evolved from crude wooden prostheses dating back millennia to advanced metals, ceramics, and polymers to augment, repair, or replace diseased and damaged tissue.³ These materials have played a pivotal and transformational role in orthopedic joint replacement, vascular grafting, plastic surgery, dental augmentation and bone fusion,^{4,5} and their usage has been explored in the field of regenerative medicine.^{6–8} While traditional materials make great structural analogues for native bone and tissue, they have trouble recapitulating the dynamics and complexity of the native biological environment.

New biomaterials for novel delivery and therapeutic design must then have marked and intentional differences from what is currently available. Addressing delivery, materials should be easily processed, administered, and able to deliver a wide range of therapeutics with controlled release profiles over various timescales. For addressing novel regenerative therapeutics, we need biomaterials that demonstrate physical and chemical cues in a biologically relevant manner. A significant body of research over recent decades has shed light on the utility of hydrogels for this very matter.^{9–13}

Hydrogels are 3D networks of crosslinked macromolecules that can entrap substantial amounts of water through surface tension and capillary forces. They may be categorized into chemical and physical networks. Chemically crosslinked hydrogels, described as polymeric chains interconnected by permanent non-reversible bonds, have been fabricated through a wide range of chemistries.^{14–20} Chemical crosslinking can be easily tuned to alter the mechanical properties of the final material and has been commonly employed when tough and stable hydrogels are required.²¹ However, due to the covalent nature of the crosslink, these hydrogels are often

brittle, lack transparency, and cannot self-heal after network failure.²² Moreover, the use of metal catalysts, photoinitiators, and UV-light coupled with incomplete conversion of reactive functional groups limits biocompatibility of these systems.²³ Further, a poorly defined structure with network defects and large equilibrium swelling volumes hamper *in vivo* material performance. In contrast, physical hydrogels rely on the formation of transient crosslinks between polymer chains. Because gelation is driven by molecular self-assembly, it avoids the deleterious implications of chemical crosslinks (brittleness, swelling, *etc.*) at the price of mechanically weaker systems. Moreover, physical crosslinks afford tailored viscoelasticity, an important property for biological recapitulation and injection.^{24,25}

Physical crosslinking is made available through the utilization of supramolecular chemistry—chemistry beyond the molecule.^{26–31} These crosslinks generally fall into two broad classes: (1) entanglements between one-dimensional fibrous assemblies of supramolecular monomers²⁷ and (2) supramolecular crosslinks between polymeric precursors.³² The former approach has been famously employed with self-assembling peptide amphiphiles^{33,34} and functionalized amphiphilic benzene-tricarboxamides (BTA).³⁵ Many excellent reviews have been written to describe these low molecular weight gelators (LMWG), so they will not be covered here.^{27,28,36}

Employing supramolecular crosslinks between polymeric chains effectively couples the beneficial properties of polymeric based hydrogels with the diversity and orthogonality of supramolecular chemistry.^{31,32} Typical supramolecular crosslinks employable in aqueous media include host-guest, ionic, metal-ligand, shielded hydrogen bond, and protein based interactions. Similar to chemical crosslinking, altering the density of crosslinks directly affects the mechanical properties



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on the immune response. She uses an injectable hydrogel platform and advanced immunology assays to study the humoral and cell mediated immune responses. Simultaneously, she is also developing novel materials platforms for sustained drug delivery.



Eric A. Appel

Eric Appel is an Assistant Professor of Materials Science & Engineering at Stanford University. He received his BS + MS (2008) in Chemistry and Polymer Science from Cal Poly San Luis Obispo, and PhD (2013) in Chemistry from Cambridge University. For his PhD work he received a Graduate Student Award from the Materials Research Society and a Jon Weaver PhD Prize from Macro Group UK. He was

awarded a Ruth L. Kirschstein National Research Service Award and a Wellcome Trust Fellowship to pursue postdoctoral research with Robert Langer at MIT. In 2016, he moved to Stanford to establish his independent research efforts in the design, synthesis, characterization and application of supramolecular biomaterials for a number of biomedical applications.

of resulting hydrogel. Dissimilar, however, is the transient nature of the crosslink; both the rate and degree of association can be controlled through the judicious selection of the crosslinking moiety. These parameters have direct implications in the mechanical and viscoelastic response of the material.

Due to the modularity of crosslinks, facile gelation mechanism, beneficial mechanical properties, and outstanding ability to replicate biological viscoelasticity, supramolecular polymeric hydrogels are increasingly employed as advanced biomaterials for applications that range from sustained therapeutic delivery to 3D-bioprinting for regenerative medicine.^{31,32} We wrote this review to highlight emerging biological applications of supramolecular polymers and specify why their chemical properties make them great candidates for clinical translation. More directly, this review intends to illustrate the toolbox of supramolecular chemistries for researchers trying to address current biomedical challenges while highlighting emerging biomedical applications for researchers working on developing new supramolecular interactions (Fig. 1). We intend to do this by first creating a design space of supramole-

cular polymeric hydrogels. We will describe the chemical moieties of crosslinks and their respective kinetic and thermodynamic parameters. We will then describe the macroscopic mechanical properties these interactions afford and how they can be controlled. After introducing the design space, we will review how the chemistries and mechanical properties of these materials are leveraged in emerging biomedical applications.

2 Chemistry of the crosslink

The past two decades have seen an explosion of research employing specific, directional, non-covalent moieties, including hydrogen bonding, metal–ligand coordination, host–guest complexation, and ionic interactions. The non-covalent interactions are realized through different binding mechanisms with corresponding associative equilibrium constants (K_{eq}) and associative (k_a) and dissociative (k_d) rate constants. These thermodynamic and kinetic parameters offer insight into the binding mechanisms and provide a handle for the judicious

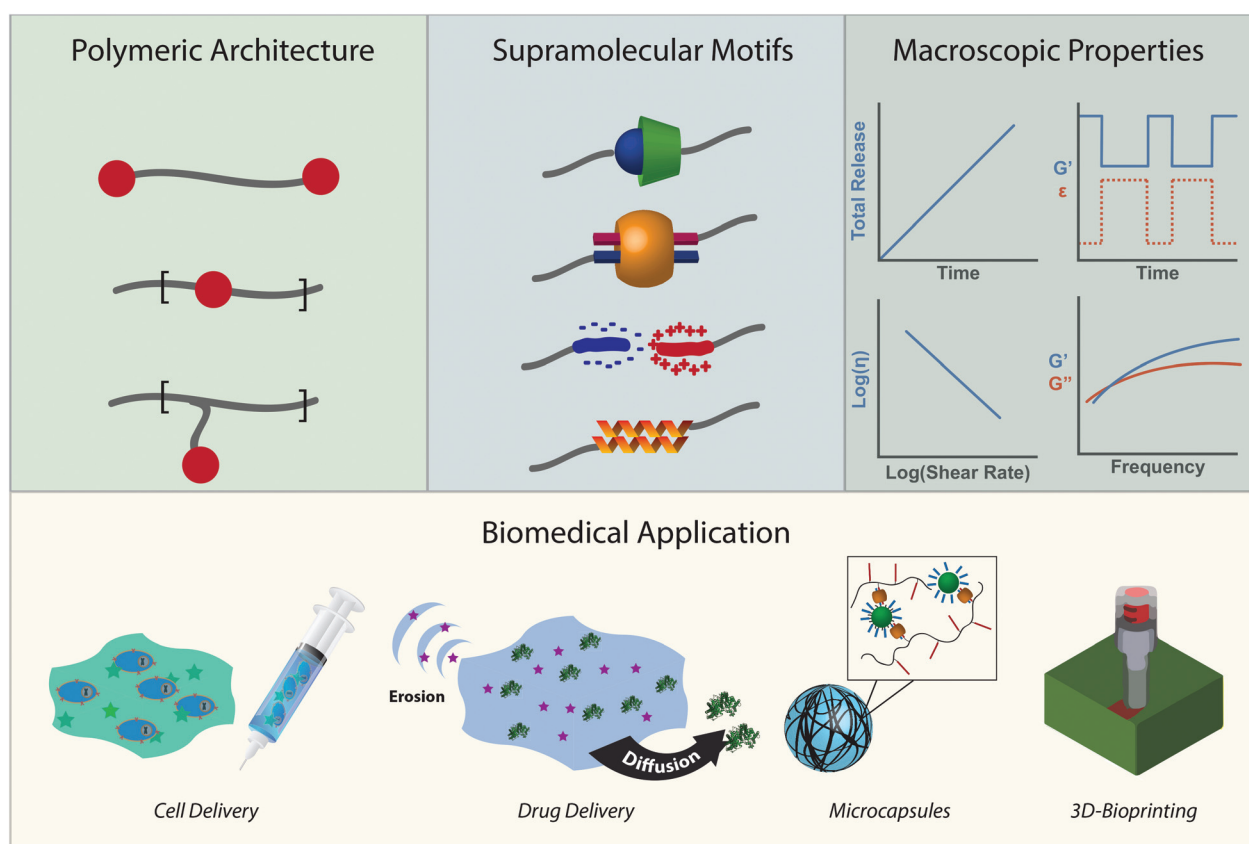


Fig. 1 When designing novel supramolecular polymeric biomaterials, researchers must first choose a polymeric architecture. Illustrated in descending order are a telechelic design, network embedding, and grafting from a linear chain. In this section, the red dot represents the supramolecular motif, chosen for its kinetic and thermodynamic properties. Among the many interactions available, we have illustrated, in a descending list, cyclodextrin, CB[8] ternary, ionic, and protein based coiled-coil interactions. The judicious choice of the architecture and motif affords very specific and unique material properties, such as (left to right) extended release, self-healing, shear-thinning, and frequency-dependent mechanical properties. These considerations make up the toolbox for supramolecular polymeric biomaterial design. Control over the structural architecture and chemical motifs enable fine control over the macroscopic properties, allowing for systematic development of advanced biomaterials. Displayed above are but a few of the applications discussed in this review: cellular injection, cargo delivery on different timescales, microcapsules, and 3D-bioprinting.

choice of a supramolecular system for a biomaterial. Generally, K_{eq} , a thermodynamic constant, will relate the *degree* of association in a system. The kinetic parameters, k_a and k_d , describe the *binding dynamics* of the system. With only these thermodynamic and kinetic parameters, one can begin to envision the degree of connectivity with an understanding of how transient that connectivity is. This connectivity, however, only comprises part of a physical hydrogel's complexity. Beyond the transient nature of the crosslinks, polymer molecular weight, crosslink density, chain rigidity, and polymer concentration must be considered when designing a supramolecular biomaterial. The upcoming section will provide a brief background and discussion on thermodynamic and kinetic parameters, followed by highlights of the currently employed supramolecular interactions and the design handle one has over their thermodynamic and kinetic properties.

2.1 Visualizing thermodynamics and kinetics

When looking at the reaction and free energy diagram of the supramolecular interaction (Fig. 2), we can visualize both the thermodynamic and kinetic parameters. The equilibrium constant is directly related to the relative free energy levels of the unbound and bound supramolecular complex and is described through eqn (1). As the bound complex becomes more energetically favorable, the degree of association increases. Indeed, the degree of association is proportional to $K_{\text{eq}} \times C^{\frac{1}{2}}$.³² Hydrogel formation occurs only when the degree of association

has reached a critical ratio so that a contiguous network forms. Fig. 2 also illustrates that the supramolecular complex spends time as an active (bound) and inactive (unbound) crosslink. The rates of interchange between these states is a direct relation of the forward and reverse activation energy (E_{a} , $E_{\text{a,d}}$). The Arrhenius relationship, shown in eqn (2), describes the temperature dependence of the reaction rate, where A represents the pre-exponential factor and R is the universal gas constant. The associative and dissociative rate constant, respectively, are directly related to the change in free energy between the transition state and the unbound and bound complex. It is not uncommon for the energetic barriers of supramolecular association to approach or fall below values of RT . In these cases, k_a is limited by the rate of transport through the medium, and the process is described as diffusion-limited.

$$\Delta G^\circ = -RT \ln(K_{\text{eq}}) \quad (1)$$

$$k_{\text{a,d}} = A \exp\left(-\frac{E_{\text{a,a,d}}}{RT}\right) \quad (2)$$

These parameters are implicitly tied to the performance and functionality of the biomaterial. An in-depth understanding of their behavior allows for the judicious selection of a supramolecular moiety when trying to design a new system. However it is important to understand that choice of polymeric architecture can drastically effect the process of association between chains. A separate thermodynamic treatment employing statistical mechanics can be used to describe a global energetic minimum for interchain connectivity. When $E_{\text{a,d}}$ is large (such as with certain metal–ligand interactions), associating chains may initially crosslink into kinetically trapped states that do not have enough thermal energy to reach a global energetic minimum. If $E_{\text{a,d}}$ is neither excessively large or small, associating polymeric chains can dissociate and will diffuse through the network, a phenomenon observed in model protein hydrogels.³⁷ This enables the transition from an initial kinetically impeded state of connectivity towards a global energetic minimum and would physically manifest in morphological and rheological changes over time. In the proceeding section we will discuss and review different examples of supramolecular interactions and available polymeric architectures, and begin to relate them back to their fundamental kinetic and thermodynamic parameters.

2.2 The chemical toolbox of the supramolecular crosslink

Supramolecular polymeric hydrogels have been prepared using a wide variety of supramolecular motifs, including host–guest complexation, H-bonding, metal–ligand coordination, electrostatic, and protein based interactions. The interactions most suitable for use in aqueous media are described below. Briefly, more attention in the literature has been given to the thermodynamics and kinetics of macrocyclic host–guest interactions. As such, the chemistry of host–guest interactions will be discussed within the lens of their thermodynamic and kinetic parameters. The following sections on hydrogen bonding,

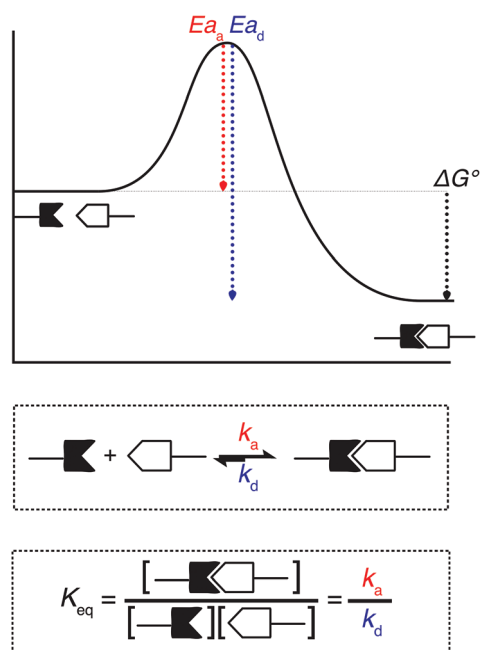


Fig. 2 A visualization of the thermodynamics and kinetics of supramolecular association. (a) Reaction coordinate diagram of a host–guest interaction illustrating the activation energy and free energy of association; factors that are correlated to the values k_a , k_d , and K_{eq} . (b) Equation illustrating the kinetic parameters of association. (c) Equation illustrating the thermodynamic parameters of association.

metal–ligand coordination, electrostatic and protein based interactions will describe trends in polymeric architecture and molecular interaction and how modulation of both parameters affects the process of association.

2.2.1 Macrocyclic host–guest chemistry. The host–guest complexation motif can be visualized as an interaction between a cyclic “host” and an insertable guest that becomes supramolecularly bound inside. Two of the most common motifs for supramolecular polymeric biomaterials are cyclodextrins (CD) and cucurbit[n]urils (CB[n]), although this class of interaction also includes crown ethers, cyclophanes, catenanes, porphyrins, cryptophanes, and carcerands. Both cyclodextrins and cucurbit[n]urils are amenable to aqueous use because the interaction between host and guest occurs in a hydrophobic cavity that minimizes polar solvent interactions. Further, both molecules are nontoxic and demonstrate biocompatibility, affording use in biological applications.

2.2.1.1 Cyclodextrin. Cyclodextrins are cyclic oligosaccharides in which D -glucose units are coupled through α -1,4-glycosidic linkages.³⁸ α -, β -, and γ -CDs, composed respectively of 6, 7, and 8 D -glucose repeat units, are most commonly encountered and generally act as hosts to hydrophobic guests.³⁹ As different CDs contain different sized cavities, they bind different types of guests. Most commonly, α -CD will bind linear alkyl chains while β -CD binds “spherical” diamondoid guests like adamantane and its derivatives.³⁹ Their 3D structure can be represented as a truncated cone with a solubilizing hydrophilic surface composed of hydroxyl groups and a more hydrophobic interior. While hydrophobic and van der Waals interactions between the host and guest in the inner cavity provide the driving force for host–guest interactions, hydroxyl groups on the surface promote self-association mediated by hydrogen bonding.

From a high-level description, these respective interactions can be leveraged to form hydrogels in two ways. The first method takes advantage of the host–guest interaction as a dynamic crosslink; CD and respective guests are grafted onto a polymer backbone (Fig. 3a). In this usage, hydrogels will form in systems where a polymer is grafted with both guest and CD,^{40–42} polymers grafted with either CD or guest are allowed to mix,^{43–47} or linear polymers grafted with a guest are mixed with a CD dimer.^{48,49} The host–guest dynamic crosslink can also be employed in hybrid hydrogels, where a linear polymer functionalized with either CD or a guest is mixed with a nanoparticle⁵⁰ or vesicle⁵¹ that has been functionalized with its complement (Fig. 3b). In a recent study employing a mixture of polymers grafted with either CD or guest, Burdick demonstrates microstructural evolution of a homogenous hydrogel into porous structures.⁴⁷ This is indicative of an initial formation of a kinetically impeded state (homogenous mixture) which subsequently rearranges to a more energetically favorable conformation (porous structure). Implementation of these different strategies, coupled with control over polymer size distribution and topology, may affect the tendency to form kinetically impeded states, however this remains a largely unexplored area of research.

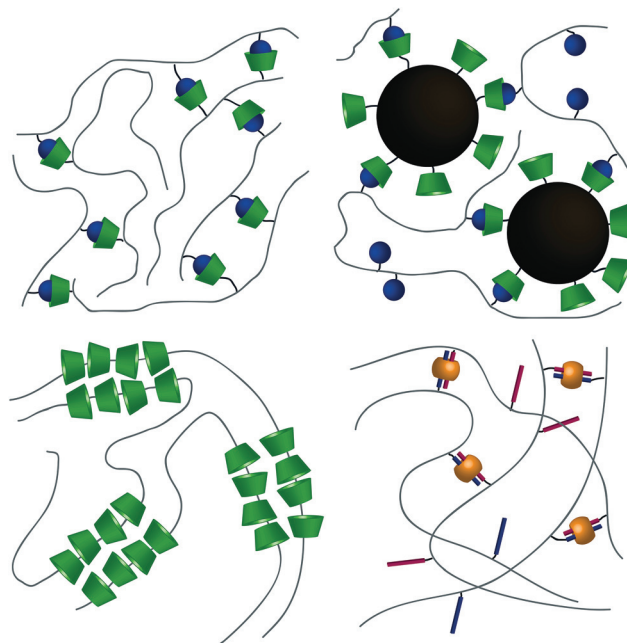


Fig. 3 Schematic illustration of the different ways that hydrogels can be formed through host–guest interactions. (a) A typical host–guest interaction, with hosts grafted to one polymer and guests grafted to another. While CD is depicted here it could just as easily be CB[n] ($n = 6, 7$). (b) A host grafted onto a nanoparticle, with guests grafted onto polymers. (c) CDs that thread polymeric backbones form poly(pseudo)rotoxanes and that have dynamic crosslinks between their surfaces (d) The CB[8] specific ternary interaction where CB[8] can act as a host for two similar or different guests.

The second method takes advantage of CD's ability to thread linear polymers, forming poly(pseudo)rotoxanes.^{52,53} When implemented into poly(pseudo)rotoxanes, CDs have been demonstrated to thread nonfunctional PEG,^{54,55} poly(propylene-glycol),⁵⁶ aliphatic polyesters,^{57,58} poly(ϵ -lysine),⁵⁹ and poly(vinyl alcohol)s⁶⁰ in aqueous solvents (Fig. 3c). While this threading alone does not lead to the formation of hydrogels, the threaded CDs self-associate through hydrogen bonding interactions, forming the physical crosslinks between the linear chains.⁶¹

2.2.1.2 Cucurbit[n]uril. Cucurbit[n]urils (CB[n], $n = 5–8, 19$) are macrocyclic oligomers containing repeat units of glycoluril connected through methylene bridges. CB[n]'s visual structure can be represented by barrels of increasing radius.⁶² CB[n]'s chemical structure is understood as a hydrophobic cavity surrounded by polar carbonyl functional units at either portal. Much like CDs, CB[n]s bind guest through hydrophobic and van der Waals interactions in the hydrophobic cavity. However, unlike CDs, CB[n]s interactions are aided by the release of “high-energy” water molecules trapped in the hydrophobic cavity.^{63,64} Further, CB[n]s have a tendency to bind metal or organic cations through ion–dipole interactions on the carbonyl portal.

Increasing repeat number (n) increases the CB[n]'s portal and cavity size, altering what type of molecule the CB[n] host

tends to bind with. CB[5], the smallest CB[*n*], can encapsulate a variety of gas molecules as well as small solvents, but will not be discussed further as this plays little relevance in the creation of supramolecular biomaterials.⁶⁵ Of greater relevance, CB[6] tends to bind neutral and positively charged organic guests, CB[7] binds larger amphiphilic guests, and CB[8] binds positively charged and large organic guests.⁶⁶ Moreover, CB[8] is unique in the fact that it can bind two guests simultaneously, forming ternary complexes.^{67,68} When guests differ, CB[8] will bind the electron deficient guest before the electron rich guest.

CB[*n*]s have been utilized to form physically crosslinked hydrogels in two ways. Similar to CDs, CB[*n*]s are appended to a polymer while guest molecules are appended to a complementary polymer. Subsequent mixing yields robust supramolecular crosslinks and hydrogel formation (Fig. 3a).^{69–71} Common guests for CB[6] hosts are diamines, such as 1,6-diaminohexane (DAH). In the second system, CB[8] is titrated into a solution of guest molecules grafted to polymers, creating ternary inclusion complex crosslinks with either the same^{72–74} or different guests (Fig. 3d).^{75–79} Common guests include amino acids, viologen (MV, first guest), and 2-naphthoxy (Np, second guest).

2.2.1.3 Kinetics and thermodynamics. Due to the shared host-guest nature of CDs and CBs, discussing their kinetics and thermodynamics in tandem affords insights into their similarities and differences and helps motivate a choice of motif for future applications. One thing that becomes immediately apparent is that CB-guest binding is much stronger than CD-guest binding, often with K_{eq} s that are many orders of magnitude larger. The thermodynamic equilibrium constant (K_{eq}) for CD range between 10^{1-5} M^{-1} with the popular β -CD@Adamantane motif at 10^4 M^{-1} and a α -CD@C₁₂ at 10^3 M^{-1} .³⁹ Equilibrium rate constants between CB[*n*]s of different sizes vary more than constants of different sized CDs. CB[6] binds guests with equilibrium constants as high as 10^{12} M^{-1} ,⁸⁰ but is centered around 10^5 for a variety of alkyl ammonium salts.⁶⁶ CB[7] includes the strongest association interactions and contain equilibrium constants as high as 10^{17} M^{-1} ,⁸¹ but are typically between 10^{5-9} M^{-1} .⁶⁶ CB[8] presents a slightly more convoluted case because there are parameters needed for the first guest, heterodimer, and homodimer inclusion interactions. In the interest of materials properties, we will look at overall equilibrium constants. In the kinetic section, we will explore dissociation constants of the second guest. Typically, both first guest and second guest K_{eq} values fall between 10^{4-6} M^{-1} resulting in overall bond equilibrium constants of $10^{10-13} \text{ M}^{-2}$.⁶⁶

While many thermodynamic parameters have been reported for CB and CD based host-guest systems, quantitative determination of association and dissociation rate constants have proved challenging due to short relaxation times (<1 s) of these supramolecular complexes that demand high time resolution.⁸² Common to both CD and CB (second guest), if the fit is right, k_{a} approaches the diffusion limit (within 1–2 orders of magnitude).^{77,83–88} However, the k_{a} will lower as the

host or guest is attached to a slower diffusing (larger) backbone, when there is a rate limiting shape reorganization between the host, or as recently demonstrated, when the pressure is lowered or the viscosity is increased.⁸⁸ While viscosity and pressure affects CB[8]'s k_{a} they leave k_{d} seemingly unchanged (resulting in K_{eq} alteration). In the same study, they demonstrate that when k_{a} was near the diffusion limited state, only k_{d} (and thus K_{eq}) was affected by different guest molecules, agreeing with previous discussions of CD.⁸⁷ Together, the data argues that the host affects *association*, while the guest controls *dissociation*. It is imperative to remember, however, that association in hydrogel networks happens in a range of viscosities and on large backbones, resulting in equilibrium and rate constants that may vary from those of stop-flow experiments. While CB and CD based systems both share control over their association and dissociation rate constants, when the k_{a} is rate limiting, the k_{d} is inherently tied to K_{eq} . Thus CB moieties that have large K_{eq} s will have larger dissociation constants than CD moieties. For CB[8], however, the relevant k_{d} value is tied to the K_{eq} of the second guest, which is similar or marginally larger than the K_{eq} of CDs.

2.2.2 Hydrogen bonding interactions. Hydrogen bonds are dynamic, directional interactions in which a hydrogen atom bound to an electronegative hydrogen bond donor forms weak interactions with a hydrogen bond acceptor. These supramolecular interactions are of utmost structural importance in biological systems. The three-dimensional assembly of DNA and RNA, the interconnectivity of the collagen triple helix, and the latitudinal extension of β -sheets are but a few of an extensive list of structural features that owe their complexity and function to hydrogen bonding.

A singular hydrogen bond pair renders a relatively weak interaction. However, due to their specificity and directionality, one can greatly increase the association constant through multivalent interactions. Seminal work by Meijer demonstrated that both the ordering of donor (D) and acceptor (A) units (*e.g.* DAAD/ADDA vs. ADAD/DADA) and the number of adjacent hydrogen bonding actors affected the relative association constants between complementary interacting motifs.⁸⁹ Generally, the relative strength can be thought of as an interplay between attractive and repulsive secondary interactions, leaving DAAD/ADDA pairs with higher K_{eq} values than ADAD/DADA pairs.

The equilibrium constant of the hydrogen bonding interaction is strongly solvent dependent.^{89,90} Polar solvent molecules, such as water, compete for binding sites and impede the ability of complementary units to associate. To incorporate hydrogen bonding motifs into biological materials the hydrogen bonds must then be shielded by a hydrophobic spacer. The most common structural design involves poly ethylene glycol (PEG) surrounded by a hydrophobic spacer with telechelic hydrogen bonding units. This design has been used with the ureido-pyrimidinone (UPy),^{91,92} and benzene tricarboxamide (BTA)³⁵ hydrogen bonding motif. In this motif, the supramolecular polymers assemble into long 1D fibers of the hydrogen bonding units (Fig. 4a). The hydrophilic chains can

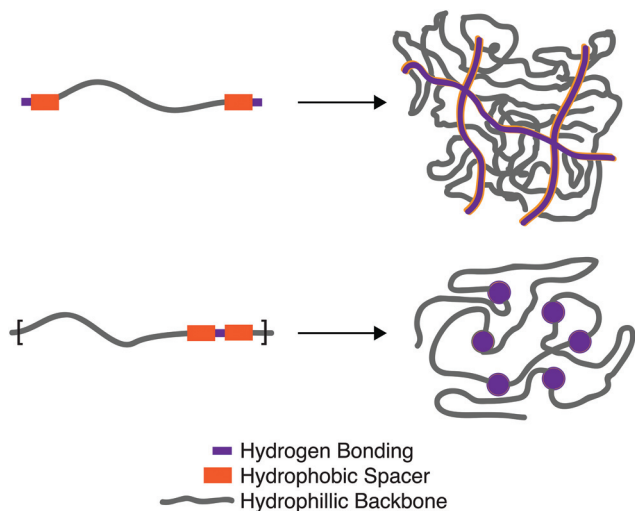


Fig. 4 Schematic illustration of the two most common forms of hydrogels formed through hydrogen bonding interactions. (a) A telechelic water soluble unit forms 1D fibers that bundle and strengthen the hydrogel. (b) Hydrogen bonding units shielded by hydrophobic spacers on a water soluble backbone dimerize to form dynamic crosslinks.

either attach two hydrogen bonding units on the same fiber, or on different fibers, acting as a crosslink between the fibrous units. A different strategy embeds hydrogen bonding units surrounded with hydrophobic spacers directly into the polymeric backbone (Fig. 4b). Instead of creating interconnected 1D fibers, dimeric hydrogen bonding units act as dynamic crosslinks. This strategy has been employed with UPy⁹³ and urea^{94,95} hydrogen bonding units. Instead of embedding the hydrogen bonding unit into the polymeric backbone, Upy groups have also been grafted onto polymeric backbones with a C₆ hydrophobic spacer.^{96–98}

Looking towards the polymeric design space there are three main components: (1) choice of the hydrogen bonding units, (2) choice and length of the hydrogen bonding spacer, (3) length of the hydrophilic chain. Choice of the hydrogen bonding unit will ultimately afford a different K_{eq} , k_a , and k_d , affecting the mechanical properties of the material. The length of the hydrogen spacer will dictate both the degree of crystallinity of the hydrogel and the degree of shielding from protic solvents. The length of the hydrophilic spacer will both describe the flexibility of chains and dictate the ratio of a polymer self-associating (loop) or associating with a different polymeric backbone (crosslink).

2.2.3 Metal–ligand interactions. Metal–ligand interactions are a class of supramolecular interactions in which two or more typically organic ligands donate a non-binding electron pair to empty orbitals in a transition metal cation. This crosslink is a derivative of the rich and well documented field of inorganic complexes. The combinatorial extent of metal–ligand pairs for crosslink applications is both daunting and enticing for the development of biological materials for advanced applications.

Perhaps the most anomalous biological example of the metal–ligand interaction is found on intertidal coastal sur-

faces, where mussels attach themselves through byssal threads terminated in adhesive plaque.⁹⁹ While the feat of underwater adhesion is outstanding by itself, the unique self-healing properties of the byssal threads have directed a field of mussel-inspired metal–ligand hydrogels. Synthetically, this design motif is incorporated into polymeric hydrogels by crosslinking telechelic catechol or histidine units on multi-arm PEG with a variety of transition metal cations, notably Fe³⁺, In³⁺, Al³⁺, V³⁺ (catechol) Zn²⁺, Cu²⁺, Co²⁺, and Ni²⁺ (histidine). Transition metals have different characteristic relaxation times with their respective ligand; judicious mixtures of ligands and metals afford predictable control over timescales of energy dissipation in bulk materials.^{100,101}

Fe³⁺ catechol interactions have stability and strength approaching covalent bonds; measured equilibrium constants approach 10⁴⁰. The mechanical properties of these hydrogel are strongly dependent on the coordination ratio of catechol : Fe³⁺, a function of hydrogel pH. Optimal mechanical properties are realized at pH 12 (coordination ratio of 3 : 1, Fig. 5a), however hydrogel formation is still realized at a pH of 8 (coordination ratio 2 : 1, Fig. 5b).¹⁰² Implementing different transition metals¹⁰³ or catechol motifs¹⁰⁴ afford different coordination ratios at a given pH, leading to more robust and translatable synthetic protocols. Histidine-transition metal hydrogels are often facilitated at neutral pH (coordination ratio 2 : 1). It has been noted that increasing the coordination ratio by lowering the relative concentration of transition metal imparts more rigidity on the network.¹⁰⁵

While strong, mussel-inspired hydrogels are often brittle with little elastic recovery force, an attempt to address this issue involves altering the transition metal bound to a catechol (vanadium and aluminum).¹⁰³ This tends to alter the coordination number at a given pH, affording control over the degree of elastic and viscous character but not necessarily overcoming the flaws of a multi-arm PEG system. Alternatively, Hawker demonstrated that interpenetrating networks could be formed between Fe-catechol crosslinked PEG and loosely crosslinked acrylamide.¹⁰⁶ This dynamic crosslink provides stiffness and energy dissipation at strain rates above the characteristic relaxation times and minimal reinforcement at slow rates. Redesigning the molecular architecture of the catechol crosslink, Ma and coworkers chelated iron with methacrylamide functionalized catechols and polymerized acrylamide, render-

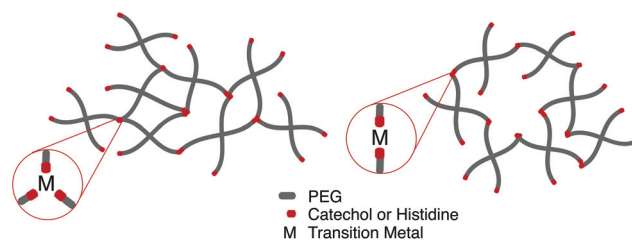


Fig. 5 Depending on the choice of chelating unit (catechol, histidine), transition metal, and pH, hydrogels with coordination ratio (a) 3 : 1 and (b) 2 : 1 will form.

ing an acrylamide network with Fe-catechol crosslinks.¹⁰⁷ These hydrogels are scalable and maintain demonstrable extensibility, however the synthetic process incorporates a non-insignificant amount of DMSO in the hydrogel.

Alternate examples of metal–ligand supramolecular crosslinks include the appendage of ligands to linear polymer backbones with titration of a transition metal crosslink. Polyacrylic acid homo- and copolymers have been used with Fe³⁺ to fabricate hydrogels with high toughness and photo-redox responsive sol–gel transitions.^{108,109}

2.2.4 Electrostatic interactions. Hydrogels that implement electrostatic (ionic) interactions form extremely strong and stimuli-responsive materials. While the underlying nature of these interactions (disordered electrostatic complexes) lack the traditional ordering and predictable stoichiometry of traditional supramolecular motifs, many of the design consideration and resulting material properties are similar to those of traditional supramolecular interactions. As such, a discussion of this chemistry will be included in the review. Several of the most prominent examples fall into three categories: (1) tethered 2D nanosheets, (2) ABA triblock copolymers (where A is charged, B is neutral), and (3) random copolymerization of polyampholytes.

In general, tethered 2D nanosheets form strong, high water content (97%) and elastic hydrogels.^{110,111} Clay nanosheets are implemented and functionalized with a surfactant, impeding aggregation and increasing anionic surface area. These nanosheets are then mixed with PEG containing dendritic,¹¹⁰ or linear¹¹¹ telechelic guanadinium (cationic) groups. The telechelic PEG functions as a supramolecular tether; mechanical properties of the gel are affected by its macromolecular architecture. As the ionic interactions are multivalent, gels become stronger with increasing length of the charge section (or increasing dendrimer generation). Moreover, increasing the length of the PEG spacer (to an extent) increases the probability that the PEG tethers two nanosheets (increasing the modulus) rather than forming a closed loop with itself (lowering the modulus). ABA triblock copolymers represent a strong class of medium water content (85–90%) hydrogels with mechanical responsiveness to salt content, stoichiometry of charge, pK_a of ionic moiety and pH (depending on the ionic A block used). In this polymer architecture, A represents ionic charge (anionic or cationic) while B represents a neutral PEG midblock. This class of material is further separated into two groups: ABA triblock copolymers that are mixed with an oppositely charged homopolymer (C)¹¹² and ABA triblock copolymers that are mixed with an oppositely charged triblock copolymer (CBC).^{113,114} While the ABA–C system form a network of interconnected polyelectrolyte micelles stabilized by a corona of neutral PEG, the ABA–CBC tend to form ordered body center cubic arrays of complex coacervates (polymer-rich regions formed *via* liquid–liquid phase separation) tethered by PEG chains (Fig. 6). ABA–CBC systems tend to be stronger and more ordered than ABA–C systems because they have an increased number of PEG bridges between domains. Both structures tend towards amorphous behavior and display

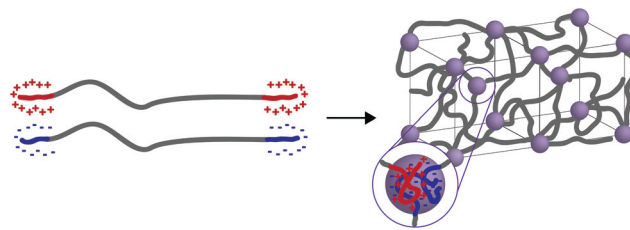


Fig. 6 When ABA and CBC triblock copolymers (A and C have opposite charges) are mixed they form ordered body center cubic (BCC) arrays of complex coacervates tethered by PEG chains.

weaker mechanical properties as either polymer concentration decreases or salt concentration increases.

The polyampholyte motif consists of polymers containing both cationic and anionic monomers and represents a class of very tough, modular, lower water content (50–70%) hydrogels intended for use as structural biomaterials.¹¹⁵ The strength of the electrostatic interaction is proportional to the multivalency of the interaction. By employing a random polymerization, segments of the polymer backbone will contain alternating cationic and anionic domains and repeating cationic (and anionic) domains. The alternating domains will have interactions with lower multivalency, thus acting as weak crosslinks, while the repeating domains will have high multivalency, acting as strong crosslinks. This is akin to utilizing a weak and strong transition metals in metal–ligand interactions; the mixture of strong and weak interactions allows for different and distinct timescales for energy dissipation, imparting strength, self-healing, and toughness into the materials. This hydrogel becomes weaker by employing either a lower concentration, or utilizing more hydrophilic monomers. While not explored in this setting, one could imagine that employing monomers with strategically chosen reactivity ratios could affect the timescales of viscoelasticity in the material.

2.2.5 Protein-based hydrogels. Utilizing proteins and peptides acquired from recombinant or solid-phase techniques requires more complicated and expensive synthetic protocols. However, their implementation as supramolecular crosslinks provides many benefits. Proteins and peptides yield perfect monodispersity and sequence control of the associating amino acids. As subtle changes in amino acid lead to different thermodynamic and kinetic properties, these interactions can be utilized to determine structure–property relationships for relevant mechanical properties, such as erosion. Moreover, protein and peptide interactions are stimuli responsive in biologically relevant settings to biologically relevant molecules, affording temporal release of molecular cargo. These interactions are also found in biologically relevant settings, such as the cellular microenvironment (*vide infra*), allowing for a simpler interface *in vivo*.

Strategies to use proteins and peptides either involve a pure multiblock recombinant protein, or the grafting of proteins and peptides to synthetic backbones. Common strategies include appending a histidine tag (his₆) to the peptide with a

synthetic polymer backbone that chelates Ni^{2+} , thiol-ene chemistry with a cysteine on the protein and a double bond on the backbone, and substitution reactions with lysine residues on the protein and activated carboxylic acids on the polymer backbone.¹¹⁶

The leucine zipper, an example of a recombinant protein, is a helical peptide with a heptad periodicity represented as (abc-defg)_n.¹¹⁷ The a and d residues are generally hydrophobic and frequently leucine, while e and g are charged. The a and d residues land on the same helical face and dimerize with other leucine zippers through hydrophobic interactions. Through recombinant engineering, the leucine zipper motif is incorporated into supramolecular hydrogels through the synthesis of an ABA triblock polypeptide, in which A groups represent the leucine zipper and B groups are a flexible water soluble peptide domain.¹¹⁸ The self-assembly of these triblock polypeptides are illustrated in Fig. 7. Flexibility in *n*,¹¹⁹ the number of heptad repeats, and subtle changes in heptad amino acids¹²⁰ can change the equilibrium number of associating helices (*i.e.* dimer *vs.* trimer *vs.* tetramer) and thermal stability of those associations. Gel stability and strength can be increased by reducing the number of primary loops (*vide infra*), a larger problem with ABA coiled-coil hydrogels. This can be overcome by increasing the B block length¹²¹ or by imagining a novel ABC triblock copolypeptide, where A and C blocks can only, respectively, form coiled-coil interactions with A and C blocks.¹²² A simpler usage of the leucine zipper coil functionalizes a water-soluble polymer with the helical polypeptide, a technique that promotes gelation with >99% water content.¹²³ While these hydrogels tend to be weaker, they demonstrate incredible environmental responsiveness that can be modulated by clever choices of amino acid residues; their mechanical properties are generally affected by pH, temperature, and salt concentration.

The coiled-coil motif has also been incorporated into supramolecular hydrogels through use in multiblock (AB)_n polypeptides, where A is the coiled-coil block and B is a flexible polyelectrolyte block.^{37,124} Like the telechelic leucine zipper, the synthetic design space includes the number of heptad repeat units in A, heptad amino acid choice, and length of the flexible polyelectrolyte block. Introduction of cysteine residues on the N and C terminal of the multiblock polypeptide affords oxi-

dative chain extension and introduces linear entanglements into the hydrogel, affording viscoelastic responses on different timescales.¹²⁴ Appending a thermoresponsive polymer, such as *N*-isopropylacrylamide (NIPAM), to the N and C terminal of the multiblock polypeptide introduces a secondary network reinforcement at temperatures above the lower critical solution temperature (LCST).^{125–127} This technique affords network strengthening after injection into physiologically relevant environments and has also been employed with associating WW and proline rich hydrogels developed by Heilshorn.^{128–130}

The use of proteins as supramolecular crosslinks also allows for environmental response to other proteins or ligands. In a landmark study, Urugami grafted an antigen and a complimentary antibody to separate acrylamide backbones which, upon mixing, association into hydrogels.¹³¹ Grafting the antigen onto the acrylamide backbone results in slight denaturation and a lower K_{eq} (10^8 M^{-1}) and faster k_{d} (10^4 s^{-1}) than the free antigen with the antibody.¹³² Because free antigen binds more competitively than grafted antigen, the hydrogel reversibly swells in the presence of free antigen, releasing molecular cargo. Weber was able to form antibiotic responsive hydrogels by grafting recombinantly engineered bacterial gyrase subunit B (GyrB) to linear polyacrylamide with a histidine tag.¹³³ Coumermycin, an antibiotic ligand, serves as a crosslink between two GyrB units. Addition of a competitively binding antibiotic that only associates with a single GyrB results in hydrogel degradation.

In addition to environmental responsiveness, appropriate use of protein-polysaccharide interactions can better mimic the interactions of proteins and glycosaminoglycans in the extracellular matrix (ECM).¹³⁴ In this regard, the Kiick group has functionalized 4-arm PEG polymers with telechelic low molecular weight heparin (LMWH) or heparin interacting proteins (HIP). Weak hydrogels (97.5% water) form upon mixture of the PEG-LMWH and PEG-HIP, which have equilibrium association constants on the order of 10^6 M^{-1} and dissociation rate constants on the order of 10^3 s^{-1} .^{135,136} Mechanical properties can be controlled through changing concentration of the molar ratio of LMWH:HIP. Weak hydrogels can also be formed through the interactions of 4-arm PEG-LMWH and heparin-binding growth factors (VEGF).¹³⁷ This strategy could be used to employ hydrogels with targeted degradation under the presence of VEGF receptors.

While peptide and protein based hydrogels afford incredible environmental responsiveness, peptide interactions can also be used to demonstrate strain resilient, shear-thinning hydrogels with tunable mechanical moduli. Burdick cleverly utilized the dimerization domain of a protein kinase with the anchoring domain of the kinase anchoring protein to create a “dock-and-lock” hydrogel that formed with simple mixing. The dimerization domain formed the A block of an ABA triblock polypeptide that would self-associate (“dock”). A multi-arm PEG conjugated with the anchoring domain on the ends of the arms subsequently reinforced the dimerization (“lock”).¹³⁸ Through altering the molar ratio of docking and locking groups, the number of PEG arms, and the concentration of

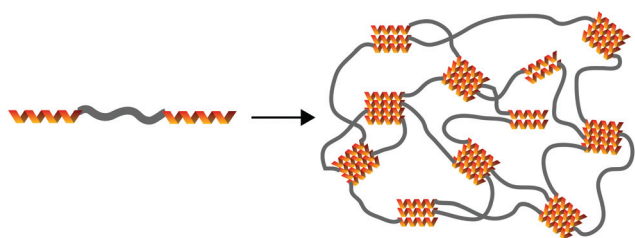


Fig. 7 Demonstration of leucine zipper hydrogels forming from an ABA triblock copolypeptide. Here, the equilibrium number of associating helices is 3, although this can be changed by altering the amino acid of the heptad or the number of heptads in the A block.

polymer, the mechanical properties and degradation rates of the hydrogel could be controlled while demonstrating resistance to yielding at strain rates of 400%.

2.2.6 Polymer–nanoparticle hydrogels. Another promising strategy for creating hydrogels that is not strictly based on supramolecular interactions utilizes polymer–nanoparticle interactions. These hydrogels utilize dynamic and multivalent interactions between polymer chains and nanoparticles to form non-covalent crosslinks that hold the gel network together. While a great advantage of these materials is the ability to form gels simply by mixing the two components, the interfacial chemistries must be matched to form effective crosslinks. However, since these interactions are not strictly directional, these polymer–nanoparticle hydrogels will not be discussed in depth in this review. That being said, it is important to note that these polymer–nanoparticle interactions have been used to make materials with similar properties as those made with aforementioned supramolecular interactions and have potential for drug delivery, surface adhesion, and regenerative medicine applications.^{139–141}

3 Macroscopic mechanical properties

Self-assembly using the preceding non-covalent chemistries is the primary mechanism for crosslinking of hydrogels and has been leveraged to impart dynamic and transient mechanical properties to biomaterials. These properties include the ability to flow under an applied shear strain (shear-thinning) and to reform the mechanical properties of the network once that strain is released (self-healing). Furthermore, they can be finely engineered for a specific application by carefully choosing appropriate supramolecular interacting chemistries with the appropriate interaction thermodynamics (K_{eq}) and kinetics (k_{a} , k_{d}). Naturally, these interactions will also govern the overall strength, stiffness, and elasticity of the material as well as the final microstructure. While each of these considerations provides an additional variable to tune, the increased complexity also allows for many creative materials strategies with precisely designed macroscopic properties. For example, due to the dynamism of supramolecular interactions, hydrogels made with supramolecular crosslinks typically will not swell when submerged in water—the stresses are dissipated through rearrangement of the non-covalent network. In the following sections, we attempt to highlight how these chemistries have been used to control gel strength, shear-thinning, self-healing, and solute release kinetics.

3.1 Viscoelastic materials

Supramolecular biomaterials are often designed to mimic the mechanical properties of biological materials. From a structural standpoint, biological materials are commonly formed from self-assembling, non-covalent interactions between small molecules or polymers. These biomaterials exhibit viscous and elastic behaviors and are therefore classified as viscoelastic materials. Due to these characteristics, engineered biomater-

ials seeking to mimic biological materials must match the elastic behavior as well as the viscous behavior of the material to impart time-dependent mechanical properties.¹⁴²

While there are many useful in-depth discussions on viscoelasticity,^{142–145} the development of models that correlate bulk rheological properties to supramolecular structure remains a complex challenge and active area of research. The following discussion presents the basic rheological grammar commonly used in characterizing and designing supramolecular biomaterials.

Oscillatory rheometry is a powerful way to probe the molecular origins of a material's viscoelasticity and also the primary way of quantifying the elastic and viscous properties. In oscillatory rheometry, a sinusoidal shear strain is applied to the material, while the corresponding stresses are monitored. An ideal elastic material will have a stress response exactly in-phase with the applied strain ($\delta = 0$), while an ideal viscous material will be exactly out of phase ($\delta = \frac{\pi}{2}$). For viscoelastic materials, the response lays somewhere in between ($0 < \delta < \frac{\pi}{2}$) and researchers typically report the storage modulus (G'), the loss modulus (G''), and the ratio between G'' and G' ($\tan(\delta)$).¹⁴⁴

While mathematically G' and G'' represent moduli multiplied by the in-phase or out-of-phase factor of the stress–strain response, they can be loosely thought of as the amount of stored energy (G') and the amount of dissipated energy (G'') by the material. $\tan(\delta)$ represents the ratio between G'' and G' and is often treated as the relative elasticity of the material. As a result, G' and $\tan(\delta)$ are commonly reported together in order to summarize the material elasticity, while providing information on how fluid-like the material is. These values are obtained through common rheological measurements like frequency sweeps (Fig. 8b and d), or strain sweeps. Frequency sweeps are one of the most common measurements reported for characterizing viscoelastic materials and Fig. 8d illustrates an example plot labelled with a crossover point and a plateau modulus. Notably, the viscoelastic moduli exhibit frequency dependent values that are related to the transient crosslinking interactions. In particular, the crossover point indicates the timescale of viscoelastic network relaxation while the plateau modulus represents a pseudo-equilibrium of the crosslinked network. Both of these parameters are directly influenced by the network's degree of crosslinking and the K_{eq} of the crosslink interactions. As the degree of crosslinking is increased, the crossover point will shift towards lower frequencies, until there is no visible crossover point in the frequency range tested. A widely used method of qualifying a material as a gel is to demonstrate no crossover point in a standard frequency range (0.1–100 rad s⁻¹). When the K_{eq} of the crosslinks approach covalent-like values, the plateau modulus will increase and extend towards lower frequencies, approaching a frequency sweep profile of an elastic material (Fig. 8b). A rich understanding of how microstructures are represented by these oscillatory measurements is invaluable when characterizing and engineering new viscoelastic materials.

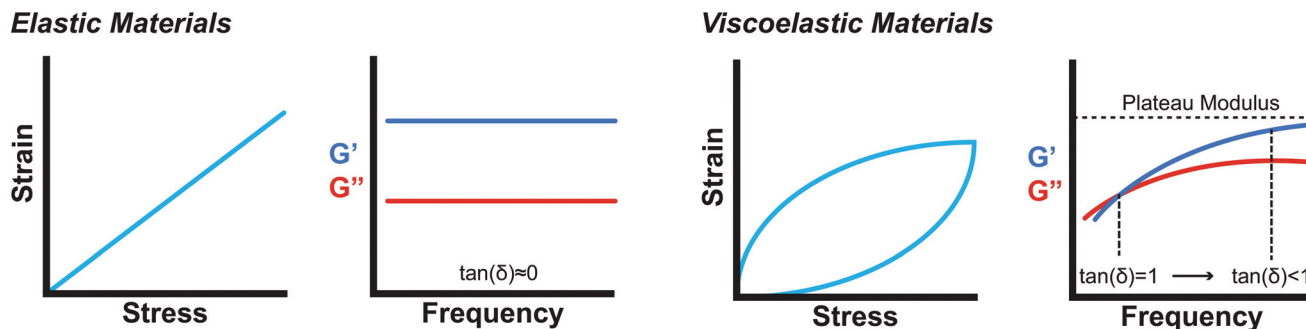


Fig. 8 (a., c) Comparison of ideally elastic materials with viscoelastic materials. While elastic materials have a linear strain response to stress, viscoelastic materials exhibit a hysteresis loop, where energy is being lost during the cycle. (b) and (d) represent typical log–log plots of frequency sweeps for elastic materials and viscoelastic materials over a standard experimental range (0.1–100 rad s⁻¹), respectively. Viscoelastic materials exhibit a frequency dependence on the modulus and a crossover point where $\tan(\delta) = 1$ that represents the timescale for network relaxation.

While oscillatory rheometry is capable of examining a wide range of moduli and time scales typically desired for biomaterials, the development of microrheology has enabled local probing measurements on heterogeneous samples, smaller sample volumes over significantly larger frequency ranges, and the possibility of high throughput screening.¹⁴⁶ These methods include atomic force microscopy,¹⁴⁷ optical interferometry,¹⁴³ video-particle tracking,^{148–151} and magnetic twisting cytometry.¹⁵² However, current microrheological techniques also suffer from several challenges, including the limitation that the materials must be partially transparent and a significant amount of computational analysis must be done to extract data. Furthermore, while microrheology excels at analyzing low viscosity samples, it has trouble with very stiff materials that limit movement of probe particles.

Using these characterization techniques, many different strategies have been employed to develop supramolecular materials with robust elastic properties. Poly(pseudo)rotaxane gels formed with α -CD and PEG chains achieved a G' of 500 Pa, $\tan(\delta) = 0.02$ with linear viscoelastic responses to strains upwards of 300%.¹⁵³ These gels utilized host–guest interactions between the α -CD and bis(2,4-dinitrophenyl)-capped PEG chains to form topological restrictions that acted as the crosslinkers. Additionally, double network hydrogels composed of interpenetrating polymer networks achieve high moduli through the combined elastic contributions from both networks. Commonly, the two networks exhibit drastically different mechanical properties, where one network might act as a sacrificial dynamic network to dissipate stress and increase stretchability of the material. Ping Gong and co-workers used bacterial cellulose (BC) and gelatin to form a double network gel with a tensile modulus of 2.9 MPa. The BC provided a hydrogen bonding network between cellulose fibers, while the gelatin provided a strong covalently bonded network. The supramolecular BC network itself showed weak mechanical properties, while the gelatin on its own was brittle. The formation of a double network allowed the final material to be strong, while maintaining stretchability.¹⁵⁴

Supramolecular motifs also allow for independent modulation of network strength and network stretchability. Alginate

hydrogels can be crosslinked by covalent and ionic interactions. Mooney demonstrated that the incorporation of ionic crosslinking allows for independent tuning of hydrogel modulus and toughness (energy stored before fracture). While increasing covalent crosslink density increased the modulus, it led to brittle, non-stretchable gels. Yet, by increasing ionic crosslink density, the resultant gels had higher moduli and increased stretchability.¹⁵⁵

Whereas consideration of interaction thermodynamics provides intuition on how strong each crosslink is in the network, the examples above utilized the dynamic property of the bonds to impart additional mechanical properties. This concept will be explored further in discussions about shear-thinning and self-healing. Moreover, beyond the chemistry of the non-covalent interactions, the examples illustrate how creatively combining the interactions can significantly influence the final modulus and relative elasticity.

3.2 Self-healing and shear-thinning

Injectable therapeutics and drug delivery strategies rely on biomaterials that can be easily pushed through a syringe or catheter. To allow for injection, these materials are designed to decrease in viscosity under an applied shear force (shear-thinning). For drug delivery materials, they must also rapidly reform their initial structure with minimal changes to mechanical properties when that stress is released (self-healing) in order to reduce burst release of cargo. Materials formed with supramolecular interactions offer both shear-thinning and self-healing properties by nature of the dynamic non-covalent bonds that act as the primary driving force for self-assembly. As a result, many developing biomaterials utilize supramolecular interactions to maintain injectability.

In the majority of cases, shear-thinning behavior has been fundamentally characterized as either a shear-induced splitting of non-covalent interactions, or a localized region of high strain rate affecting flow properties.¹⁵⁶ The degree to which a material shear-thins and self-heals is a direct consequence of the polymer architecture and bond association and dissociation kinetics. Consequently, a comprehensive understanding of the supramolecular chemistry is essential towards

imbuing materials with these properties. It should be noted that covalently crosslinked materials can also exhibit shear-thinning properties through dynamic covalent bonds that are thermodynamically controlled rather than the traditional kinetically controlled bonds.¹⁵⁷ Yet, in covalently crosslinked materials, as well as most non-covalently crosslinked materials, the viscosity differential upon shearing is limited to 1–3 orders of magnitude. In both covalent and non-covalently crosslinked materials, controlling these properties requires understanding the chemistry and kinetics of the intramolecular bonds, as well as the mechanism of bond dissociation and association. The following discussions will focus on supramolecular materials.

While there is no single unifying parameter than can be used to compare between shear-thinning abilities across different mechanisms, chemistries, and materials platforms, the relaxation time of the network is often used as a point of reference. This relaxation time encompasses the dynamics of network rearrangement and can provide insight into the material's mechanical behavior regardless of the specific chemistry used. Craig and coworkers expanded on this idea and identified an untangled, a semi-dilute, and an entangled regime that all have competing relaxation mechanisms and lifetimes that determine the final shear-thinning abilities.^{158,159} Furthermore, it has been demonstrated that materials can shear-thin or shear-thicken depending on the kinetics of bond dissociation. Faster bond dissociation was attributed to the supramolecular bonds rapidly associating and dissociating, which led to network disentanglement. Slower bond dissociation led to intrachain crosslinks being converted into interchain crosslinks upon shearing, resulting in an increase in entanglement.⁷⁷

Despite the lack of a unifying parameter, the rheological experiments used to measure self-healing and shear-thinning are well defined. Typically, demonstration of self-healing is done using either a step-strain experiment, where high and low strains are alternately applied and the moduli are measured in oscillatory mode, or a step-shear experiment, where a high and low shear rate are alternately applied and the viscosity is measured in flow mode. These experiments probe the ability for the material to recover its original structure after network scission and characterize how quickly the network heals. While step-strain experiments probe the elastic network by measuring the shear moduli, step-shear experiments monitor the shear viscosity of the bulk material. Although both experiments can characterize the time-scale of self-healing, a step-strain experiment is insufficient when characterizing a material for injectability, a process that is fundamentally flow-based. Moreover, data points are often “lost” between the strain transitions due to the time restraints of oscillatory mode (data must be averaged over several cycles) and it is harder to relate modulus and shear strain to a material passing through a needle. On the other hand, step-shear experiments measure the viscosity based on a varying shear rate—native parameters when considering injectability. Therefore, step-shear experiments are better for characterizing injectability, while step-strain experi-

ments provide information on the network structure and elasticity under high shear strains.

Shear-thinning is fully characterized using a steady-shear experiment, where a constantly increasing flow rate is applied at a fixed rate and the viscosity is measured. In simple terms, the viscosity (η) can often be experimentally related to the applied shear rate ($\dot{\gamma}$) through a power law relationship (3).

$$\eta = k\dot{\gamma}^{-a} \quad (3)$$

This relationship provides insight into the degree of shear-thinning through the power law index (a). A value of $a = 0$ represents a Newtonian fluid, such as water or honey, that does not shear-thin. As the value of a increases, the degree of shear-thinning increases. This feature is particularly important when considering the stress required to inject a supramolecular hydrogel. Fig. 9a illustrates the calculated pressure (P) required to push 1 mL of material through a 1 mL syringe in 10 seconds. As a general trend, more pressure is required to push material out of a higher gauge needle (lower needle diameter).

The magnitude of this relationship ($\frac{dP}{d(\text{gauge})}$) is dependent on the power law index, where materials with higher power law indices exhibit less dependence on needle gauge (Fig. 9b). This lack of dependence is desirable as higher gauge needles afford less invasive procedures. Newtonian materials like honey may be injected through lower gauge needles, but becomes increasingly more difficult to push through higher gauge needles as indicated by the dotted line (the maximum achievable pressure applied to a 1 mL syringe by 98% of subjects from ref. 164). In contrast, MV/Np@CB[8] and PNP hydrogels exhibit significant shear-thinning, with minimal pressure dependency on needle gauge (Fig. 9a). Notably, for a 31 gauge needle typically used to inject insulin, MV/Np@CB[8] hydrogels approach the injectability of blood, while exhibiting moduli ranging up to several hundreds of pascals under working conditions.⁷⁸ It is important to point out that for injectable applications, both step-shear and steady-shear experiments are crucial to demonstrate appropriate injectability properties. In a steady-shear experiment, low shear rate data provides insight into the material viscosity *in operando* and high shear rate data characterizes injection viscosity (Fig. 9d). The corresponding log–log plot is used to determine the power law index, allowing for comparison of shear-thinning ability between materials. Studies of shear-thinning materials regularly only perform a step-strain experiment, which lacks information about how easily the material would actually flow through a needle and diminishes the ability to truly judge or engineer injectability.

The physical chemistries behind disentanglement and dissociation processes have been extensively studied and exploited through using various bonding chemistries and bonding motifs.¹⁶⁵ For supramolecular biomaterials, shear-thinning materials have been loosely categorized under three groups: (1) protein-based hydrogels, (2) hydrogel nanocomposites, and (3) host–guest based hydrogels.¹⁶⁶

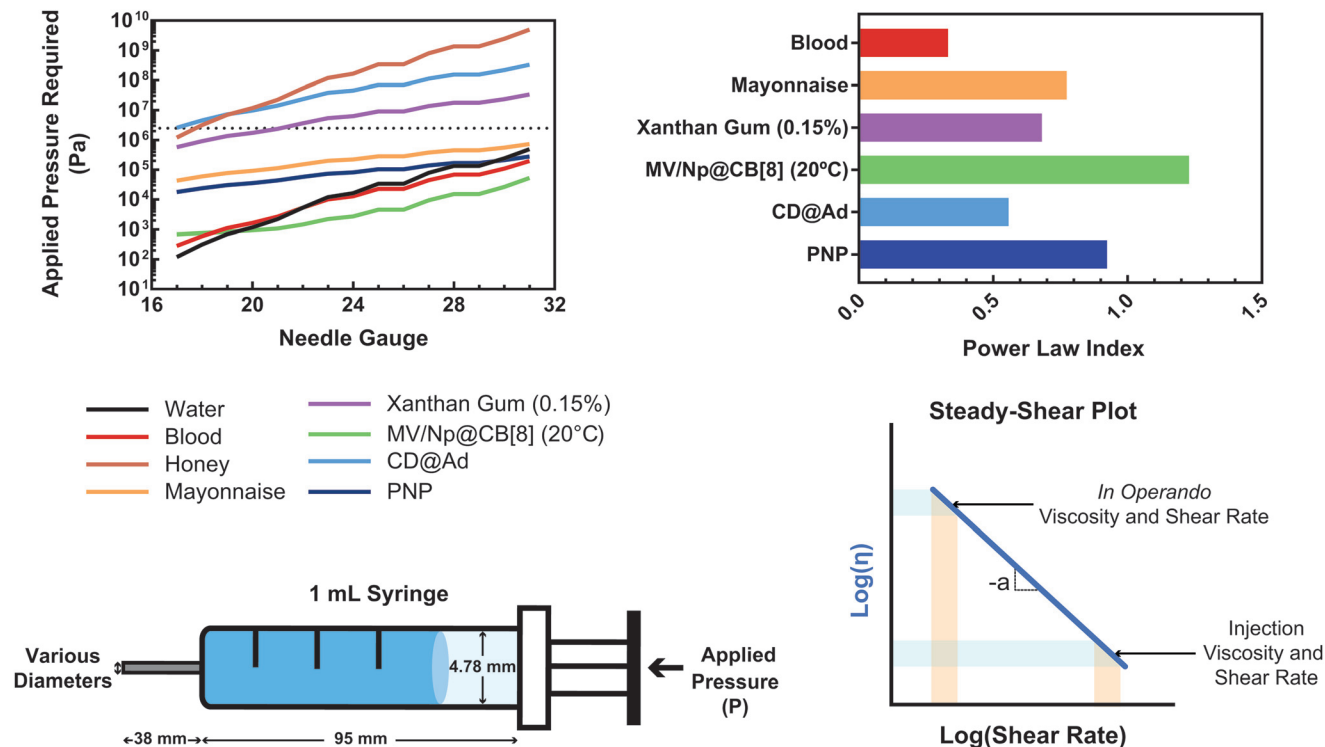


Fig. 9 (a) The calculated pressure required to push 1 mL of material [water, blood,¹⁶⁰ honey, mayonnaise,¹⁶¹ xanthan gum (0.15%),¹⁶² HEC-Np + PVA@CB[8] (0.1%),⁷⁸ Ad-MeHA/CD-MeHA (10%),¹⁶³ and HPMC-C₁₂/PEG-PLA PNPs¹⁵⁹] through a 1 mL syringe over 10 seconds. The dotted line represents the average pressure subjects were able to apply to a 1 mL syringe from ref. 164. Plateaus in the plot result from gauges that have the same inner diameter. (b) A plot of the power law index (a) of the shear-thinning materials pushed through the syringe. All data were obtained at 25 °C unless otherwise stated. (c) Calculations were done assuming a horizontal needle exhibiting lamellar flow and pressure loss from the syringe and needle walls. (d) Example of a steady-shear plot, where viscosity is plotted vs. shear rate in a log–log graph. Viscosities at low shear rates are indicative of *in operando* viscosities, while high shear rate viscosities exemplify injection conditions. The power law index values were calculated from the referenced data from viscosity points around 10 s⁻¹.

Protein–protein interactions used to develop shear-thinning and self-healing hydrogels leverage hydrogen bonding between specific β -sheet structures to provide the non-covalent, dynamic interactions. Specifically, intracellular WW domains bind with Pro-Pro-X-Tyr motifs with K_{eq} values as high as 10⁶ M⁻¹. These components self-assemble to crosslink the network ($G' \sim 50$ Pa) and self-heal within 5–30 minutes depending on the strength of the interacting peptide pairs.^{128,167} As mentioned before, leucine zipper motifs have been utilized for making shear-thinning hydrogels. In these motifs, the bundle formation of helical structures can be disrupted at high or low pH, which allows for stimulus controlled shear-thinning and self-healing of the structure.¹⁶⁸

Hydrogels that form through interactions with nanoparticles or nanodisks provide another set of shear-thinning hydrogels, though these interactions are not strictly supramolecular due to the lack of directionality. In these materials, the primary network structure is usually adsorbed onto the surface of the nanoparticle or nanodisk through hydrophobic interactions, hydrogen bonding, and ionic interactions. PEG based gels have been created through physically crosslinking the PEG chains with silicate nanodisk surfaces. It is imperative in these structures that each polymer chain interconnects several nano-

disks or nanoparticles. Schmidt and coworkers demonstrate this requirement by combining LAPONITE® clay nanodisks with PEG chains of different molecular weight. Using small angle neutron scattering (SANS) and rheology, they were able to show that the onset of shear-thinning decreases as the PEG chain molecular weight is decreased, since the chains lose their ability to effectively interconnect multiple disks.¹⁶⁹ While clay nanodisk based nanocomposites utilize ionic interactions, purely hydrophobic interactions can also be used to create shear-thinning, self-healing gels. Diblock copolymers of PEG-*b*-poly(lactic acid) (PEG-PLA) can be nanoprecipitated to form polymer nanoparticles and act as crosslinks in a cellulosic network. Previously, Langer and Appel showed that hydroxypropylmethyl cellulose (HPMC) can interact with PEG-PLA polymer nanoparticles to form hydrogels. To strengthen this interaction while maintaining injectability, pendant alkyl groups could be added to the HPMC backbone.¹³⁹

Host–guest interactions also provide facile methods to tune the shear-thinning and self-healing abilities of hydrogels. Burdick and coworkers have extensively studied how to leverage host–guest interactions between β -CD and adamantane to form injectable hydrogels. In particular, they designed hyaluronic acid hydrogels with β -CD and adamantane pendant groups

to drive supramolecular self-assembly of the hydrogel. The combination of HA with CD was chosen due to both being abundantly used in biomedical applications and the modularity of the interaction. By changing the concentration of β -CD and adamantane modified HA, the authors were able to tune the shear-thinning and self-healing properties as well as overall gel modulus.^{43,170,171} These CD based polymeric materials have continued to be explored for 3-D printing of biomaterials due to their injectability.¹⁶³ Similarly, CB based complexes have also been used as supramolecular crosslinks. In 2010 Scherman reported the first example of a hydrogel based on CB[8] host-guest inclusion complexes. These host-guest interactions involved two hosts, forming a ternary complex that had independent kinetics for each host. The pore size, mechanical properties, and thermal reversibility of these materials as well as their shear-thinning, self-healing properties could be modulated through tuning the concentration of CB[8] present.⁷⁷

3.3 Cargo release kinetics

Supramolecular polymeric biomaterials have increasingly been developed for drug delivery applications. One of the main advantages of using supramolecular hydrogels for release is the ability to finely control the kinetics of cargo release. When designing a new material for a specific desired kinetic profile, it is important to consider the drug-network interactions and the overall network stability over time.

3.3.1 Cargo diffusion. Solute release from a polymeric network is highly dependent on mesh size and solute-network interactions. Typical mesh sizes range from 5 to 100 nm (ref. 172) and direct characterization techniques of the mesh size include small-angle X-ray scattering (SAXS), small-angle neutron scattering (SANS), and surface adsorption analysis. If the mesh size of the network is much larger than the size of the solute, then the kinetics are diffusion dominated. In this case, the diffusivity, D , of the diffusing species can be modeled by the Stokes-Einstein equation $D = \frac{k_b T}{6\pi\eta r}$.¹⁷³ Once the mesh size is close to the solute size, solute diffusion and determination of the diffusivity becomes more complex. If the mesh size is smaller than the cargo, no diffusion occurs. However, supramolecular networks are composed of dynamic crosslinks that serve as molecular gates for the meshes (Fig. 10a). The rate at which this gate opens and closes is dependent on the k_a and k_d of the supramolecular interaction (a function of $E_{a,d}$; eqn (1)). This is to say that a material with supramolecular crosslinks, where k_a and k_d can be controlled independently of K_{eq} , can be engineered to demonstrate different release profiles (Fig. 10b).¹⁷⁴

Scherman exploited this phenomena using CB[8] based hydrogels with different second guest moieties. These second guest moieties displayed a range of k_a and k_d values, while exhibiting identical K_{eq} values. Interestingly, this preparation leads to a set of gels that have the same plateau G' due to the identical crosslink K_{eq} , but different release kinetics depend-

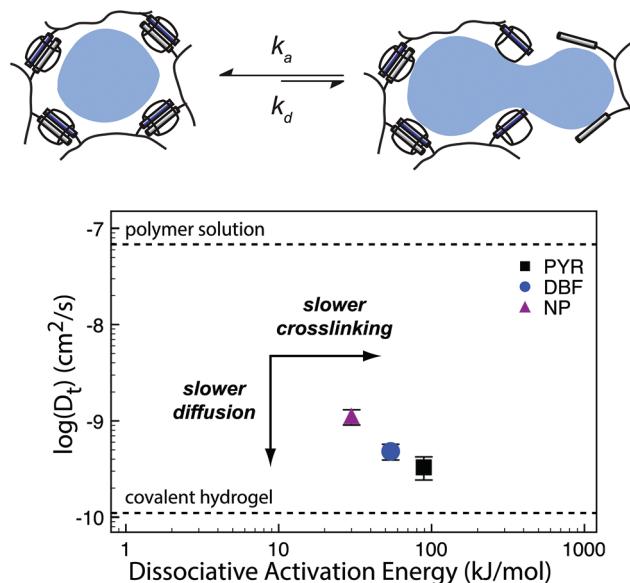


Fig. 10 (a) Supramolecular hydrogels that use non-covalent crosslinks, such as the CB[8] ternary complexes shown above, exhibit time-dependent mesh sizes that depend on the kinetics of the crosslink interactions. These hydrogels can exhibit identical crosslink thermodynamics, but varying kinetics due to the different guest moieties. (b) A supramolecular interaction can have different dissociation kinetics, but similar binding thermodynamics. Using the example of hydrogels prepared with CB[8] ternary complex crosslinks with data adapted from ref. 175 we demonstrate that different diffusion profiles for the same cargo are achieved for materials of similar moduli due to varying guest moieties (PYR = pyrone (slow); DBF = dibenzofuran (medium); Np = naphthalene (fast)). Moreover, scattering experiments illustrated that these gels were isostructural.

ing on the rate of the crosslink kinetics (k_a and k_d). This phenomenon is due to faster kinetics resulting in larger effective mesh sizes and therefore faster release of cargo.¹⁷⁵ These results illustrate an important distinction between interaction strength and interaction kinetics, and how they must be independently considered when engineering complex properties like cargo diffusion. While reports of G' and $\tan(\delta)$ can provide insight into the overall elasticity of a material, it is not sufficient to draw conclusions about release kinetics solely from those values. These complex relations have not been explored thoroughly and further research would offer more insight into engineering solute diffusion in a number of important areas.

A different strategy to tune the diffusion of the solutes involves deliberately engineering solute-network interactions. This method provides an additional interaction to impede solute diffusion kinetics and can involve ionic interactions, hydrogen bonding, and hydrophobic interactions, to name a few. Guldberg and coworkers utilized negatively charged alginate hydrogels to delivery cationic, heparin binding growth factors. The cargo release was primarily controlled by hydrogel degradation and the protein-network interaction provided additional stability to the protein encapsulated in the gel.^{176,177} Using a CD/Ad host-guest based HA hydrogel,

Burdick and coworkers demonstrated control over solute release kinetics by modifying the CD:Ad pendant group stoichiometry.¹⁷⁸ While CD typically acts as a host for Ad, when CD is in excess it will be available to reversibly trap solute molecules in its hydrophobic cavity as well. Tuning the CD:Ad ratio as well as the solute affinity to CD allowed for fine control over release kinetics.

3.3.2 Network erosion and dissolution. Beyond tuning the diffusivity of the solute in the network, it is also important to consider the kinetics of material degradation. Typically, release from a supramolecular hydrogel occurs through a surface erosion or dissolution mechanism (Fig. 11). Surface erosion or dissolution occurs when loss of network structure is rate limited at the surface of the material. This loss of structure can occur through the scission of chemical bonds (surface erosion), or through the solvation of polymer chains attached to the network by supramolecular interactions (dissolution). Surface degradation is relatively slow and linear with respect to time, leading to a near zero-order release profile and is desirable for long-term delivery strategies. On the other hand, bulk erosion occurs when erosion takes place at the surface and in the bulk simultaneously. However, in the case of supramolecular hydrogels, bulk erosion occurs at such a negligible rate that for the purposes of cargo delivery it will not be discussed here.

Surface based degradation is often desired in supramolecular polymeric material systems and is a direct result of the K_{eq} and kinetic rate constants, as well as the polymer molecular weight, degree of connectivity, and open surface area. In the β -CD-Ad-HA gel system described previously, the hydrogels exhibited an initial burst release of bovine serum albumin (BSA) due to residual BSA on the surface, followed by a nearly linear release rate. These release rates were tunable by varying the initial amount of BSA. The combination of a shear-thinning, self-healing material with surface dominated erosion kinetics makes these gels excellent candidates for extended drug delivery (*vide infra*).⁴³ Previously discussed cyclodextrin poly(pseudo)rotaxane motifs have also been shown to form hydrogels with surface dominated release kinetics. In these studies, the authors loaded PEG- α -CD poly(pseudo)rotaxane

hydrogels with BSA and lysozyme and measured kinetics of release while keeping track of the exposed surface area. The final results were best fit to zero order release kinetics, insinuating surface dominated kinetics.⁵⁵

4 Biomaterial applications

Given the toolbox of chemical binding moieties and design criteria for macroscopic mechanical properties, one can begin to engineer new biomaterials for several biomedical applications. In the following sections, we will describe how researchers have used these tools to design biomaterials for drug delivery, immunology, regenerative medicine, and 3D-bioprinting.

4.1 Drug delivery

Drug delivery using polymers began in the 1960s, when researchers began to explore biocompatible materials to manipulate the pharmacokinetics and pharmacodynamics of a delivered drug.¹⁷⁹ Since then, the field has progressed in producing a variety of strategies for “smart” delivery, where the materials allow the drug to overcome certain biological barriers for delivery. These advances were largely motivated by improving efficacy of the delivered drugs, requiring a lower dosage and therefore cost of therapy, and increasing patient compliance. Currently, there is a focus on creating advanced delivery systems, which involve additional functionality to delivery systems. These functionalities include extended cargo release and stimuli responsiveness, which both fall under the category of controlled drug delivery.

Supramolecular polymeric biomaterials provide significant advantages towards realizing advanced delivery systems. The main criteria required for effective controlled release systems include localization of the payload, stable therapeutic release of cargo, no toxicity arising from the delivery system, stability of the encapsulated drug, amenability to stimuli responsive release, and a non-harmful fate for the material at end-of-use. In light of these criteria, supramolecular hydrogels and microcapsules utilizing supramolecular motifs are excellent candidates as a general platforms for engineering delivery materials.

When applying supramolecular materials for drug delivery, localization of the payload is contingent on how the material is delivered to the site of interest. This most often translates to designing hydrogels that are injectable, or using microcapsules. Injectability allows the material to be administered relatively easily and implies fast shear-thinning and self-healing kinetics in the case of hydrogels. Fast self-healing kinetics reduce burst release, allowing more of the payload to be released at a kinetic rate governed by the engineered microstructure and transient behavior of the material. Stable therapeutic release of cargo is related to the overall release kinetics and whether the kinetic profile is consistently at therapeutic levels. Toxicity can be reduced by choosing components of the material that are highly biocompatible, such as those “generally recognized as safe” by the Food and Drug Administration (GRAS-listed). Due to the modularity of supramolecular inter-

Surface Erosion or Dissolution

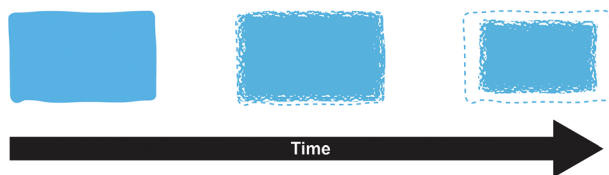


Fig. 11 Surface degradation (erosion or dissolution) occurs when chemical bonds or supramolecular interactions at the surface are broken. Surface erosion occurs when chemical bonds are broken, usually through a hydrolysis reaction, while surface dissolution occurs when non-covalently crosslinked polymer chains detach from the surface. Surface degradation kinetics are directly proportional to the surface area and arise when the rate of degradation at the surface exceeds bulk erosion.

actions, there is a significant amount of flexibility in choosing components that can result in the same final macroscopic properties, while maintaining minimal toxicity. Furthermore, as demonstrated in a previous section, the modulus and dynamic properties can be tuned through increasing the amount of supramolecular interactions, or devising ways to tailor the thermodynamics/kinetics of these interactions (*vide supra*). This flexibility allows for materials to have moduli and viscoelastic properties most compatible with the surrounding biological interface. To address drug stability in the material, strategies have utilized non-covalent drug-network interactions that can protect the drug from being exposed to enzymes and maintain a specific conformation. Stimulated release is also a property that is amenable to supramolecular polymeric systems that use pH, temperature, ionic, or other external stimuli to change the release behavior (*vide supra*).

4.1.1 Hydrogels. Polymeric hydrogels are the most widely utilized systems used for drug delivery due to their high water content, innate biocompatibility, tunable mechanical properties, and ability to protect cargo from enzymes and proteins.^{180,181} However, traditional hydrogels made with covalent crosslinks require surgical implantation and do not reliably provide long-term delivery beyond a couple days. Consequently, the materials solution to these challenges is to use physically crosslinked hydrogels, where the crosslinks are dynamic. These gels are imbued with useful functional properties, including the ability to be injected through syringes, or catheters and to immediately reform to provide a minimally invasive implantation method. Below, we will discuss several notable strategies in designing supramolecular hydrogels for drug release.

Host-guest interactions provide a facile way of independently controlling thermodynamics and kinetics of the interactions. As described above, Burdick has developed a β -CD-Ad-HA system that forms shear-thinning, self-healing hydrogels with tunable strength. In particular, they designed hyaluronic acid hydrogels with β -CD and adamantane pendant groups to drive supramolecular self-assembly of the hydrogels.^{170,171,178,182} By changing the concentration of β -CD and adamantane modified HA, the authors were able to tune the shear-thinning, self-healing properties as well as the overall gel strength. These initial studies also demonstrated the ability to tune the release of BSA to beyond 60 days. Specifically, by changing the hydrogels from a 2.5 wt% to 10 wt% gel, the time it took to achieve 90% cumulative release doubled.⁴³

Using this same system, Burdick and coworkers also demonstrated modular tuning of release.¹⁷⁸ The authors were able to tune the cumulative release profile through either varying the total polymer wt%, the Ad : CD ratio, or the affinity of the peptide cargo. All release profiles achieved sustained release of the two peptides for up to three weeks, while exhibiting different controllable parameters that can be used to tailor the material for a wide range of payloads. This work culminated in tuning the HA-CD gels to deliver two clinical therapeutics of different affinities with CD for two weeks. Collectively, these results illustrate how hydrogels dynamically

crosslinked with CD complexes can be used to create injectable materials capable of releasing peptide drugs over weeks, while also remaining versatile as a materials platform to be tuned for a wide range of cargo.^{178,183}

In another example of hydrogels based on host-guest interactions, Scherman and coworkers developed a system based on CB[8] self-assembly with two different guests.⁷⁷ The authors modified cellulose derivatives with either viologen or naphthoyl derivatives and demonstrated that the presence of CB[8] induced gelation due to the formation of ternary complexes. By tuning the crosslink density, the authors were able to control pore sizes, mechanical strength, and drug release kinetics.^{75,76} Extremely sustained release of BSA was observed over 5 months from a hydrogel containing only 1.5 wt% of polymer. Furthermore, the bioactivity of the proteins were maintained for over a month, demonstrating the potential for using these hydrogels for long-term controlled release.^{78,79}

Kim has also shown hydrogel formation using CB[6] and alkylammonium guests derived from 1,6-diaminohexane (DAH) and spermine (SPM).⁶⁹ These gels were made by grafting (allyloxy)₁₂CB[6] to a thiol-functionalized HA through a thiol-ene reaction. By mixing the CB[6] containing polymer with DAH-containing HA, a hydrogel was produced after several minutes. These gels exhibited a storage modulus as high as 3.4 kPa for a hydrogel comprised of 2 wt% polymer. Moreover, cytocompatibility studies demonstrated high cell viability, enzymatic degradability, and negligible cytotoxicity of the hydrogels. When injected under the skin of mice, the hydrogel formed within a few minutes and was stable for longer than two weeks. More recently, the authors have been investigating the same CB[6] based platform for long term delivery of engineered mesenchymal stem cells (eMSCs).⁷¹ In this work the authors used CB[6]-HA combined with DAH-HA and drug conjugated HA. The eMSCs produced enhanced green fluorescence protein in mice for more than 60 days, demonstrating the hydrogel's feasibility in providing long term cell therapies. This work exemplifies how the modularity of supramolecular chemistries enable the researchers to finely tailor their materials platform for cell and protein delivery. This flexibility has allowed for supramolecular materials to solve many challenges in biomedical fields such as immunology and regenerative medicine (*vide infra*).

Protein interactions have also been employed to assemble gels for drug delivery. Kiick illustrated this strategy by using heparin-modified PEG copolymers to drive assembly when introduced to proteins or peptides.¹³⁴ Star PEG polymers terminated with heparin-binding peptides (HBP) were mixed with heparin-modified star PEG polymers to form robust gels with $G' \sim 200$ Pa ($\tan(\delta) \sim 0.05$) over $0.1\text{--}100$ s⁻¹. Contrary to the previous drug delivery systems, the solute release kinetics were mediated by matrix erosion as the star PEG polymers eroded off the matrix. By changing the ratio of heparin containing PEG to PEG-HBP, they were able to obtain a wide range of cumulative release profiles over a two week period. Notably, the modulus and relative elasticity of these heparinized hydrogels can be easily tuned by modulating the heparin : HBP ratio.

This tunability in conjunction with heparin's biological versatility has allowed heparin based assembly motifs to be used for a variety of different drug delivery strategies.¹⁸⁴

While not strictly based on directional supramolecular interactions, nanocomposites formed from polymer-nanoparticle (PNP) interactions have also been explored for their potential for injectable, controlled release platforms. Langer developed shear-thinning injectable hydrogels that exploit dynamic non-covalent crosslinks between polymers and nanoparticles.¹⁸⁵ These PNP gels were made of hydrophobically modified (hydroxypropyl)methyl cellulose (HPMC-C_x) that acted as the main network component and nanoparticles composed of poly(ethylene glycol)-*block*-poly(lactic acid) (PEG-*b*-PLA). When mixed, the HPMC-C_x would rapidly and selectively adsorb onto the nanoparticles, forming the physically and dynamically crosslinked gel. Using this materials platform, they demonstrated hierarchical drug release by encapsulating cargo inside the nanoparticles in addition to the bulk of the PNP gel.¹⁸⁹ This co-loading allowed for control over two different mechanisms of drug release—Fickian diffusion from the bulk and erosive release from the nanoparticles. Lastly, they demonstrated *in vivo* functional validation by simultaneously delivering a model hydrophobic cargo and dye labeled BSA.

4.1.2 Microcapsules. In addition to supramolecular hydrogels, drug delivery has also been approached by using supramolecular chemistries to form microcapsules (micron sized particles with core-shell architectures). A major biomaterials challenge is solubilizing and stabilizing pharmaceuticals for delivery.¹⁸⁶ Current strategies include using co-crystals, pro-drug formulations, salt formation, and drug complexation with a molecular capsule. For molecular capsules, the capsule formation can involve sacrificial core templates, hydrogen bonding, electrostatic interactions, covalent bonds, and host-guest interactions. While many methods to make capsules involve layer-by-layer (LBL) assembly of polyelectrolytes, development of the assembly process is typically driven by improving processing methods rather than significant focus on the non-covalent interactions and will not be focused on here.^{187–190} On the other hand, host-guest chemistries provide a straightforward method for molecular recognition and encapsulation of a chosen drug for delivery. The main challenges and considerations when using host-guest interactions for the formation of capsules include aqueous solubility, kinetics of dissolution, ease of synthesis, and material biocompatibility.

Cyclodextrins have typically been used for capsule formation due to their GRAS-listed status, industrial scale production, and biocompatibility.^{39,191,192} Captisol uses β -CD derivatives to solubilize a number of different drugs and Stella and coworkers have demonstrated that a variety of charged drugs can be encapsulated by using different β -CD derivatives.¹⁹³ Extending LBL assembly technologies for construction of CD multilayers,¹⁹⁴ Auzély-Velty and coworkers developed hollow capsules using HA- β -CD to deliver paclitaxel, an anti-cancer drug.¹⁹⁵ β -CD formed host-guest complexes with the

hydrophobic paclitaxel, greatly enhancing water solubility of paclitaxel, which is a major problem for clinical administration. Release of paclitaxel is controlled by the K_{eq} of the host-guest complex and the authors demonstrated controlled release kinetics, while maintaining efficacy towards killing breast cancer cells. Moreover, the ability for β -CD to pre-complex with paclitaxel circumvents reduced encapsulation efficiency typically seen in capsule formation using LBL techniques, where cargo is loaded after fabrication of the capsule. Beyond CD, calix[4]arenes have also been explored since the 1990s for their use in capsule formation and molecular recognition. These capsules involve the formation of dimeric architectures with two calix[4]arenes encapsulating various small molecule guests and hope to overcome the low K_{eq} values and relatively poor selectivity of CD complexes.¹⁹⁶ Some examples of improving calix[4]arene capsule solubility involve attaching large benzylic groups to the lower rim, or alkyl substituted aromatics along the urea groups of the upper rim. Liu and coworkers demonstrated the formation of supramolecular capsules based on *p*-sulfonatocalix[4]arene and asymmetric viologen host-guest complexes to release doxorubicin.¹⁹⁷ Notably, these capsules were shown to be stimuli responsive to temperature, redox, and host-guest inclusion. This multi-responsiveness is exemplary of the supramolecular host-guest interactions. Due to the enthalpy-driven viologen-calix[4]arene interaction, the interactions are weakened with increasing temperature. Furthermore, redox responsiveness was included by deliberately using a viologen guest, which are known to transform into radical cations or neutral molecules *via* redox reactions. This study not only demonstrates the use of calix[4]arenes in designing capsules for drug encapsulation and delivery, but also exemplifies how the inherent modularity of host-guest interactions enable novel and creative materials.¹⁹⁷

CB-based complexes have also been used to make capsules and provide additional tunability. While CB complexes have relatively slow guest dissociation kinetics and more rigid structures than other hosts, CB[5] and CB[7] have relatively high water solubilities of 20–30 mM in neutral water and have demonstrated suitable biocompatibility.^{198–200} Accordingly, CB[7] has been widely studied for encapsulating drugs, while maintaining bioactivity.^{201–204} Isaacs and coworkers demonstrated that acyclic CB-based containers increased the solubility of ten different insoluble drugs by several orders of magnitude. Furthermore, the CB materials exhibited low *in vitro* toxicity in the presence of human liver, kidney, and monocyte cells, and also increased concentrations of paclitaxel encapsulated in the capsules.²⁰⁵ In addition to CB[5] and CB[7], there has been considerable interest in using CB[8] for capsule formation due to the ability for CB[8] to form higher stoichiometric complexes.^{206–209} Abell and coworkers used CB[8] to form 1 : 1 : 1 ternary complexes through multiple supramolecular interactions between an electron-deficient methyl viologen (MV) and an electron-rich naphthol (Np) derivative (Fig. 12). By leveraging the ternary structure formation of CB[8] host-guest interactions, the authors were able to “handcuff” viologen modified gold nanoparticles to a polymer network con-

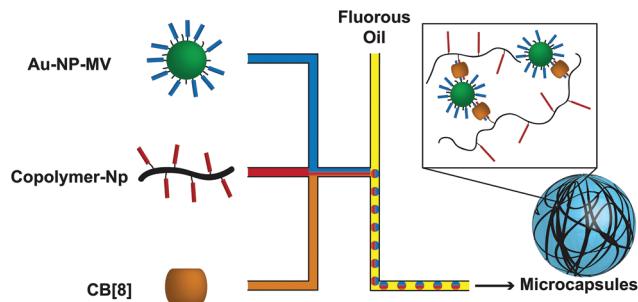


Fig. 12 CB[8] ternary complexes were used to form microcapsules using microfluidics processing. The MV functionalized Au NPs were mixed with a copolymer functionalized with Np pendant groups and a CB[8] solution for efficient and facile formation of microcapsules. Microfluidics processing allowed for formation of monodisperse capsules with diameters of $59.6 \pm 0.8 \mu\text{m}$.

taining naphthol pendant groups to form controlled dispersions of NPs in the network. Furthermore, the highly stable and strong 1:1:1 ternary complexes allowed the microcapsules to be stable at 100 °C and lower pressures (20 Pa). Similarly to the use of viologens in multifunctional theranostic particles (*vide supra*), the use of MV allowed for stimuli responsive degradation of the capsules by reduction of the MV group. The use of microfluidics allowed the authors to obtain monodisperse, high throughput, one-step fabrication of these CB[8] microcapsules, demonstrating the efficiency and scalability of the process (Fig. 12).²⁰⁷

As a whole, these examples illustrate how supramolecular chemistries can improve current microcapsule designs. In particular, the innate modularity and tunability of host-guest interactions enables many different functionalities that are amenable to many current and future microcapsule designs.

4.2 Immunology

Over the last 30 years our understanding of the immune system has increased tremendously, enabling new approaches for engineering immune control. Biomaterials are a powerful tool for designing new therapeutic strategies to modulate the immune system. Below we outline the few examples of immune modulating platforms that exploit the supramolecular systems that have been described in this review. To elucidate the potential for growth in the use of supramolecular polymeric biomaterials for immune applications, we will also provide examples that utilize the highly specific, directional, and strong supramolecular interactions in materials platforms not explored previously in this review. These materials have been used in many areas of this field such as vaccine design, cancer immunotherapies, and general immunomodulation. The research applications that follow indicate the great potential for the use of supramolecular materials for understanding and influencing the immune system.

Supramolecular materials held together by host-guest chemistries have been used to provide controlled release of cytokines and therapeutics for immunomodulation. Physically

crosslinked hydrogels are desirable materials for cargo delivery because they are both easily loaded and self-heal after injection. Heilshorn and Soranno take advantage of these properties when creating an injectable hyaluronic acid hydrogel grafted with adamantane and CD to deliver interleukin-10 (IL-10) and anti-transforming growth factor β into chronic kidney disease affected kidneys. This treatment successfully produced local delivery depots that reduced macrophage infiltration in a kidney disease model, though the combined delivery of these molecules did not improve long-term fibrosis.²¹⁰ Similarly, Burdick employed the “dock-and-lock” hydrogels (*vide supra*) for the delivery of IL-10 to treat chronic kidney disease. The gels delivered IL-10 locally for up to 30 days leading to a decrease in fibrosis in mice.²¹¹ Host-guest interactions are also valuable for their modularity. A study used CB[8] to create multipartite vaccines, circumventing challenging synthetic protocols associated with covalently bonding conjugate vaccines. The components of their model antitumor vaccine (tumor antigen, T-helper cell epitope, and TLR2 ligand) were functionalized with either a first or second guest and mixed with CB[8] to create the linked “supramolecular” vaccines. This supramolecular approach to producing conjugate vaccines provides previously unattainable adaptability and ease of synthesis.²¹²

Hydrophobic and entropically driven interactions are also useful supramolecular motifs for immunomodulating applications. Kiyono adapted a cholesteryl-functionalized pullulan (CHP) for intranasal vaccination. CHP self-assembles into physically crosslinked nanogels in water through hydrophobic interactions and traps cargo as it assembles. A similar CHP nanogel successfully improved humoral and T cell responses to the NY-ESO-1 human tumor antigen in a clinical study.^{213,214} Kiyono incorporated cationic functionality on the cholesteryl bearing pullulan to improve nasal uptake of a subunit fragment of *Clostridium botulinum* type-A neurotoxin from the CHP nanogels. This vaccine improved systemic and mucosal immune responses in mice and successfully avoided central nervous system accumulation of the antigen.²¹⁵ A different study developed a self-assembling nanogel from pyridine functionalized poly(hydroxyethyl methacrylate) (pHEMA-pyridine). The polymer, dispersed when unperturbed, self-assembled into nanogels in the presence of protein due to noncovalent interactions. The study demonstrated that these self-assembled nanogels released protein cargo intracellularly in dendritic cells and led to the activation of CD8⁺ T cells. They describe a new platform for delivery of protein antigens for achieving cell mediated immune responses.²¹⁶

Targeted drug delivery to the lymph nodes (LNs), tumor sites, or other tissues of interest remains an important goal of immunoengineering. A particularly innovative application for self-assembling polymer micelles involved the design of amphiphilic polymers with a retinoic acid hydrophobic region in the core and a cationic polyethylenimine hydrophilic outer region. After the polymers self-assembled into micelles, the cationic charges were shielded with hyaluronic acid (HA), limiting the toxicity. After the particles passively diffused to tumor

sites, the high expression of hyaluronidase in tumor micro-environments degraded the HA coating, unshielding the cationic charges and leading to tumor specific cell necrosis. This necrosis activated an immune response against the tumor, therefore achieving an immune modulation effect.²¹⁷ Targeting LNs can also be particularly valuable when designing vaccines in order to maximize the interaction necessary for immune cell activation. Irvine and collaborators were inspired by the supramolecular interactions that allow commonly used LN tracers to bind to serum albumin in order to “hitch-hike” to the LNs. They created polymers with three components; an albumin binding domain, PEG, and the peptide antigen. These polymers formed micelles that were able to bind to albumin and be transported into the LNs in mice. This strategy increased LN accumulation of their model vaccine and led to a 30-fold increase T cell priming.²¹⁸

Rational peptide design enables immunoengineers to create peptide based vaccines that self-assemble like LMWGs, affording unique characteristics for improved immune responses. One approach for preparing supramolecular peptide vaccines employs the well-known β -sheet motif to create highly modular nanofibers and cylindrical micelles.²¹⁹ By conjugating immunological epitopes to the self-assembling β -sheets, engineers have precise control over the epitope content, dose, and the multivalency of their vaccine. Chong emphasized the utility of precise epitope display while conjugating a CD4⁺ T cell epitope and B cell epitope to their supramolecular peptide platform in specific ratios. These ratios affected the immune phenotype, favoring T follicular helper, Th1, or Th2 responses.²²⁰ These technologies provide a strategy to improve the immunogenicity of subunit vaccines without introducing inflammatory adjuvants. Another way to design peptide vaccines requires conjugating the peptides to a lipid tail that will self-assemble into micelles. It is predicted that these micelles provide high local concentrations of the peptide antigen, protect the peptide from degradation, and increase antigen uptake by dendritic cells. A study showing the preliminary efficacy of peptide micelles for protecting against tumors used micelles with an ovalbumin peptide as a prophylactic tumor vaccine. The mice were inoculated with ovalbumin expressing lymphoma cells after receiving three doses of the vaccine. The peptide micelle treatment stimulated a CD8⁺ T cell response against the tumor challenge and improved the protection from the tumor compared to using a traditional adjuvant system.²²¹

The above examples provide a snapshot into the potential impact of supramolecular materials in the design of immune modulating therapeutics. The materials can be used alone or as delivery vehicles to produce state-of-the-art therapies. The relatively few studies that utilize the supramolecular systems we focused on in this review demonstrate that this field is still in its infancy. The examples of additional supramolecular materials used for immune applications reveal the advantages of these interactions in the context of modulating the immune system, and show that there are many directions to explore in developing new biomaterials for this field. For immunoengi-

neering applications, supramolecular material features that have been most impactful are their modularity, injectability, loading efficiencies, and stimuli responsiveness. The design parameters are endless in this space; we believe there is significant room for innovation in both prophylactic and therapeutic immune modulating therapies by employing supramolecular polymeric biomaterials.

4.3 Regenerative medicine

In a succinct editorial, Mason and Dunnill describe regenerative medicine as a tool to replace or regenerate human cells, tissues, or organs, restoring or establishing normal functions.²²² Translationally minded research in the field parses this out into the combinatorial delivery of soluble factors, cells, and scaffolds. The design and employment of novel biomaterials requires intimate knowledge of how cells interact with their physical environment (mechanotransduction) and the temporal presentation of soluble factors.²²³

There are many excellent reviews describing the intricacies of mechanotransduction; we will only briefly discuss material handles on cellular fate.^{224–226} It has been demonstrated that stem cells are specifically responsive to their ability to generate traction forces through integrin mediating binding with their microenvironment. This can be modulated mechanically through changes in both matrix elasticity²²⁷ and viscoelastic relaxation timescales.²²⁸ Moreover, changes in dimensionality²²⁹—culturing cells above (2D) or within (3D) a matrix—and presentation of the ligand—density and spatial clustering²³⁰—affect how cells pull on their matrix and can have profound effects on cell morphology and fate. Like the viscoelastic compliance, the degradability of matrices also change the temporal interaction of cellular materials with their ligands.²³¹ Similarly, there are a collection of great reviews on cellular interaction with soluble factors for regenerative medicine.^{232–234} In regards to soluble factor delivery, it is becoming more apparent that spatiotemporal control of multiple agents over extended timeframes will be required to affect cellular fate and regenerative processes.²³⁵ In addition to these environmental factors, it is important to understand how these materials will be delivered into the body. While surgical implantation relaxes material property constraints, the use of non-invasive protocols through injection provide a more translational alternative.

Supramolecular polymeric materials make a great platform for addressing these needs due to their facile modulation and dynamic nature. The modulus of hydrogels can be affected by changing polymer concentration, backbone architecture, degree of crosslinking, and moiety of the crosslink. The time-scale of viscoelasticity can be modulated by selecting and combining crosslinks with different dissociation rate constants. Moreover, it has been demonstrated that precursors to the hydrogels are non-toxic to cells, allowing for high viability encapsulation into 3D networks. Because these materials are synthetic, it is simple to incorporate biological functionality (degradable motifs and RGD adhesion domains) through polymeric backbone modification or by incorporation into the

actual backbone through recombinant engineering. Due to the dynamic nature of the crosslinks, engineered materials are shear-thinning, and can be either injected directly or through a catheter. The following section will describe the initial efforts mainly clustered into the following two areas: using supramolecular polymeric hydrogel to (1) deliver soluble factors and to (2) create a 3D microenvironment suitable for injection.

4.3.1 Delivery of soluble factors. The two primary methods of soluble factor incorporation into hydrogels are through encapsulation and tethering (either physical or chemical). The release of soluble factors is then a factor of diffusion (if the pore size of the material is sufficient) and material degradability. Because the crosslinks that define the pore sizes of physically crosslinked hydrogels are transient, there is design space to control macroscopic diffusion through the association and dissociation rate constants of physical crosslinks.⁷⁵

Within the scope of encapsulation, Erythropoietin (EPO), a hormone that can prevent apoptosis while enhancing neo-vascularization for myocardial infarctions (MI), was delivered *via* injection in an α -CD PEG-PCL-PEG hydrogel.²³⁶ While sustained release of EPO led to improvement of cardiac function in a MI rat model, the authors argue that its localization into an injected hydrogel reduced systemic off-target effects, such as polycythaemia. A telechelic UPy system has also been employed to deliver soluble factors through a catheter for MI, and deliver 2 growth factors, HGF and IGF.^{237,238} A porcine study demonstrated improved cardiac function and cardiomyocyte generation over a bolus injection of the respective growth factors. Rather than being employed as cargo, growth factors, such as VEGF, can be employed as the actual physical crosslink of a material due to their affinity with LMWH.¹³⁷ Using VEGF as a physical crosslink resulted in triggered degradation of the material in the presence of VEGF receptors, found on cells, opening the ability to design materials for regenerative medicine with cellular mediated release profiles.

4.3.2 Delivery of cells *via* injection. The use of hydrogels for the delivery of stem cells for regenerative medicine has the ability to localize the delivered cells to a targeted area while initially providing an engineered microenvironment that promotes cell viability. While covalent hydrogels are traditionally brittle and cannot be passed through a syringe, the use of supramolecular crosslinks affords direct injection.

Using α -CD poly(pseudo)rotoxane hydrogel networks, mesenchymal stem cells (MSC) were injected into rats and a rabbit MI model.^{239,240} While it seems important to note that in the rabbit model the components were mixed during the injection process using a Duploject applicator, both demonstrated beneficial cell morphology, limited material cytotoxicity, and better regenerative profiles than with simple bolus injection.

Kim and coworkers have initiated studies employing CB[6] and guests grafted onto polymeric backbones for cellular delivery through *in situ* gelation.^{69–71} When cells are mixed with a hydrogel precursor solution (either CB[6]-grafted or guest-grafted polymer) and injected simultaneously with their pre-

cursor counterpart, a hydrogel network results *in situ*; this strategy is used to deliver a 3D cellular microenvironment in animal models. Because the supramolecular hydrogel is synthetically created, chemical motifs can be introduced to release soluble factors to the cells in the engineered microenvironment. Tethering a steroid that promotes chondrogenesis to the polymeric backbone *via* a hydrolytically cleavable ester while physically encapsulating a growth factor afforded independent control of two soluble factors over radically different timescales.⁷⁰ Creative design with the modularity of a supramolecular system allowed control over release and hydrogel mechanics, promoting chondrogenesis through injection into a mouse model. Moreover, a very similar material was used to inject engineered MSCs in a cancer model (Fig. 13).⁷¹ This system released two small molecules, dexamethasone and retanoic acid, that acted as cues for transgene expression of the engineered MSCs. To achieve different release profiles, the cues for transgene expression were directly grafted to the polymeric backbone or appended to a soluble CB[6] (letting the cue-CB[6] attach to a free guest on the polymer backbone). The release of the two agents was then mediated by hydrolysis or the dissociation kinetics of the CB[6] guest pair. The use of this supramolecular hydrogel afforded rapid *in situ* gelation while demonstrating effective transgene expression of engineered MSCs for 5 weeks and increased cellular viability over Matrigel, a traditionally used extracellular matrix for *in vitro* cell culture. The use of this system retarded tumor growth and increased average lifespan over the delivery of engineered MSCs in Matrigel. While these supramolecular motifs afforded synthetic control of release kinetics and hydrogel viscoelasticity, they ultimately were not able to demonstrate the direct injection of a 3D microenvironment; cells in a hydrogel environment formed before injection. To accomplish this, researchers must look to employ material systems with larger power law indexes (larger values of *a*) (Fig. 9a and b).

Using associating protein hydrogels to accomplish this, Heilshorn employed the WW and proline rich domains for stem cell encapsulation and delivery.^{128,241} Specifically, the use of recombinant proteins afforded direct insertion of cell adhesion motifs into the polymer backbone, enabling encapsulation of a wide range of cell types. As is, this engineered system increased cellular retention over both alginate and collagen post-injection in a mouse model. Incorporating a polymeric segment with a LCST below body temperature provided a thermally induced second gelation, increasing the modulus of the material and overall cellular retention post injection.¹²⁹

4.4 3D-Bioprinting

The field of regenerative engineering aims to replace or regenerate human cells, tissues, or organs, restoring or establishing normal functions. Complex spatio-temporal control in the fabrication process of materials for regenerative engineering is necessary to replicate the complexities of biological tissue and is only afforded through 3D-bioprinting; precise layer-by-layer depositions of biological materials, biochemical cues, and living cells.²⁴² 3D-bioprinting is a complex and multidisciplinary

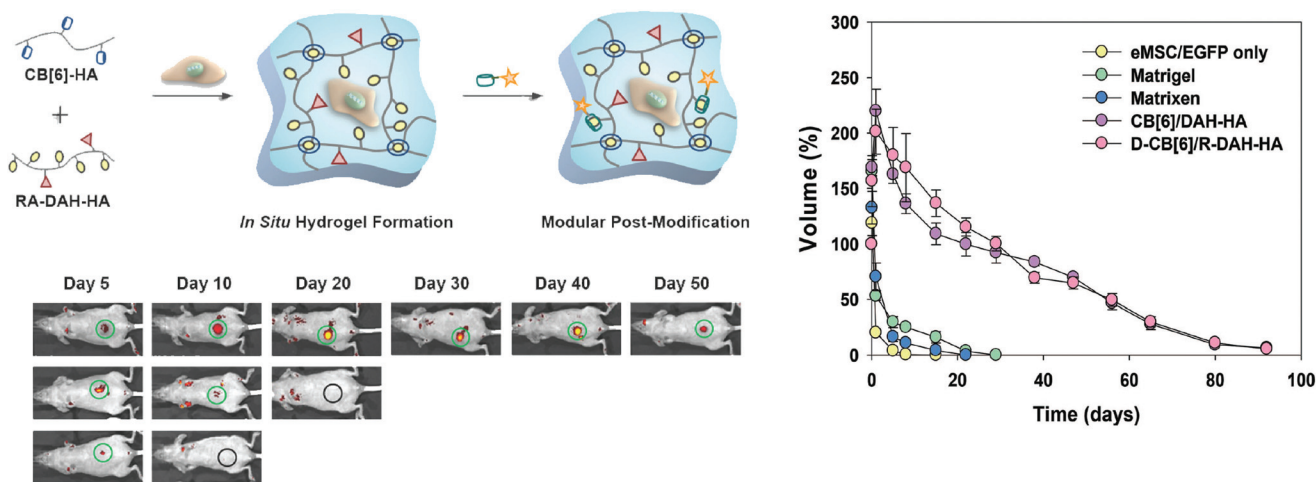


Fig. 13 The usage of post-modified CB[6] based hydrogels to deliver engineered mesenchymal stem cells (eMSC). (a) General scheme: eMSCs are mixed with CB[6] (host) functionalized HA and diaminohexane (DAH, guest) and retinoic acid (RA, transgene expression agent) to form a hydrogel with excess guest molecules. The excess guest molecules then react with addition of dexamethasone (D) functionalized CB[6]. (b) Real time bioluminescence imaging of enhanced green fluorescent protein (EGFP) expression in eMSCs in (i) the hydrogel system, (ii) matrigel, and (iii) serum free media to 50 days. (c) Degradation profile of hydrogel samples in mice over 3 months. The post-modification of D-CB[6] shows no negative effects on degradation profile. Figure reproduced with permission from ref. 71, Wiley.

any problem that involves advances in imaging, biological design, materials selection, cell selection, printer selection, and application engineering. While these have been reviewed elsewhere,^{242–244} we will briefly discuss printer technology, material requirements, and demonstrate the benefits of supramolecular crosslinks in advanced bio-inks.

The most common 3D-bioprinting machines utilize either laser assisted, inkjet, or microextrusion printing technologies.²⁴⁵ Laser assisted bioprinting involves focusing laser pulses onto an absorbing layer of a material, generating a high-pressure bubble that directs cell-containing materials to a collector. Inkjet printers use thermal or acoustic forces to eject drops of liquid onto a substrate. Microextrusion printers deposit continuous beads of material through a nozzle with pneumatic or mechanical (piston or screw) force. The benefits afforded by dynamic crosslinks are most closely associated with aiding printability in microextrusion printing because it requires printing a viscous material through narrow nozzles under force. This is not to say that other printing methods should not employ supramolecular bio-inks. For example, an inkjet bioprinter containing two cartridges of liquid polymer with separate but complimentary supramolecular crosslink components could print supramolecular materials. However, the benefits of using supramolecular materials for this application are no different than the benefits of using supramolecular interactions for traditional regenerative engineering approaches. For this reason, we will continue to describe the benefits of dynamic crosslinks in extrusion based systems.

The spatio-temporal control afforded by 3D-bioprinting requires that the material demonstrate printability, a novel material property similar to injectability.²⁴⁶ In addition to being biocompatible, having controlled degradation kinetics and mechanical properties, and having dynamic biomimetic

materials properties, bio-inks must also be suitable to be extruded from a nozzle under pressure and form defined printed architectures at high resolution while maintaining high cell viability.

While readily available biomaterials have been explored for printing, they are met with many problems. Agarose, methylcellulose, gelatin, and collagen possess gel transition temperatures and must be printed at high temperatures potentially dangerous to cells.²⁴⁷ Further, they form weakly bound structures that often need a secondary crosslinking step. While use of alginate forms robust hydrogels, it is printed as a viscous fluid in which cells experience both sedimentation which can lead to printer clogging and substantial shear forces during extrusion.^{248,249}

To address issues of printability of alginate materials, Heilshorn employed tailored supramolecular interactions on the alginate backbone.²⁵⁰ Alginate was functionalized with a proline rich domain and mixed with alginate functionalized with a WW domain, forming weak (20 Pa), shear-thinning hydrogels. This pre-gel was printed into a bath of CaCl_2 , forming a robust alginate hydrogel (4 kPa) whose material properties largely matched those of crosslinked alginate without peptide crosslinks. By printing alginate as a weak gel rather than a viscous liquid, they demonstrated homogeneity of cells in the cartridge, increased cell membrane integrity during printing, and >90% viability of hASCs and 3T3s a week after printing. While originally demonstrated with complimentary peptide sequences, alginate printability ostensibly would increase with other, more modular supramolecular crosslinks, offering the ability to correlate macroscopic pre-gel properties with cell printability.

Similarly, Burdick modified hyaluronic acid (HA), a utilized bioink, with either β -CD or Ad and printed the hydrogel into a

support gel of near identical material—a unique printing strategy afforded by the shear-thinning of the extruded gel and the rapid self-healing of both the extruded gel and the displaced support network (Fig. 14).²⁵¹ Appending methacrylate units to the HA polymers in the cartridge or HA support gel gives the ability to reinforce the printed architecture with a photo-crosslinking step and subsequently wash away the support. This strategy led to a complex 3D architecture unobtainable without printing into a medium. The washing process, however, involves disrupting all (print and support) host-guest interactions, leaving only the covalent methacrylate crosslinks in the printed structure. In a similar fashion, methacrylate units can be appended to only the support supramolecular gel. Here the support is reinforced and the printed structure is removed through flow applied at the inlet and outlet of the channel by the printer, leaving a support gel with supramolecular crosslinks and well defined channels. Excitingly, the printer could pattern MSCs in the extruded gel, demonstrating >90% viability immediately after printing and >80% viability after 3 days in culture. While printing into a support gel had many advan-

tages, it ultimately could not produce a printed design with physical crosslinks removed from the support. To address this, Burdick explored printing HA appended with methacrylate and either CD or Ad motifs directly onto a substrate with a subsequent photo-crosslinking step.¹⁶³ Here they explore how changes in crosslink density, crosslink nature (supramolecular vs. chemical), and curing method affect the printability and mechanical properties of the printed gel. While they could functionalize printed scaffolds with RGD and seed the scaffolds with 3T3 fibroblasts, direct printing of cells from the printer was not demonstrated.

Rather than modifying an already used bioink, Duncan and collaborators created a new ink backbone with appended dynamic biological crosslinks.²⁵² Here they separated two components, a polypeptide with a grafted nucleic acid sequence (A) complimentary to the sticky ends of a DNA crosslinker (B), into separate printer valves. Strong, optically transparent hydrogels formed in seconds upon co-release of bioink A and B through the printer microvalves. Further, because these hydrogels are composed of peptides with DNA crosslinks they demonstrate biodegradability in the presence of specific nucleases and proteases. Because the dissociation rate constant is much smaller for multivalent, complimentary hydrogen bonding, similar levels of shear-thinning are not recognized for this system as they are for Burdick's, removing the possibility of directly printing gels. This limits the resolution of the system to that of the droplet, 500 μm (as opposed to 35 μm for printing into a support gel and 100 μm for free-standing CD@Ad gels). They demonstrated the printability of AtT-20 cells at exceptional viability (>98%) but did not demonstrate the printing of cell lines more relevant to regenerative medicine applications.

Future development in this field will benefit from utilizing supramolecular crosslinks to tease out structure property relationships between bio-ink and printability. Scalable, tailorable new bioinks should be created that are shear-thinning and maintain high cell viability. With this knowledge, hydrogel properties can be specifically engineered for use in a 3D printer for applications in regenerative medicine.

5 Conclusions

This review illustrates the design toolbox for supramolecular polymeric hydrogels—water soluble polymeric chains that assemble into contiguous networks through physical crosslinks (host-guest, hydrogen bonding, metal-ligand, electrostatic, and protein based interactions). Different from conventional crosslinking, supramolecular crosslinks exploit specific, directional, tunable, and reversible non-covalent interactions. Accordingly, supramolecular polymeric hydrogels exhibit unique and desirable properties not found in covalent systems. Through careful choice of crosslinking motif and polymeric architecture, novel biomaterials can be systematically built to have engineered viscoelasticity and release profiles, with shear-thinning and rapid self-healing properties.

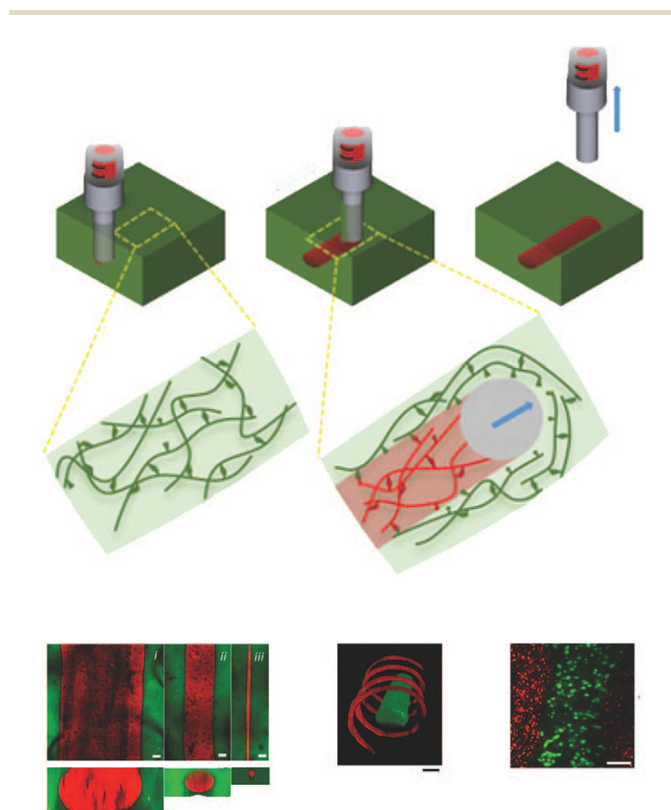


Fig. 14 Burdick's process of 3D printing into a support gel. (a) Illustration of the injection process where the injected gel (red) contains the same CD host-guest chemistry as the support gel (green). (b) Printing different resolutions of rhodamine (red) labeled gel into a fluorescein (green) labeled support gel through a 20 (i), 27 (ii), and 34 (iii) gauge needle (d) Confocal image of two different inks (rhodamine and fluorescein) printed in different geometries. (d) Printing of mesenchymal stem cells (green) into a support gel containing 3T3 fibroblasts. Scalebars in (b): 100 μm , (c), (d): 200 μm . Figure reproduced with permission from ref. 251, Wiley.

Supramolecular polymeric hydrogels are increasingly used as novel biomaterials because their macroscopic material properties can be precisely engineered. The specific, directional, tunable, and reversible nature of a supramolecular interaction facilitates creative solutions to complicated immunological problems. Controlled viscoelasticity enables rheological properties similar to *in vivo* biological tissue, making these materials excellent candidates for controlled microenvironments. Shear-thinning and rapid self-healing allow mechanically strong gels to flow through high gauge needles during injection and then reform *in situ* with minimal cargo loss. Physical crosslinks unperturbed by traditional network swelling afford controlled and extended release through kinetically controlled mesh dynamics. The controlled viscoelasticity, shear-thinning, and self-healing properties enable pneumatic and mechanical printers to extrude well defined cellular microenvironments with incredible precision.

While many studies have developed the synthetic toolbox for designing supramolecular polymeric biomaterials, exploration of the relationship between crosslinking chemistries and material properties has only just begun. The ability to precisely correlate, model, and engineer the relationship between network architecture and supramolecular kinetics and thermodynamics to defined rheological and mechanical properties is presented as a future goal. Work towards this understanding will enable facile implementation of these chemistries into the biomedical community.

Conflicts of interest

There are no conflicts to declare.

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