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Engineering Tough, Energy-Dissipating Soft Materials via Sacrificial Chemical Bonds

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ABSTRACT

Impact-resistant materials such as Kevlar provide impact protection but lack flexibility and processability due to their highly crystalline, rigid structures. In contrast, the natural world is replete with examples of soft materials that can resist applied load. For example, sea cucumbers and other echinoderms modulate the stiffness of their outer skin in response to predators. This dynamic change in mechanical properties is regulated by transient, non-covalent interactions between the collagenic protein fibers that comprise the dermis. Upon the application of force, these transient interactions break, thereby acting as sacrificial bonds and providing a means of energy dissipation without damaging the protein fibers. This mechanism has been mimicked in synthetic materials, typically based on covalent polymers augmented with non-covalent bonding motifs. In this review, we survey the current state-of-the-art in the field of energy-dissipating soft materials. Specifically, we present recent (post-2017) highlights from the primary literature, organized by the chemical nature of the weak sacrificial bonds to outline the chemical toolkit available for programming material properties. We conclude by highlighting some opportunities to advance the development of soft yet tough energy-dissipating soft materials that harness the sacrificial bond.

1. Introduction

The ability of natural materials to withstand mechanical force without damage is crucial for the survival of living organisms; consider the thousands of impacts endured by a human foot during a day's walking or by a woodpecker's skull during a day's drilling.¹⁻³ The resilience of these materials is even more remarkable when one considers that *strength* (force withstood by a material) and *toughness* (resistance to fracture) are generally considered to be mutually exclusive in canonical materials,^{4,5} especially soft materials.^{6,7} Consequently, nature has evolved various approaches to prevent irreversible (plastic) material damage under applied load, including architectural hierarchy, porosity, and the use of composite structures that utilize small amounts of soft materials to absorb and dissipate energy (Figure 1).^{3,5} For example, the extraordinary toughness of abalone shells and nacre—the iridescent internal layer of mollusc shells known also as mother of pearl—derives from their 5% biopolymer content rather than their major aragonite component.⁸ This biopolymer comprises a mixture of proteins such as lustrin, which reversibly folds and unfolds in response to force, and polysaccharides such as chitin, which engages in polyvalent hydrogen-bonding. Similarly, collagen molecules endow bone with its hallmark toughness by forming thin, glue-like filaments that bridge mineral components (Figure 1, bottom right).^{9,10} Individual collagen fibers and fibrils within the filament can slip against each other due to the breakage and subsequent re-formation of sacrificial non-covalent interactions bind these filaments together.

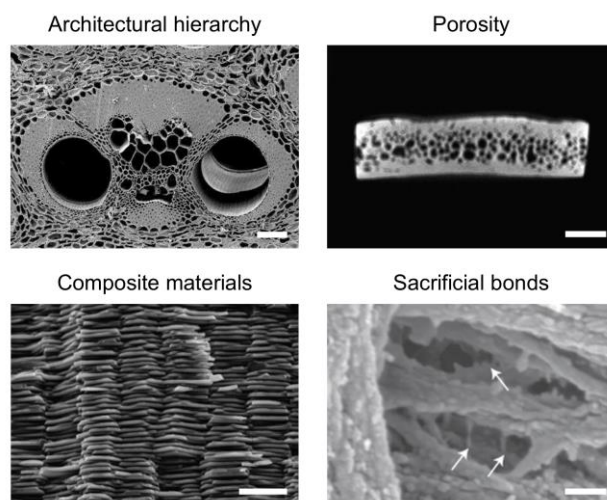


Figure 1. Biological strategies to achieve impact resistance in natural materials. Clockwise from top left: architectural hierarchy in bamboo's culm wall (SEM, scale bar: 100 μm ; adapted from ref. 11, copyright 2010 Budapest University of Technology and Economics); porosity in a turtle's shell (X-ray CT, scale bar: 2 mm; reprinted from ref. 12 with permission from Elsevier, copyright 2009); composite materials such as nacre in seawater snails (SEM, scale bar: 10 μm ; reprinted from ref. 13 with permission from Elsevier, copyright 2008), and collagen-based sacrificial bonds [white arrows] within fractured bone (AFM, scale bar: 200 nm; adapted from ref. 14 with permission from Springer Nature, copyright 2005).

The soft components of these composite materials harness *sacrificial bonds* to absorb and dissipate energy.¹⁵ Sacrificial bonds are comparatively weak bonds within a polymer chain or network that, under applied force, break before the (typically covalent) bonds within the strong polymer network.^{16,17} Under mechanical stress, rupture of these weak bonds dissipates energy while protecting the integrity of the strong matrix.⁶ If the fission and formation of these sacrificial bonds is reversible, their presence can enhance a material's toughness under multiple cycles of loading and unloading. In this way, natural biomaterials harness structural failure, through the rupture of deliberately weak sacrificial bonds, to tolerate mechanical force.¹

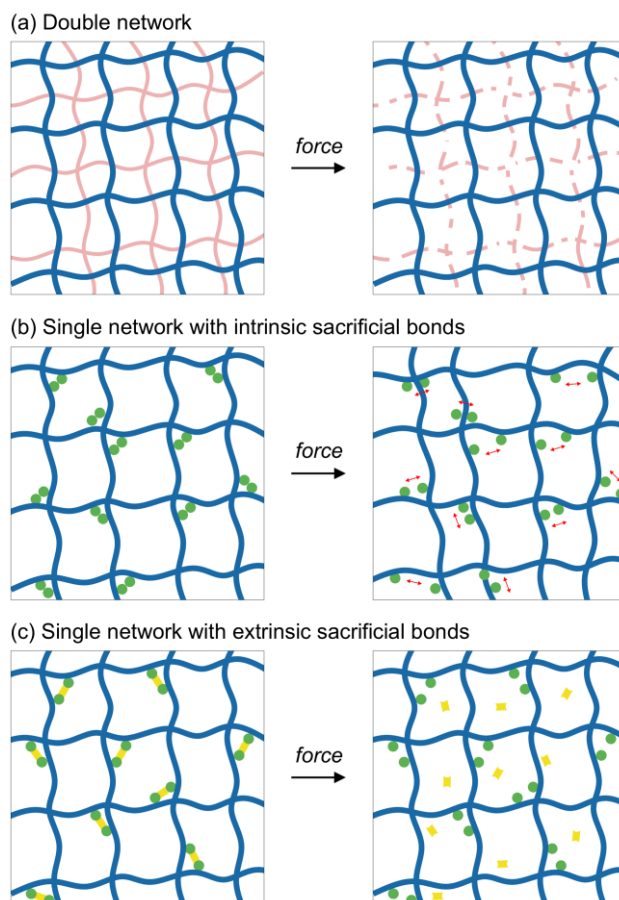


Figure 2. Strategies to incorporate sacrificial bonds into synthetic materials. (a) In double network materials, application of force breaks a weaker sacrificial network (pink), thereby protecting the stronger structural network (blue); these materials were reviewed recently elsewhere.¹⁸ (b, c) In single network materials, applying force breaks the weak, sacrificial bonds that exist between interacting motifs (green) on the structural network (blue); such materials are the focus of this review. Intrinsic sacrificial bonds (b) are those resulting from direct interaction between polymer chains, whereas extrinsic sacrificial bonds (c) rely on small molecule mediators (yellow) to define intra-chain interactions.

Due to the attractiveness of strong yet tough soft materials, there has been substantial research effort expended towards incorporating sacrificial mechanisms into synthetic materials. These efforts can broadly be categorized based on whether weak sacrificial bonds are incorporated as a separate material component or as part of the primary

material matrix (Figure 2).¹⁹ In the former strategy, materials containing two interpenetrating polymer networks rely on the preferential rupture of the weaker sacrificial network to dissipate energy while maintaining the integrity of the stronger interpenetrated matrix (Figure 2a).^{19,20} Such materials, known as double network materials,²¹ are not discussed further in this review due to an excellent recent discussion by Gong and coworkers.¹⁸ Instead, this review focuses on the latter strategy, whereby a single polymer network is responsible both for the material's strength and toughness, aided by weak sacrificial interactions that, at rest, hold together strong polymer chains, but rupture preferentially under mechanical stress (Figure 2b, c). This approach has also garnered substantial research interest, as excellently summarized by Hu and coworkers in 2017.¹⁷

Our aim in this review is to present recent (post-2017) highlights from the primary literature, organized by the chemical nature of the weak sacrificial bonds. This organizational scheme was motivated by our belief that the key challenges in the field have evolved from *preparing* energy-dissipating materials to *designing* specific properties into those materials. Reviews by others have focused on mesoscale design strategies, including the nature of the polymer backbone itself.²²⁻²⁶ Instead, we offer herein a chemist's perspective on the design of sacrificial interactions and thus hope to provide the reader with a better sense of the chemical toolkit available for programming material properties (Figure 3). We begin by defining some key terms used to describe materials properties and then introduce recent advances in the theoretical treatment of sacrificial bond-containing materials. The bulk of the review, describing recent experimental progress, follows. We conclude by highlighting some outstanding challenges and proposing potential research avenues to continue progress towards soft materials with extraordinary strength and toughness enabled by the sacrificial bond.

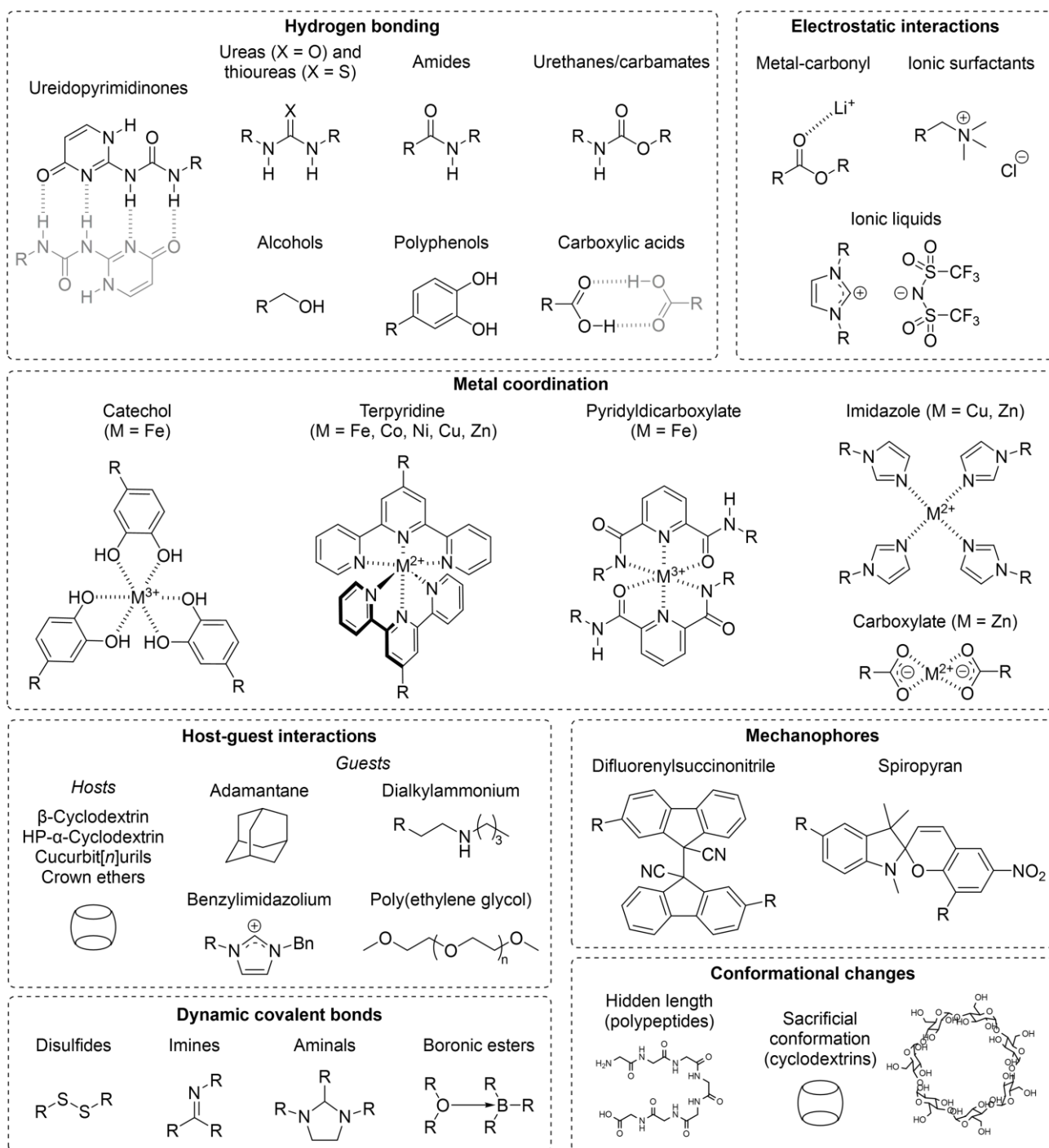


Figure 3. Chemical toolkit described in this review. In all structures, R denotes an arbitrary group and differs from structure to structure.

2. Terminology for Mechanical Properties

In this section, we briefly define the terms used in this review to describe the mechanical properties of materials. Many rheological properties focus on how materials respond to external applied forces; the measure of amount of force applied to a given area of material is denoted *stress*. *Tensile stress* is the most reported type of stress, measured by stretching a material along the axis of measurement. This stretching of the material is a form of *deformation*; the

extent of deformation (i.e., the ratio of a material dimension under deformation vs. at rest) is defined as *strain*. Thus, strain is a direct measure of physical change in a material and may be associated with a change in length (*axial strain*) or change in volume (*volumetric strain*). Experimentally, the rate at which a material is forcibly deformed (i.e., the *strain rate*) plays a substantial role in the material's stress response. At small strains, the relationship between force applied (stress) and material deformation (strain) is linear and *elastic* (that is, reversible); in this case, the stress-strain relationship adheres to *Hooke's law* and the ratio of tensile stress over axial strain is described by the *Young's (elastic) modulus*. This constant indicates the extent of deformation under a given applied stress and is characteristic of each material, ranging from $\sim 10^3$ Pa for gels to $\sim 10^9$ Pa for metals. The *limit of proportionality* denotes the maximum stress at which Hooke's law applied; beyond this limit, stress is no longer directly proportional to strain.

The resistance of a material to deformation may be characterized by different terms depending on the mode of failure. *Tensile strength* is defined as the maximum stress that a material can withstand when pulled before permanent deformations occur, such as irreversible shape changes or material fission. Some materials tend to crack or fracture when stress is applied; *fracture stress* is the stress at which a material fails due to fractures, and the amount of energy that leads to crack propagation is known as *fracture energy*.

Viscoelastic materials behave differently depending on the applied strain rate; this time-dependent response to stress leads to different regimes in which the material exhibits more solid-like or liquid-like behavior at different strain rates. Viscoelastic materials with the ability to stretch and return to original shape without damage have high *stretchability*, also termed *extensibility*. Typically, this ability is limited: the *maximum elongation* measures the maximum (axial) strain before permanent damage occurs. Materials subjected to constant stress over time might experience a gradual, time-dependent deformation, called *creep elongation*.

Energy dissipation is commonly quantified using cyclic loading-unloading experiments, whereby materials are subjected to an increasing external load which is gradually removed (unloaded). Stress-strain plots of the loading and unloading phases differ: the loading cycle represents the strain energy required to generate a given stress within the material under measurement, while the unloading cycle represents the residual elastic energy of the material at that stress. This residual elastic energy is typically lower due to the energy dissipation via heat or fission of sacrificial bonds; consequently, the stress-strain plot of a loading-unloading cycle exhibits a *hysteresis loop* whose

area denotes the total energy dissipated by the material. After a waiting period (*relaxation time*), this loading-unloading cycle is repeated and compared to the initial cycle. If a material does not return to its original length after removal of the external load, the material has deformed and its recovered length is called the *residual length*.

For viscoelastic materials, rheological measurements also quantify two key measures of a material's response to external stress: the storage and loss moduli. The *storage modulus* (G') represents a material's ability to store elastic energy, while the *loss modulus* (G'') represents its ability to dissipate energy as heat. The ratio of G'' to G' , known as $\tan \delta$, quantifies a material's ability to absorb and dissipate energy: values of $\tan \delta$ less than 1 denote a more elastic material, while values greater than 1 denote a predominance of viscous behavior and a more plastic material.

3. Theoretical Studies

Much interest in sacrificial bond-containing materials stems from their non-linear mechanical properties. However, this non-linearity complicates efforts to faithfully describe such materials using canonical polymer physics descriptions such as the worm-like chain (WLC) model, which assumes that the motion of each segment in a semiflexible polymer chain is correlated to some extent with its neighboring segments.¹ In particular, theoretical treatments of energy-dissipating materials must consider the topology of the strong polymer network before and after load is applied, as well as the rate dependence of sacrificial bond rupture and formation. Thus, the faithful modeling of energy-dissipating materials requires consideration of both thermodynamic and kinetic considerations.

Kroy and coworkers have approached this problem by considering the inelasticity intrinsic to biomechanics.¹ Their initial model proposed a 'glassy' WLC model that accounts for the reduced dynamics of a polymer under confinement in a network,²⁷ and was subsequently updated to include transient crosslinks, in a revised *inelastic glassy WLC* (iGWLC) model (Figure 4).^{28,29} The presence of transient crosslinks, akin to sacrificial bonds, are parametrized with a single time-dependent variable that represents the fraction of such bonds that are 'closed'. Despite the simplicity of this treatment, the authors are able to reproduce non-linear properties including hysteresis and strain-rate dependent mechanical response.¹

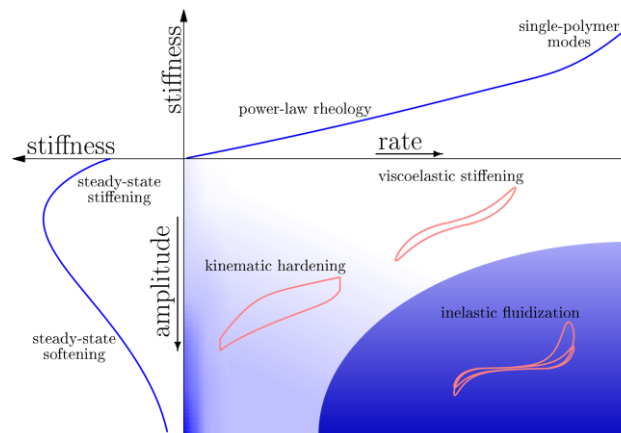


Figure 4. Kroy and coworkers' inelastic glassy wormlike chain (iGWLC) model reproduces the non-linear mechanical response of biopolymers. At low rates where power-law rheology is exhibited (top panel, log scales), the model predicts entropic stiffening of the polymer backbone at low amplitudes (left panel, linear scale). This stiffening is superseded at higher amplitudes by exponential bond softening. At high rates and high amplitudes, rapid stiffening is followed by fluidization. Reprinted from ref. 29. Copyright 2012, Wolff et al., licensed under CC BY.

Incorporating thermodynamic details of specific chemical bonds was achieved by Elbanna and coworkers, who developed a polymer network model (based on the WLC) supplemented with rate-dependent damage behavior.³⁰ In their model, sacrificial bonds are described using a transition state theory (TST) approach that considers each bond to have two stable states (associated or dissociated), separated by a transition state with an accompanying activation energy (E_a). This E_a term contains a contribution that accounts for the force-dependent difference in nuclear coordinate between a given bond and the transition state. A term is also incorporated to account for the rate of sacrificial bond-breaking. Using this model, the authors found that increasing the concentration of sacrificial crosslinkers does not lead to a monotonic increase in the peak force withstood by the polymer network, but instead a maximum was reached due to the prevalence of short chain lengths between adjacent crosslinks. The authors also used their model to probe the relationship between the velocity at which tensile force is applied to a material and peak force withstood by that material. However, the re-formation of sacrificial bonds was not investigated by the authors in that report.³⁰ A similar, damage-induced, energy dissipation mechanism was incorporated into a recent model by Sun, Yin, and Liew that sought to correlate microscale structural damage with macroscale mechanical properties in double-network materials.³¹

The topic of breaking and re-forming sacrificial bonds has been modelled through various approaches. Suhail, Banerjee, and Rajesh developed a course-grained kinetic model to describe energy dissipation in fibrils of collagen.³² Their model not only accounts for the reformation of sacrificial bonds, but also considers how *hidden*

length—segments of a polymer chain released upon rupture of an intra-chain sacrificial bond—affects energy dissipation under cyclic loading. The authors compare their kinetic model against experimental data and find that several aspects of the stress-strain response, including cycle-dependent energy dissipation (known also as moving hysteresis loops), are well reproduced.

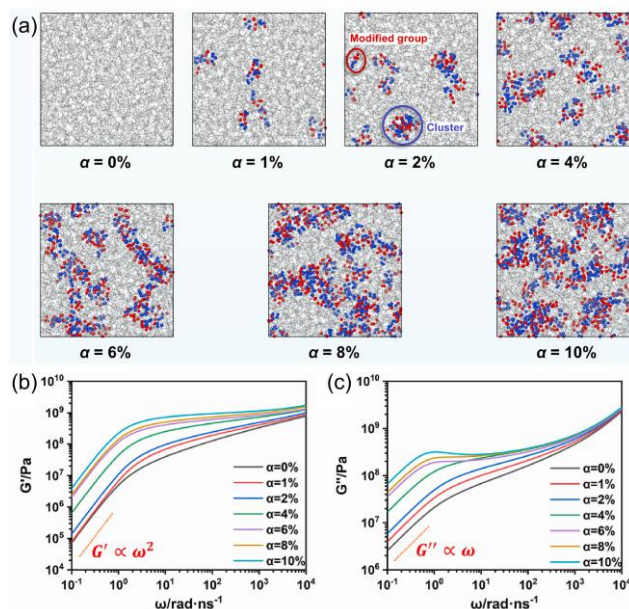


Figure 5. (a) Molecular dynamics (MD) simulations of poly(isoprene) with different monomer ratios (α) of an hydrogen bond (HB) competent motif. Red and blue spheres denote oxygen and nitrogen atoms, respectively, of the HB motif. (b, c) Variation of (b) storage modulus, G' , and (c) loss modulus, G'' , as a function of frequency (ω). Reprinted from ref. 33 with permission from Elsevier, copyright 2023.

An alternative approach by Liu and coworkers employed molecular dynamics (MD) simulations to investigate the breaking and re-forming of specific chemical hydrogen-bonding motifs in polymer networks based on poly(isoprene) (Figure 5).^{33,34} Polymer backbones were investigated with different extents of modification with a hydrogen bond-capable motif, either 3-amino-1,2,4-triazole³³ or a tetraalanine peptide.³⁴ Fully atomistic MD studies allowed the authors to observe the extent of sacrificial bond clustering in these materials and reproduced the finding of Elbanna and coworkers³⁰ that desirable mechanical properties initially increase with increasing sacrificial bond content before reaching a maximum and decreasing as the number of sacrificial bond motifs continues to increase. Atomistic MD studies were also employed by Xie *et al.* in their recent development of a multiscale model to describe the fracture mechanics of self-healing, energy-dissipating composite materials such as nacre and bone.³⁵ MD simulations on the nanoscale were coupled with shear-lag and crack-bridging models at the meso- and macroscales, respectively, to produce an integrated framework for predicting materials properties based on structural parameters.

One emerging area for experimental investigation is the incorporation of multiple different types of sacrificial bond within the same material (as will be discussed later). Such materials are also being explored at the vanguard of current theoretical efforts. In 2022, Yu *et al.* reported a coarse-grained MD treatment of networks of carbon nanotube fibers crosslinked with two distinct sets of sacrificial bonds, one much weaker than the other.³⁶ The authors explore several features of their simulated material, including conductivity and Young's modulus, as a function of both overall crosslinking density and the ratio of strong and weak crosslinks. Weak crosslinks are observed to enhance the toughness of the doubly-crosslinked networks, while stronger crosslinks contribute little to material toughness but underlie material strength. Analysis of the spatial distribution of broken crosslinks after stress suggests that heterogeneous crosslinking leads to stress concentrations within the material, resulting in synergistic rupture and a marked decrease in overall resistance to deformation. These results emphasize the importance of the relative interaction strengths and spatial distribution of multiple sacrificial bonds within a single material.

4. Hydrogen Bonding

Hydrogen bonds (HBs) are particularly attractive candidates for sacrificial bonds, due to their tunable binding strength, directional interactions, and readily reversible binding modality.³⁷ Biological materials, such as silk and muscles, often utilize HBs to crosslink polypeptides to form a primary structural network.³⁸ Following a similar strategy, several synthetic materials have been reported in the past two decades that exhibit excellent energy dissipating and mechanical properties.^{39–41} The most commonly employed approach has been to modify polymer chains with prosthetic HB-capable motifs (for example, ureidopyrimidinones and amides) to define sacrificial interactions between polymers.^{42–45} Additionally, some studies have leveraged intrinsic HB acceptors and donors in the chemical structure of the strong polymer network (for example, poly(vinyl alcohol) and poly(ether-thiourea)) to augment the prosthetic hydrogen-bonding motifs.

4.1. Ureidopyrimidinones

Inspired by DNA base-pairing, Meijer and coworkers designed the ureidopyrimidinone (UPy) motif (Figure 6), a self-complementary quadruple H-bond unit that is attractive for materials design due to its strong binding affinity and facile synthetic accessibility.⁴⁶ In the solution state, UPy exists in an equilibrium between two isomers, 2-ureido-4[1*H*]-pyrimidinone and 2-ureido-4[1*H*]-pyrimidinol, due to keto-enol tautomerization (Figure 6, top). The pyrimidinone derivative has an *AADD* (*A* = acceptor, *D* = donor) H-bonding pattern and the other derivative has a

DADA H-bonding pattern. Both conformers are interconvertible and the ratio of the two tautomers in a specific environment depends on the nature of the substituents at the R_1 and R_2 positions (Figure 6). The presence of *n*-alkyl chains at the R_1 and R_2 positions favors the *AADD* tautomer, while in the presence of any electron-withdrawing group, the *DADA* tautomer predominates. Both tautomers are self-complementary, that is, a given tautomer can associate with itself while forming four HBs. Due to this quadruple hydrogen bond, UPy exhibits high dimerization constants ($K_a > 10^5 \text{ M}^{-1}$ for the *DADA* tautomer and $5.7 \times 10^7 \text{ M}^{-1}$ for the *AADD* tautomer).⁴⁷

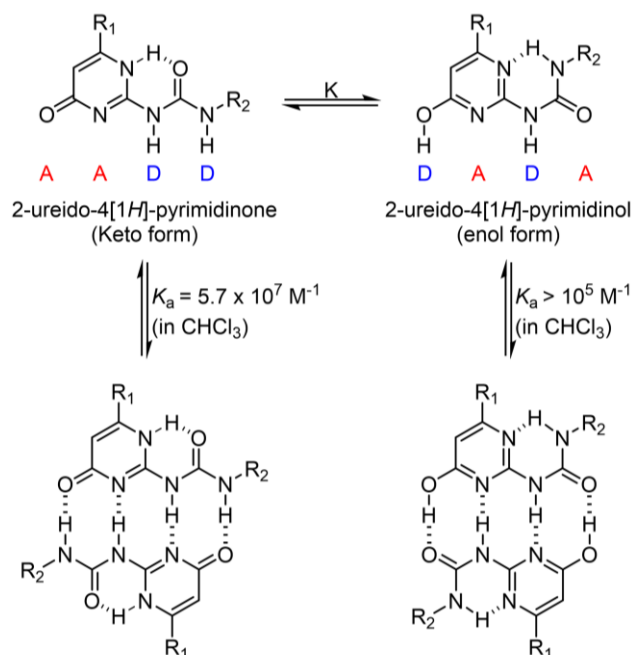


Figure 6. UPy exists as two tautomers at equilibrium (top). Each tautomer can respectively dimerize due to self-complementary quadruple hydrogen bonding to form homodimers (bottom). Reprinted from ref. 46 with permission from Elsevier, copyright 2023.

Li and coworkers employed the UPy motif to design bioinspired, HB-crosslinked healable polymers with remarkable energy dissipating and healing properties.⁴⁴ Poly(propylene glycol) (PPG) or poly(tetramethylene ether glycol) (PTMEG) chains (soft segment) were combined with cyclohexylmethanemethane-4,4'-diisocyanate (HMDI) or isophorone diisocyanate (IPDI) (hard segment) as well as UPy units (Figure 7). The reported polymer has several units, such as UPy, urea, and urethane, capable of forming hierarchical H-bonding networks that lead to hard-soft nanophase segregation. Regulating these nanophases allowed the mechanical and energy dissipating properties of the bulk material to be modified. The samples prepared with PTMEG and HMDI showed elongations of up to 1960% at their breakage point. To investigate the energy dissipation properties, cyclic tensile tests were conducted. Samples showed hysteresis loops as well as energy dissipation at low and high strain levels. At low strain levels,

energy dissipation occurs via the breakage of weak HBs in urethane while at high strain levels, energy is dissipated via the breakage of strong HBs between urea and UPy. In addition to energy dissipation and self-healing, the material showed an exceptionally high toughness of 345 MJ/m^3 , which compares favorably to synthetic materials such as Kevlar (33 MJ/m^3) and natural materials such as spider silk ($50\text{--}500 \text{ MJ/m}^3$).⁴⁸

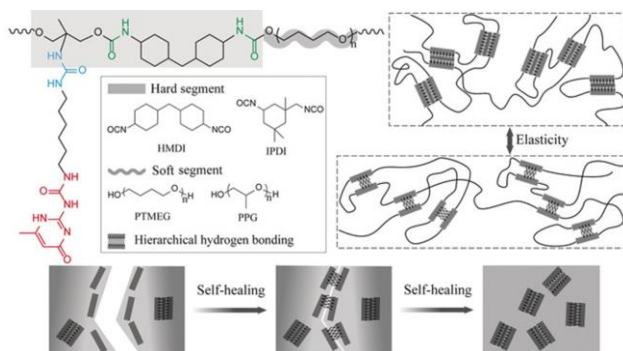


Figure 7. Copolymers of hard and soft segments generate hierarchical H-bonding domains that are responsible for self-healing and energy dissipative properties. Figure adapted from ref. 44. Copyright 2018 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.

Biobased elastomers are attractive materials due to their low modulus, high elasticity, and excellent recovery upon removal of external load,⁴⁹ but their mechanical weakness and limited molecular structures reduce the scope of their potential applications.^{50,51} To overcome this challenge, Guo and coworkers employed a sacrificial bond strategy to conserve material integrity.⁵² Elastomers were synthesized by dissolving various ratios of UPy-based methacrylate (UPyMA) and biobased polyester elastomer (BPE) in DMF (Figure 8a). To confirm the network structure and UPy dimerization, variable temperature (30 to 150 °C) FTIR tests were conducted. The elastomers with higher proportions of UPyMA showed better shape memory behavior (restoration within 30 s), suggesting that increased HB density increases the energy dissipation and shape recovery of the materials.⁵² Stress-strain plots of polymers with 0% and 30% UPyMA showed that the addition of UPy increased the Young's modulus from 0.6 to 3.5 MPa and tensile strength from 0.9 to 5.2 MPa (at 100% strain). Moreover, cyclic loading-unloading tests indicated that the amount of dissipated energy increased with increasing UPyMA content (almost 20-fold from 0% to 30% UPyMA, Figure 8b).

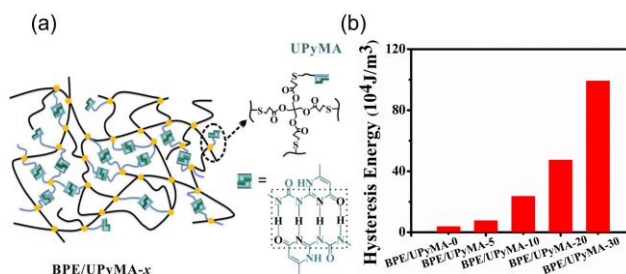


Figure 8. (a) Schematic representation of BPE/UPyMA- x polymeric networks, where x denotes the relative proportion of UPyMA, with UPy quadruple HBs. (b) Energy dissipation (hysteresis loop area) for BPE/UPyMA- x , $x = 0$ to 20. Reprinted from ref. 52 with permission from Elsevier, copyright 2019.

Impact-hardening polymers (IHPs) are a class of energy-absorbing materials with loading rate-dependent viscoelasticity, commonly applied as impact protectors.⁵³ IHPs exhibit an increase in viscosity with increasing shear stress and can be categorized by their physical form under no stress as either shear-thickening fluids (STFs) or shear-stiffening gels (SSGs). STFs, known also as non-Newtonian fluids, have limited applications due to their fluid nature and inability to hold their shape; therefore, SSGs have been more explored for applications.⁵⁴ In 2021, Yan and coworkers designed an impact-protective SSG based on a supramolecular polymeric material (SPM) comprising the triblock copolymer poly(propylene glycol)-*b*-poly(ethylene glycol)-*b*-poly(propylene glycol) (PPG-PEG-PPG) and UPy motifs.⁵⁵ Loading-unloading cycles showed an increase in hysteresis with increasing compression rates, representing a higher degree of energy dissipation. Hammer drop tests were performed to mimic the destructive impact in the real world and the SPM slowed down the hammer to rest. To demonstrate the protective properties of the SPM as a casing, an egg placed between two SPM cups and hammered several times remained intact after multiple impacts (Figure 9).

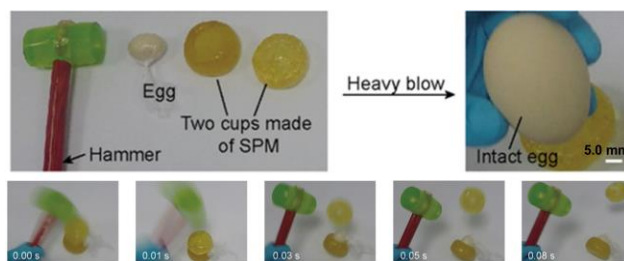


Figure 9. Hammer test on an egg encapsulated between two SPM cups. Energy dissipation by the SPM cups protected the structural integrity of the egg. Adapted with permission from ref. 55. Copyright 2020 American Chemical Society.

The role of multiple sacrificial bonds in IHPs was recently investigated by Zhang and coworkers.⁵⁴ The authors prepared a single IHP containing H-bonding UPy motifs, reversible boron-oxygen linkages, and Cu^{2+} -coordinating imidazoline ligands, as multiple energy-dissipating pathways (Figure 10a). To investigate the mechanism of energy

dissipation, loading-unloading cycles at different strain rates were performed. Samples showed greater hysteresis at higher strain rates, indicating higher energy dissipating capabilities. The authors proposed that at low strain rates, boron-oxygen bonds break, increasing the mobility of polymer chains within the network. At higher strain rates, chain motion becomes too fast for boron-oxygen bonds to break, causing chain entanglement, followed by the dissociation of HBs and metal coordination (Figure 10b). Further increasing the strain rate breaks the covalent bonds of the polymer backbone and leads to irreversible microcracks. To confirm that the sacrificial bonds break reversibly, the authors spliced the sample in two halves ($30 \times 10 \times 3$ mm) and subsequently kept them in close contact for 10 s at room temperature. These combined segments were able to withstand 400 g of external load. Hence, the authors concluded that the reported IHP showed effective impact-absorbing capabilities due to reversible dynamic bonds. This work highlights the ability of multiple non-covalent interactions to act in synergy to achieve impact resistance and, moreover, provides a rare example where the relative kinetics of strain rate and bond breaking lead to selectivity over which type of sacrificial bond is broken. Such control is desirable for envisaged applications including soft robotics and personal protective equipment.

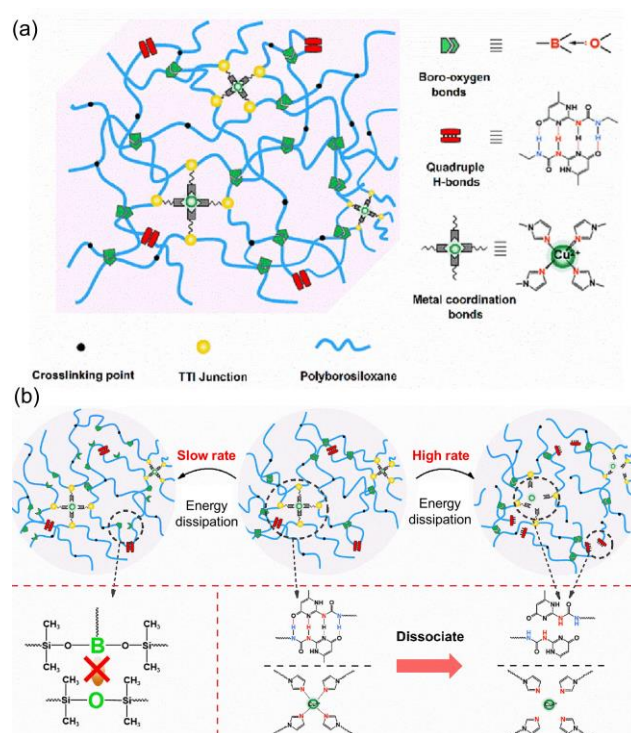


Figure 10. (a) Molecular structure of the reported impact hardening polymer (IHP) with trisubstituted triisocyanate junctions (TTI), multiple HBs, and metal-coordination sacrificial bonds. (b) Illustration of energy dissipation mechanism. Reprinted from ref. 54 with permission from Elsevier, copyright 2023.

In 2017, Ikkala and coworkers reported a similar strategy to enhance the mechanical and energy-dissipating properties of an acrylate-based polymer, but utilized three H-bonding motifs with different association strengths rather than three different bonding interactions.⁵⁶ Poly(2-hydroxyethyl methacrylate) (pHEMA) is known to have a brittle structure and poor mechanical properties despite having hydroxyl groups as HB acceptors and donors. The resultant HBs alone are not strong enough to provide toughness or dissipate substantial amounts of energy. To increase the HB density, UPy motifs were attached to a fraction of chains and the resulting copolymer, p(HEMA-*co*-UPyMA), displayed an array of single and quadruple HBs as interchain links (Figure 11). The strong UPy-mediated HBs slowed crack propagation and increased yield strength and Young's modulus, by 65% and 69%, respectively, indicating that these links act as sacrificial bonds to dissipate the crack energy. To further enhance the mechanical properties of the copolymer, carbon nanotubes (CNTs) were incorporated into the material. CNTs exhibit extraordinary thermal, electrical, and mechanical properties, and there have been several reports on harnessing these properties in the bulk.^{57,58} However, materials with CNTs incorporated non-covalently into soft materials had previously shown poor mechanical properties due to highly heterogeneous dispersion of CNTs in the polymer matrix.^{59,60} To further increase these properties of p(HEMA-*co*-UPyMA), oxidized multiwall carbon nanotubes (MWCNTs) were used as another crosslinker to strengthen the material. The surface of the oxidized CNTs presented UPy, -OH, and -COOH groups to facilitate interactions with the polymer chains (Figure 11). The resulting material showed increases in Young's modulus, tensile strength, and yield strength of 321%, 51%, and 100%, respectively. This report demonstrates that the incorporation of sacrificial bonds in nanocomposites can synergistically increase solid-state toughness and defect tolerance. This strategy of combining hierarchical HBs with CNTs and harnessing the properties of both materials mirrors natural approaches to composite materials with ultra-strong toughness and energy-dissipating properties.³ Jiang and coworkers also demonstrated the utility of UPy motifs as one of several sacrificial HB motifs in a polyacrylate hydrogel.⁶¹ Notably, the authors found that incorporating UPy at a concentration of just 0.8 mol% was sufficient to increase the Young's modulus from 74 ± 10 kPa to $1,252 \pm 108$ kPa, surpassing the mechanical properties of human tissues such as nasal cartilage (250 – 440 kPa)⁶² and skin (200 – 250 kPa).⁶³

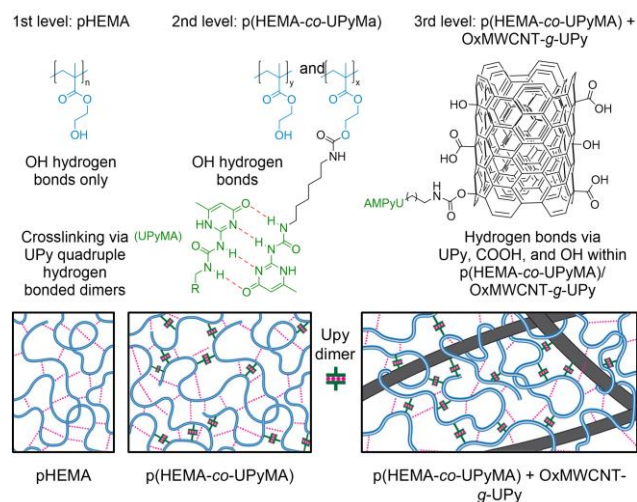


Figure 11. Different levels of hierarchical HBs in the supramolecular network. The first level of HBs is between the –OH groups of pHEMA chains (weak interactions). The second level of HBs is between the UPy motifs (strong interactions) and the last level is between p(HEMA-co-UPyMA) and OxMWCNT-g-UPy. Adapted with permission from ref. 56. Copyright 2017 American Chemical Society.

Hierarchical H-bonding was also explored by Sun and coworkers, who used carboxylic acids and UPy to crosslink polyurethane (PU) chains, supplementing the carbamate-based H-bonding inherent to the PU backbone.⁶⁴ To compare the relative contributions of each HB motif to the polymer's mechanical properties, three elastomers were prepared with varying molar ratios of UPy and DPA (3-hydroxy-2-(hydroxymethyl)-2-methylpropanoic acid), where DPA acts as a chain extender and provides a carboxylate H-bonding motif. The authors propose that weak carbamate-carbamate and carboxyl-carboxyl HBs dissociate first and contribute to the toughness of the elastomer, while the UPy-UPy dimer provides high strength (Figure 12a). The sample with the lowest UPy content (10 mol% UPy) exhibited low tensile strengths, presumably due to an insufficient number of strong HBs, while the sample with the highest UPy content (30 mol% UPy) showed an increased Young's modulus but had poor stretchability, attributed to an insufficient number of weak HBs. However, a sample with a UPy:DPA ratio of 20:80 exhibited a good balance of enhanced toughness and mechanical strength. This elastomer was characterized by cyclic loading-unloading tests at 400% strain. The first cycle showed a hysteresis loop with an area of 5.29 MJ/m³ that was significantly reduced in an immediate second cycle (no relaxation time) with 44.8% residual strain (Figure 12b). However, with increasing relaxation time, the residual strain increased and was indistinguishable from the first cycle after a relaxation time of 90 min (Figure 12c). This study demonstrates how mechanical properties can be tuned by tailoring the relative amounts of different sacrificial bonds as well as their relative interaction strengths.

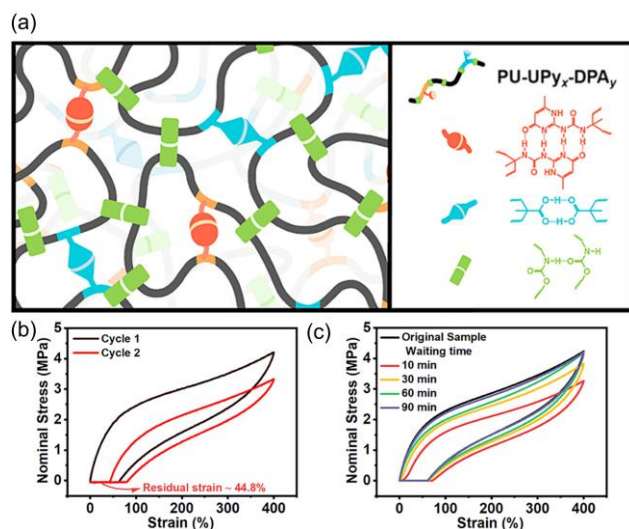


Figure 12. (a) Structural representation of PU-UPy_x-DPA_y polymer network and sacrificial HBs in the network. (b) Two successive cycles of cyclic loading-unloading test of the PU-UPy_{0.2}DPA_{0.8}. (c) Cyclic loading-unloading test with different relaxation times. Adapted with permission from ref. 64. Copyright 2023 American Chemical Society.

UPy motifs have also been explored as sacrificial bonds in elastomers, whose elastic properties make them excellent materials for applications in tires, sealants, protective coatings, and wearable electronics.^{65,66} Elastomers can tolerate large deformations and stress due to their elasticity, but they accumulate internal microcracks that expand over time and result in functional failure and material degradation. To address this problem, Cai and coworkers combined PU with UPy and imidazole motifs using thiolactone chemistry (Figure 13).⁶⁷ In brief, a thiolactone derivative, α -isocyanato- γ -thiolactone, was functionalized with pendent UPy and imidazole groups, and incorporated into PU chains via typical synthesis strategies using PTMEG and HMDI. Subsequent treatment with $\text{Zn}(\text{CF}_3\text{SO}_3)_2$ to form the metal-ligand bonds between Zn^{2+} and imidazole (Im), followed by molding and drying, afforded the polyurethane-based elastomer, PU-Im-UPy-Zn. A control sample prepared via a similar strategy, PU-Im-BA-Zn, replaced UPy with butyl acrylate to eliminate the H-bonding network. Rheological studies showed that the introduction of Zn^{2+} ions increased the storage modulus (G') two-fold compared to metal-free PU-Im-UPy and four-fold compared to the butyl acrylate control, demonstrating the synergistic cross-linking of UPy units and Zn^{2+} coordination with imidazole ligands. At higher strain rates, PU-Im-UPy and PU-Im-BA-Zn showed enhanced tensile strength but poor extensibility. In contrast, PU-Im-UPy-Zn showed enhanced toughness (62.1 MJ/m^3 , 136% vs control), tensile strength (9.1 MPa, 198% vs control), and extensibility, due to the rapid rearrangement of dynamic HBs and metal-ligand coordination bonds. This work highlights the ability of dynamic, rapidly reformed sacrificial bonds to bolster the mechanical properties of elastomers.

A 2019 report by Guo and co-workers showed that incorporation of UPy in poly(urea) elastomers improved their strength, stiffness, and toughness due to the formation of quadruple HBs.⁶⁸ Samples with UPy showed threefold higher toughness than the pristine sample. The dimerization of UPy has also been used to improve the toughness of polymer networks comprising poly(ether imide)⁶⁹ and other weak polymers.^{70,71}

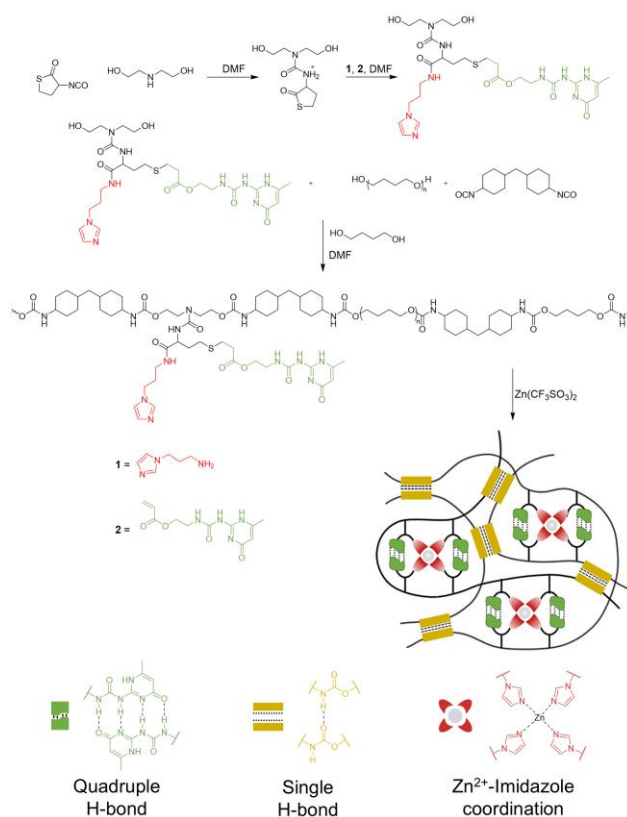


Figure 13. Synthetic route for PU-Im-UPy-Zn and sacrificial bonds involved in the network. Adapted with permission from ref. 67. Copyright 2023 American Chemical Society.

4.2. Amides

In biological systems, amides serve as the primary chemical motifs for HB formation: for example, proteins have hierarchical structures scaffolded by amide-carbonyl, amide-water, and carbonyl-water H-bonding interactions. These interactions provide structural stability to the proteins and ensure proper functioning.⁷² Accordingly, amides have been explored extensively as H-bonding motifs in synthetic polymeric materials because of their biocompatibility, straightforward synthesis, and potential to enhance mechanical and energy-dissipating properties, as discussed in an excellent recent review by Li *et al.*⁷³

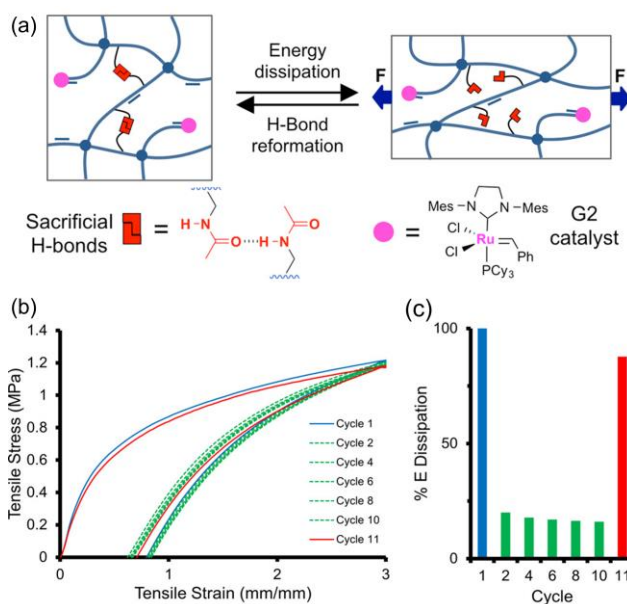


Figure 14. (a) Schematic illustration of reversible sacrificial HB breaking under external force. (b) Hysteresis loops for 11 loading-unloading cycles of ACON at 300% strain. The first ten cycles were done immediately and the last was performed after 30 min of relaxation time. (c) Energy dissipation corresponding to each cycle. Adapted with permission from ref. 45. Copyright 2015 American Chemical Society.

In 2015, Guan and co-workers combined amide functionality with a covalent self-healing polymeric network and observed enhancements in mechanical and energy-dissipating properties.⁴⁵ Specifically, secondary amide groups were attached to a cyclooctene-based polymer backbone which was cross-linked and appended with Grubbs' second-generation ruthenium catalyst (G2), to give a final material (amine-containing cyclooctene network, ACON) with both covalent and non-covalent cross-links. The HB cross-links broke under stress and provided toughness to the material, while the G2 catalyst mediated dynamic olefin metathesis at the fracture interface in the case of mechanical damage to the covalent network (Figure 14a). A control material in which the amide hydrogen was replaced with a methyl group (blocked amide-containing cyclooctene network, BACON) exhibited decreased tensile toughness (7-fold) and strain-at-break (6-fold), confirming the importance of the amide-mediated H-bonding. The reversible nature of amide sacrificial bonds was demonstrated via cyclic tensile tests, in which BACON showed an extensibility of 167% while ACON showed up to 300%. At first, 10 immediate cycles of stress loading and unloading were performed. The first cycle showed large energy dissipation, but this decreased by 80% in the second cycle and decreased further over subsequent cycles. An 11th cycle was performed after 30 min of relaxation time in ambient conditions, and ACON exhibited 90% of the energy dissipation observed in the first cycle (Figure 14b,c). This study validated amides as an effective H-bonding motif to improve the energy-dissipating properties of synthetic polymer networks, demonstrated by a battery of mechanical testing studies.

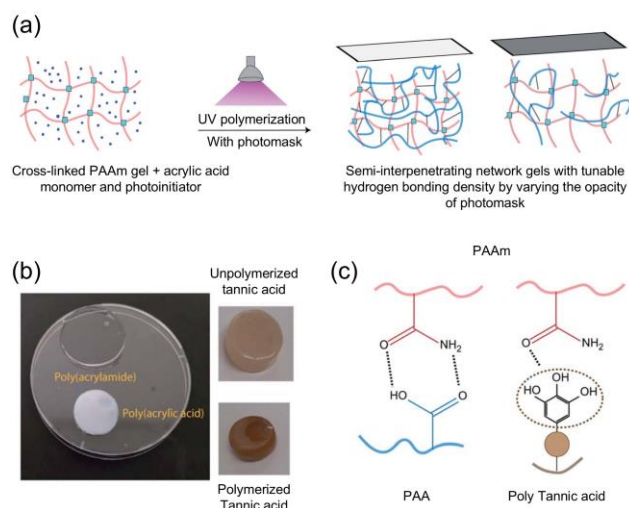


Figure 15. (a) Photopolymerization of monomers (acrylic acid, acrylamide or tannic acid) in a cross-linked PAAm network using UV light. The HB density can be easily tuned by varying the intensity of UV light with different photomasks. (b) Macroscale samples of double network hydrogels. (c) H-bonding interactions between PAAm and poly(acrylic acid) or poly(tannic acid). Figure adapted from ref. 74 with permission from the Royal Society of Chemistry, copyright 2021.

Hydrogels are particularly attractive in applications like tissue engineering due to their high modulus, softness yet toughness, resistance to mechanical shear, and biocompatibility.^{75,76} Early work by Sheiko and coworkers reported that hydrogels formed by copolymerizing *N,N*-dimethylacrylamide and methacrylic acid dissipate large amounts of energy (1.7 MJ/m^3) and exhibit a near-complete recovery of dissipative capacity after a relaxation time of 60 min.⁷⁷ Later work by Malmström and coworkers achieved tunable energy dissipation in synthetic hydrogels.⁷⁴ Three different semi-interpenetrating double networks were made by photopolymerizing two different monomers (acrylic acid and tannic acid) in cross-linked poly(acrylamide) (PAAm) hydrogels (Figure 15a, b). The two different monomers had unique H-bonding capabilities when copolymerized with the PAAm network (Figure 15c). Poly(acrylic acid) (PAA) adopts a charged state at physiological pH and formed complexes (via H-bonding) with PAAm. Poly(tannic acid) (PTA) is capable of UV polymerization, has multiple HB sites, and forms shorter, more highly branched chains compared to PAA. In comparison to unmodified PAAm, in which limited H-bonding is expected, both PAAm-PTA and PAAm-PAA showed increased energy dissipation as reflected by improved mechanical properties: PAAm-PTA showed a 19-fold increase in tensile strength and a 55-fold increase in toughness, while the PAAm-PAA hybrid exhibited 4.3- and 3.6-fold improvements, respectively. DLS studies indicated that PAAm-PTA formed larger aggregates than PAAm-PAA, which might account for the increase in toughness. The reported hydrogels have similar mechanical properties to soft tissues and comparable relaxation

times to skin. Such materials could be used to explore hydrogel applications in cellular mechanotransduction studies.

As noted earlier, polyurethanes are desirable materials whose toughness is limited due to their low inherent H-bonding strength. In contrast to Cai and coworkers' study of UPy-functionalized polyurethanes (Section 4.1),⁶⁷ Liu and coworkers employed an alternative approach that utilized H-bonding chain extenders made from the dipeptide *N*-acryloyl glycinamide (NAGA) (Figure 16).⁷⁸ These extenders were reacted with copolymers of PTMEG and either hexamethylene diisocyanate (HDI) or IPDI to generate PUHD-NAGA and PUIP-NAGA, respectively. The aim was to supplement the main chain amide HBs with the amide HBs of the NAGA extenders to make a mechanically stiff, tear-resistant, self-healing material. By varying the molar ratio of NAGA units to PTMEG segments, different mechanical properties could be optimized. For example, PUIP-NAGA with a NAGA:PTMEG ratio of 2.1 showed the highest energy dissipation (2.52 MJ/m³ at 200% strain), while a thin-film sample of PUIP-NAGA with a NAGA:PTMEG ratio of 2.6 could hold up to 4 kg of weight with minimal deformation.

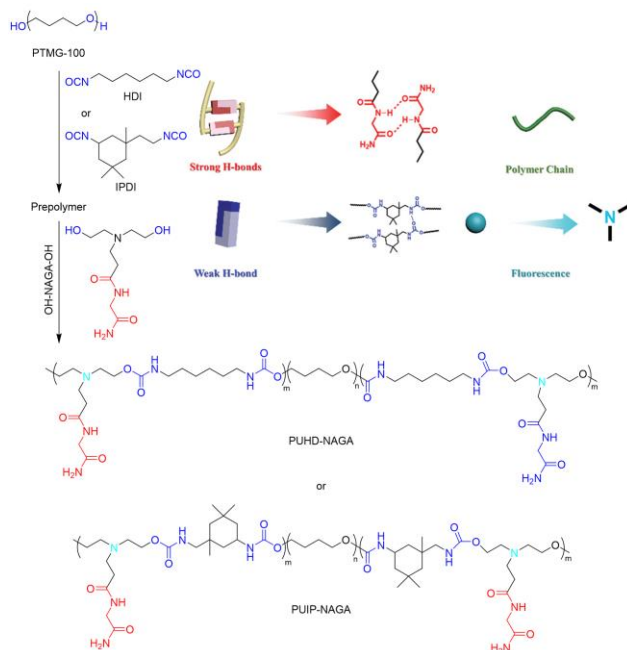


Figure 16. Synthetic schemes for PUHD-NAGA and PUIP-NAGA PU elastomers. H-bonding in the polymer network comprises two different types of HB: strong HBs between NAGA motifs and weak HBs between amides in the polyurethane backbone. Figure adapted from ref. 78. Copyright 2020 Wiley-VCH GmbH.

To achieve high toughness and tensile strength, one of the most common strategies is the crosslinking of polymeric chains, but this can cause other issues like poor reprocessibility and difficulties in degradation, which create environmental issues. Modulating the number of amide HBs was also recently explored by Shi and co-

workers in the context of curable epoxy materials based on polyurethanes.⁷⁹ The authors synthesized epoxy resins with different H-bonding capabilities by curing an epoxy-terminated monomer (IDV-EP) with three different bisamines (Figure 17a): diamino-diphenylmethane (DDM), which introduces the fewest HBs; 2-(4-aminophenyl)-*1H*-benzimidazol-5-amine (APBIA), which introduces an imidazole moiety; and 4,4-diaminobenzoyl triphenylamine (DABA), which introduces an amide motif. All three samples surpass a commercially available resin (diglycidyl ether-bisphenol A, DGEBA) in terms of mechanical properties. Notably, IDV-EP-DABA showed a tensile strength of 90 MPa and toughness of 9.73 MJ/m³, which are 1.8 and 6.7 times higher than DGEBA/DDM (Figure 17b). Moreover, the reported epoxy materials showed complete degradation when exposed to mixture of 0.1 M HCl and THF for 24 h at room temperature, opening a potential route to reprocessing.

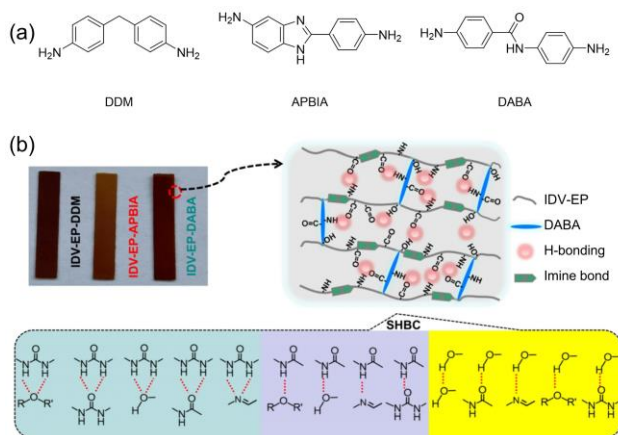


Figure 17. (a) Three different curing agents used for IDV-EP. (b) Sacrificial HBs between alcohol, amide, urea, and imine functionalities in IDV-EP polymers. Reprinted from ref. 79 with permission from Elsevier, copyright 2024.

Ureas offer many of the same benefits of amides for sacrificial bond formation due to their HB-accepting carbonyl oxygen atom and the presence of two N–H bonds, compared to the single HB donor in a traditional amide. Consequently, ureas and related structures have been utilized extensively in tough materials. Zhang and co-workers recently reported a poly(urethane-urea) based elastomer with HB-rich macrocyclic 2D polyamide (2DPA).⁸⁰ 2DPA was synthesized from melamine and 1,3,5-benzenetricarbonyl trichloride and was subsequently integrated into a soft matrix of poly(urethane-urea) (Figure 18a,b). Varying the proportion of 2DPA (1, 2, 3, or 4 wt%) gave elastomers with different numbers of sacrificial HB cross-links between the urea and urethane functional groups of poly(urethane-urea) and the amide groups of 2DPA. The pristine polymer network was soft and weak with a tensile strength of 1.1 MPa, toughness of 5.8 MJ/m³, and 641% elongation at break. All samples with 2DPA outperformed the pristine sample; the sample with 3 wt% 2DPA (PITAT@2DPA3) showed the highest tensile strength (54.6 MPa)

and toughness (116.7 MJ/m^3) with an excellent elongation at break (705%). Additionally, the reported elastomer can lift a mass of 10 kg, 40,000 times its own weight. In cyclic tensile tests, the sample with 3 wt% 2DPA showed increased energy dissipation with increased strain (strain increased from 50% to 400%, Figure 18c). When the specimen was subjected to five consecutive cycles at a constant strain of 200%, all five cycles showed energy dissipation (denoted by the hysteresis loop area) with substantial loss after the first cycle but minimal loss thereafter (Figure 18d). A recovery time of 3 h was sufficient to restore energy dissipation almost completely, to 95.6% of its original value (Figure 18e).

Sun and coworkers made a similar poly(urethane-urea)-based elastomer containing poly(caprolactone) (PCL) segments (Figure 18f).⁸¹ The reported elastomer has multiple cross-linking HBs between urethane and urea-containing acylsemicarbazide (ASC) groups, achieving a tensile strength of 72.6 MPa and 1440% of elongation at break with extremely high toughness of 375 MJ/m^3 . Wang and coworkers used a similar strategy with IPDI and PCL to synthesize an elastomer with HBs between the hard and soft segments.⁸² Their supramolecular poly(urethane-urea) showed a tensile strength of 64 MP and 2160% of elongation at break along with robust energy dissipation properties.

The urea moiety has been derivatized to further expand the palette of amide-like HB motifs for sacrificial bonds. Zhang and co-workers employed imidazolidinyl urea (IU) as an HB motif in poly(urethane)s based on PCL and hexamethylene diisocyanate (Figure 18g, h).⁸³ The resulting polymer exhibited a toughness of 168.2 MJ/m^3 . Cyclic tensile strength results showed that HBs in IU dissipate energy during stretching. The mechanical properties could be further enhanced by adding cellulose nanocrystals as additional HB partners.⁸⁴ Similarly high densities of HBs were achieved in thermoplastic PU elastomers by Cai and co-workers.⁸⁵ Like typical urea, thioureas have also shown promise as sacrificial bond motifs due to their ability to engage in H-bonding between amide-like N-H bonds and thiocarbonyl sulfur atoms. Additionally, thioureas can form dynamic covalent bonds due to the reversible nature of the thiourea linkage. Kim and coworkers leveraged both of these binding modes in a biomimetic polymeric network using thioureas as a key sacrificial motif.⁴² The H-bonding capabilities of thiourea enhanced the polymer's mechanical properties (maximum strain = 360%, toughness = 71 MJ/m^3) while its dynamic covalent character improved malleability and processibility.

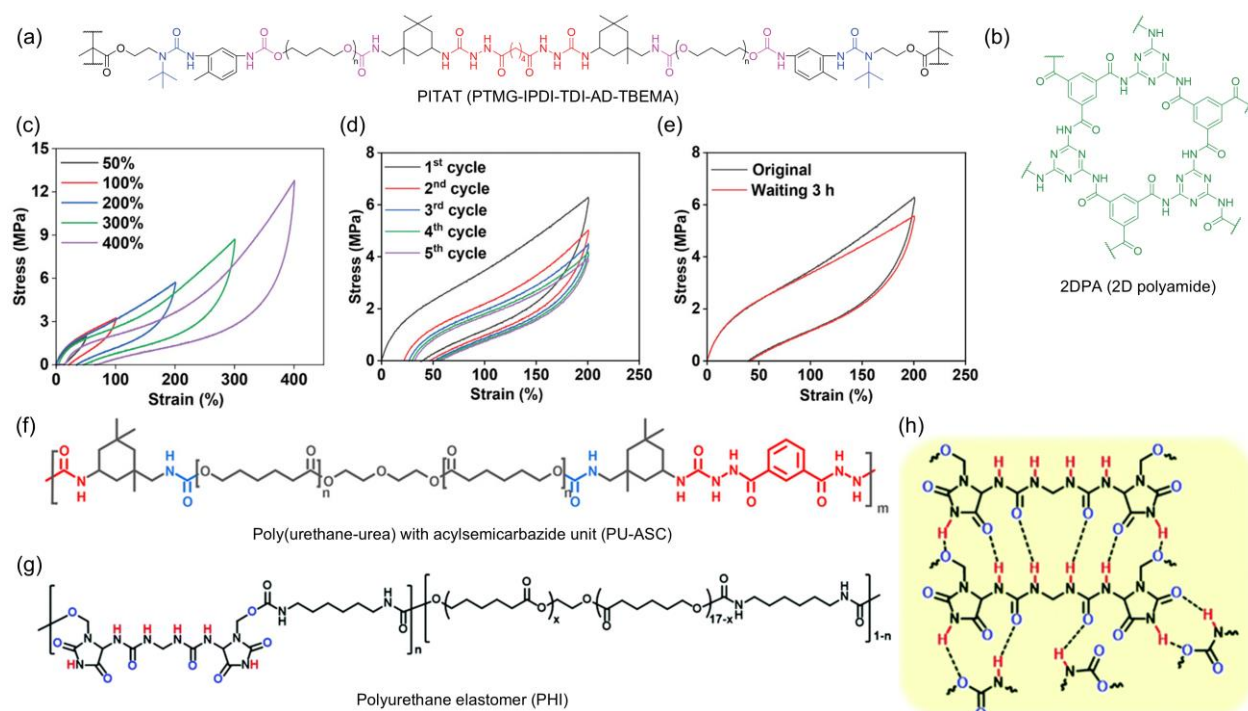


Figure 18. a) Chemical structures of poly(urethane-urea) made from polytetrahydrofuran (PTMG), isophorone diisocyanate (IPDI), toluene-2,4-diisocyanate (TDI), and 2-(*tert*-butylamino)ethyl methacrylate (TBEMA). Different H-bonding sites are represented by different colors. (b) Chemical structure of 2D polyamide (2DPA). (c–e) Cyclic tensile test results for PITAT@2DPA3 with varying (c) strain, (d) cycle number at a constant strain of 200%, and (e) recovery time. Figure adapted from ref. 80. Copyright 2024 Wiley-VCH GmbH. (f) Chemical structure of poly(urethane-urea) with acylsemicarbazide unit (PU-ASC). Adapted with permission from ref. 81. Copyright 2022 American Chemical Society. (g) Chemical structure of PU with imidazolidinyl urea (IU), hexamethylene diisocyanate (HDI), and polycaprolactone (PCL). (h) H-bonding cross-links due to the presence of IU. Figure adapted from ref. 83 with permission from the Chinese Chemical Society (CCS), Institute of Chemistry of Chinese Academy of Sciences (IC), and the Royal Society of Chemistry, copyright 2021.

4.3. Alcohols

Unlike ureidopyrimidinones and amides, in which oxygen atoms act solely as HB acceptors, alcohols include a polar O–H bond that serves as an HB donor. This labile hydrogen atom, in addition to the HB acceptor ability of the oxygen atom, makes alcohols privileged motifs capable of engaging in multiple HBs simultaneously. This ability is leveraged throughout natural materials, from the molecular recognition of sugars in the glycocalyx⁸⁶ and the structural integrity of polysaccharides such as cellulose,⁸⁷ to the strong adhesive properties of polyphenols such as catechols even in challenging aqueous environments.⁸⁸ Unsurprisingly, materials scientists have sought to leverage the O–H group as a motif for sacrificial bonds in synthetic systems too.

In 2020, Chen and coworkers reported hydrogels based on poly(vinyl alcohol) (PVA) and cellulose nanocrystals (CNC) with tannic acid (TA), a large polyphenol, as a H-bonding crosslinker (Figure 19).⁸⁹ Cyclic tensile tests showed the expected increase in the area of the hysteresis loop (i.e., dissipated energy) in the presence of HB-

mediated crosslinking between TA and PVA/CNCs. Notably, a sample with 15 wt% of TA and 20 wt% of CNC (both with respect to PVA) exhibited increased amounts of energy dissipation with increasing strain levels (0.96 MJ/m³ at 100% strain and 11.89 MJ/m³ at 600% strain). When this sample was subjected to multiple loading-unloading cycles (at 200% strain) with no relaxation time, a significant decrease in the hysteresis area was observed, suggesting that the H-bonds and PVA crystalline domains were broken. Re-testing after 180 min of relaxation time caused the hysteresis loop to recover to almost 60% of its original size.

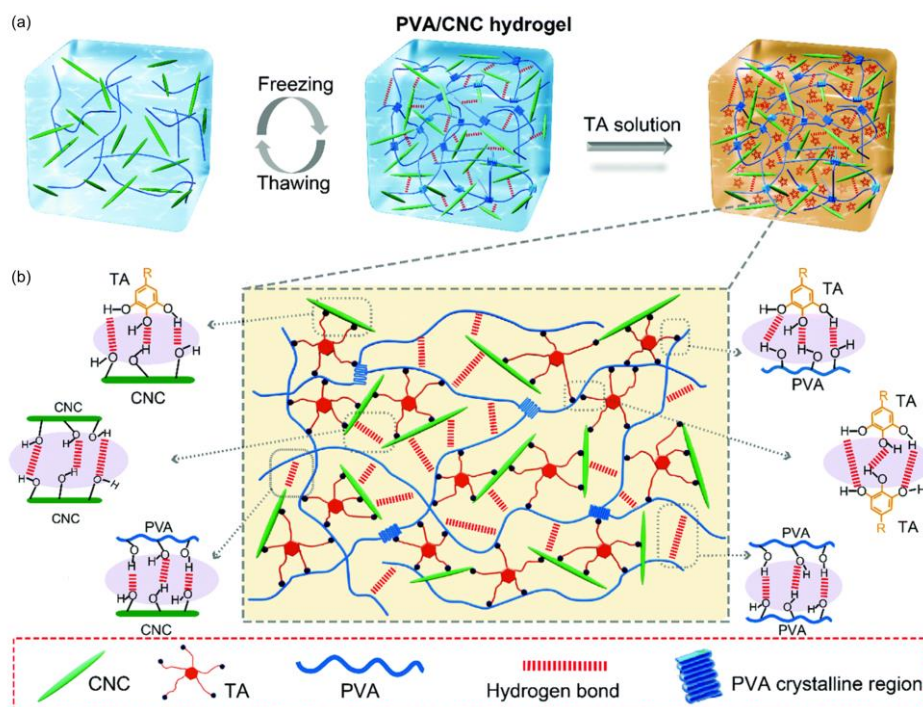


Figure 19. (a) Fabrication process of TA-PVA-CNC hydrogels. (b) Sacrificial HBs in the hydrogel network. Figure adapted from ref. 89 with permission from the Royal Society of Chemistry, copyright 2020.

In 2021, Duan and coworkers reported a strategy to make tough hydrogels using chitin and polyphenols.⁹⁰ Their approach employed two separate non-covalent networks: a strong and rigid non-covalent network to achieve high mechanical performance and a weak non-covalent network for efficient energy dissipation. Chitin acted as the strong network because of the extensive H-bonding, hydrophobic, and ionic interactions between its polymer chains, while the weak and reversible interactions between polyphenols defined a second, weaker network. Of several polyphenols screened for hydrogelation with chitin (Figure 20a), TA showed the highest compressive strength (8 times higher than that of chitin alone, Figure 20b). The authors attribute this strength to the large number of weak chitin-TA interactions acting as sacrificial bonds under stress loading.

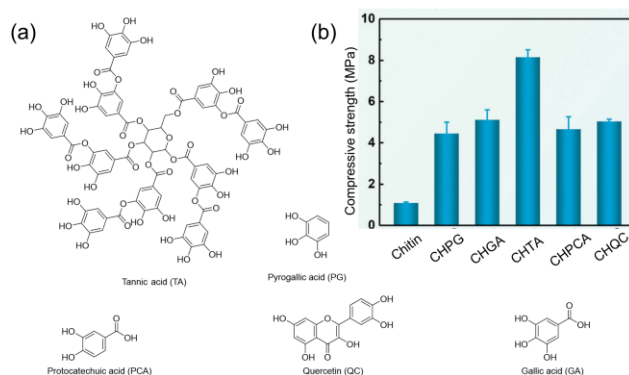


Figure 20. (a) Different polyphenols used to make chitin-polyphenol. (b) Comparison of compressive strength of chitin and chitin-polyphenol hydrogels (CH: chitin, PG: pyrogallol acid, GA: gallic acid, TA: tannic acid, PCA: protocatechuic acid, and QC: quercetin). Figure adapted from ref. 90 with permission from the Royal Society of Chemistry, copyright 2021.

In 2022, Zhou and coworkers reported an unusual strategy to build sacrificial bonds by incorporating a molecular solvent into a double network-based hydrogel (Figure 21).⁹¹ The double-network structure, termed PAAN, comprised interpenetrated polyaniline and hydrophobic-associating poly(acrylic acid) (HAPAA). Sacrificial bonds were introduced by incorporating glycerol into the PAAN gel using a solvent exchange method. The glycerol content in the resultant organohydrogels was varied by immersing PAAN samples in glycerol for different lengths of time. Increasing the immersion time enhanced the mechanical properties of the organohydrogels, which showed a fracture energy ~ 4 times higher than that of human skin (~ 3.6 kJ/m²) and ~ 14 times that of cartilage (~ 1 kJ/m²).^{92,93}

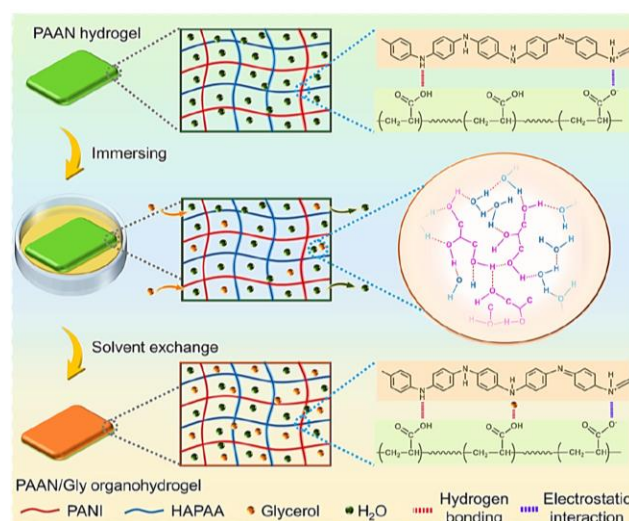


Figure 21. Chemical structure and preparation scheme for the PAAN/Gly organohydrogels with different types of sacrificial bonds involved in the hydrogel. Adapted with permission from ref. 91. Copyright 2022 American Chemical Society.

Several recent reports have also examined alcohols as HB motifs for sacrificial bonds. Zhang and coworkers showed that molecules with multiple –OH groups, such as tannic acid, can act as H-bonding bridging agents between the –OH groups of natural bamboo fiber and the carbonyl groups of the synthetic polymer poly(butylene succinate-*co*-butylene adipate) (PBSA).⁹⁴ The resulting biocomposite showed a 22% increase in tensile strength compared to the composite without tannic acid. Karak and coworkers reported a poly(ester-amide-urethane) network with different ratios of IPDI, leading to excellent elongation at break values (641% to 1876%) and tensile strengths (1125 to 1896 MJ/m³).⁹⁵ Leng and coworkers explored the role of H-bonding on the mechanical properties of cross-linked epoxy resins.⁹⁶

5. Metal Coordination

Metal-ligand interactions are formed when a Lewis base (ligand) donates electrons to a Lewis acidic (metal) center. Interestingly, the dynamic properties of resultant metal complexes can range anywhere between covalent and supramolecular systems depending on the specific metals and ligands used: in some cases, a ligand formally donates a pair of electrons to the metal center, making the resultant bond stronger than strictly⁹⁷ non-covalent interactions such as H-bonding. Furthermore, multidentate ligands introduce the potential for multiple interactions between ligands and the same metal center, in both intramolecular and intermolecular fashions, providing additional options for tuning the overall strength of their interactions. Indeed, mussels' unique ability to attach themselves to solid surfaces and avoid being swept off by intertidal waves is attributed to the polydentate coordination between iron and DOPA molecules in mussel cuticles.⁹⁸ Moreover, metal-ligand systems often possess fast bond exchange kinetics that makes them more versatile than similarly strong covalent systems.¹⁶ These properties, coupled with the vast chemical space of metal-ligand systems, have spurred substantial exploration of metal coordination chemistry in the context of self-healing polymers,^{99–104} and make such bonds attractive candidates to serve as programmable sacrificial bonds.

Inspired by the adhesive mechanism of mussels (*vide supra*),^{98,105} several groups have explored iron-catechol interactions as sacrificial bonds. Waite and coworkers examined the role of pH on iron-catechol interactions in PEG-based materials.¹⁰⁶ Upon deprotonation of the acidic catechol protons, the resulting catecholate motifs formed bidentate coordinate bonds to the iron center that could then serve as sacrificial bonds to enhance the polymer's mechanical properties (Figure 22a). The authors noted that a significant challenge in this process was that the

alkaline pH necessary for bis- and tris-complex formation between Fe^{3+} ions and catechol made the metal ions insoluble, but this was overcome by slowly adjusting the system's pH. Under acidic conditions, Fe^{3+} ions were primarily coordinated by a single catechol motif (Figure 22b, green). Increasing the pH favored additional catechol coordination, and at pH 12, complete conversion to the tris-coordinated complex was achieved, similar to that observed in mussel cuticles (Figure 22b, red). The authors compared the mechanical properties of this modified polymer with a control polymer without catechol motifs or metal-ligand interactions. Although both samples showed similar hardness (comparable G' values) under low strain values, the iron-catechol-modified gel dissipated 10-fold more energy (G'') compared to the control gel. This study validates the energy dissipating potential of metal-ligand bonds. Using similar coordination chemistry, Holten-Anderson and coworkers reported a strategy to decouple spatial and temporal hierarchy in a material.¹⁰⁷ By establishing different metal-based coordination networks in the same polymeric matrix, the authors were able to design the hierarchy of sacrificial bond rupture. This allowed the authors to program the viscoelastic properties of the bulk polymeric material.

Valentine and co-workers have noted that, in soft, wet systems such as hydrogels, high stiffness and high maximum elongation strain are often orthogonal and cannot be achieved simultaneously.¹⁰⁸ To circumvent this problem, they designed a dry polymeric system in which sacrificial metal-coordination networks yield a material with both high stiffness and extensibility. The covalent polymer network was formed by copolymerizing a short-chain bisepoxide (poly(ethylene glycol) diglycidyl ether, PEG-DE), a monoepoxide carrying a silyl-protected catechol, and a diamine crosslinker (Figure 22c). Subsequent deprotection of the silyl groups to reveal the catechol motifs, followed by chelation of Fe^{3+} ions and drying, produced the crosslinked metal-coordination network (Figure 22d). To quantify the enhancement of mechanical properties upon the introduction of iron-catechol coordination bonds, uniaxial tensile tests were used to compare silyl-protected polymers (no Fe^{3+} -OH interactions) with deprotected polymers (with free catechols capable of metal coordination): the latter were shown to exhibit a 770-fold increase in elastic modulus, a 58-fold increase in tensile strength, a 76-fold improvement on yield stress and a 92-fold increase in tensile toughness (Figure 22e). These results provide a seminal example of metal-ligand coordination being successfully deployed in dry polymer materials.

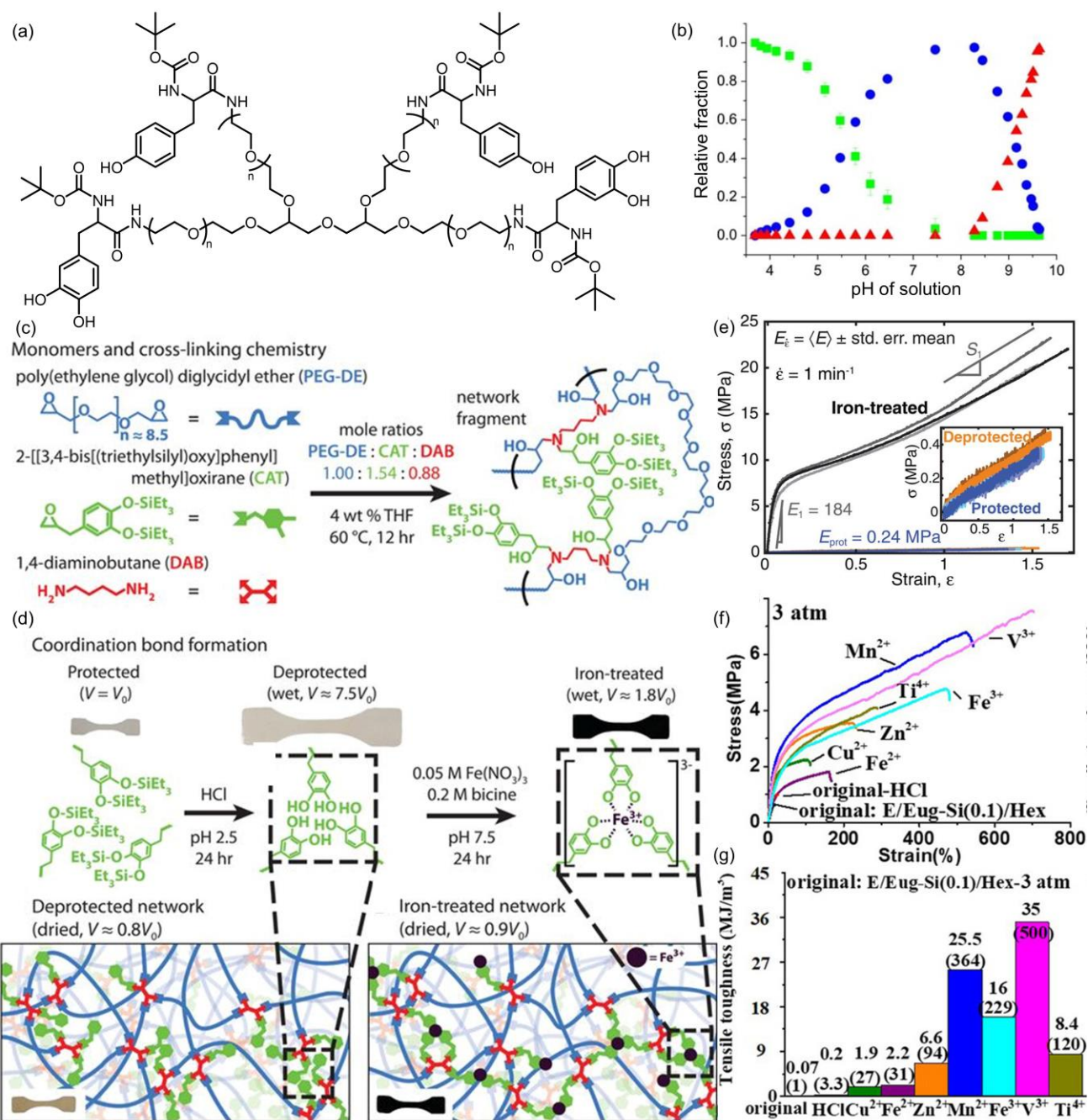


Figure 22. Metal-catechol coordination interactions as sacrificial bonds to improve mechanical properties. (a) DOPA-modified PEG units containing catechols for iron coordination. (b) Relative fractions of mono- (green), di- (blue) and tris- (red) coordinated Fe^{3+} in solution of PEG-dopa₄ at different pH values. Figures adapted from ref. 106. (c) Synthesis of catechol-containing PEG polymers. (d) Scheme showing preparation of dry, iron-containing catechol-PEG networks. (e) Uniaxial tensile tests demonstrating the impact of metal coordination on mechanical properties of catechol-PEG polymers. Figures adapted from ref. 108. Reprinted with permission from AAAS. (f) Stress-strain curves showing different mechanical properties of eugenol-based polymers. (g) Comparing toughness of different polymers with different metals or HCl in eugenol-based polymers. Figures adapted from ref. 109. Copyright 2020 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.

The coordination of metal cations beyond Fe^{3+} was employed by Na and coworkers in their study of eugenol-based polyolefins.¹⁰⁹ These polymers were prepared via copolymerization of a silyl-protected version of eugenol, a

low cost and renewable allylic alkene, and an alkene, followed by acid-catalyzed hydrolysis of the silyl ether protecting group to enable metal coordination. This increased coordination of Fe^{3+} -catechol sacrificial bonds led not only to an 8-fold increase in tensile strength and 28-fold increase in strain-at-break, but also a remarkable 229-fold increase in tensile toughness when compared to the silyl-protected polymer with no metal-coordination bonds. The authors also found that increasing the chain length of the alkene component from ethene to 1-hexene led to increased flexibility and hence better sacrificial capacity of the iron-catechol coordination. The authors also investigated the effect of replacing iron with other metals. Mn^{2+} and V^{3+} were found to show even better property enhancement compared to Fe^{3+} , with V^{3+} showing a 500-fold increase in tensile toughness compared to the coordination-free silyl-protected network (Figure 22f,g).

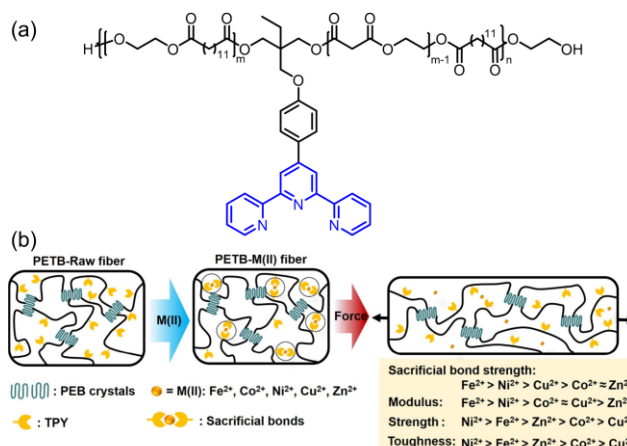


Figure 23. Effect of different metal cations on sacrificial bond strength. (a) Structure of polyester that incorporates pendent tridentate terpyridine ligands (shown in blue). (b) Schematic depiction of the sacrificial nature of the metal coordination network and the relative effect of different metals on modulus, tensile strength and toughness. Adapted with permission from ref. 110. Copyright 2024 American Chemical Society.

Recent work by Chen and coworkers has further explored how the identity of the metal component in a metal-ligand sacrificial bond can be changed to tune the mechanical properties of polymers (Figure 23).¹¹⁰ Specifically, polyesters containing pendent terpyridine (TPY; shown in blue) groups, which serve as high-affinity tridentate ligands (Figure 23a), were doped with a series of divalent transition metals (Fe^{2+} , Co^{2+} , Ni^{2+} , Cu^{2+} , Zn^{2+}) (Figure 23b). Interestingly, the identity of the metal cation did not affect the modulus, strength, and toughness of the resultant materials equivalently. The authors found that the enhancement in Young's modulus of the Fe^{2+} -polymer compared to the undoped polymer (3.3-fold), consistent with the coordination of Fe^{2+} by TPY being the highest of the metals investigated. However, the enhancement in maximum elongation was higher for Ni^{2+} compared to Fe^{2+}

(Figure 23b), which the authors attribute to the inability of the strong Fe^{2+} -TPY coordination to break before other interactions are (irreversibly) disrupted.

Metal-coordination sacrificial bonds have also been deployed to improve the mechanical properties of rubbers.^{111–113} Zhang and coworkers used the coordination of iron clusters by free hydroxy groups to reinforce rubber.¹¹⁴ Hydroxy groups were introduced into solution-polymerized styrene-butadiene rubber (SSBR) via thiol-ene click chemistry, after which the material was soaked in a FeCl_3/THF solution to incorporate the metal ions. TEM studies on model compounds suggest that Fe^{3+} ions form clusters of iron-rich segments instead of a homogenous solution. Hence, the resultant material had sulfur crosslinks, hydroxy-based H-bonding and hydroxy-based coordination with Fe^{3+} ions in the iron cluster (Figure 24a). The presence of Fe^{3+} ions led to a marked increase in mechanical properties, with an almost 7-fold increase in tensile strength and 9-fold increase in maximum elongation strain compared to unadulterated SSBR (Figure 24b). Furthermore, metal coordination enhanced the extent of energy dissipation, as demonstrated by cyclic tensile loading experiments. While the unadulterated polymer and metal-free hydroxy-containing polymer show only 12% and 21% relaxation of the initial stress, respectively, Fe^{3+} -doped polymers showed 71% relaxation of the initial stress. These enhancements were achieved with only 4.2 wt% iron, whereas achieving this strength in SBR modified by other methods would require a much higher content of fillers.¹¹⁵ Moreover, control experiments without metal ions showed limited improvement in tensile strength and modulus compared to unadulterated SBR, suggesting that the hydroxy-based H-bonding alone does not have a drastic effect on mechanical properties, thereby validating the efficacy of metal coordination as a sacrificial bond strategy in rubbers.

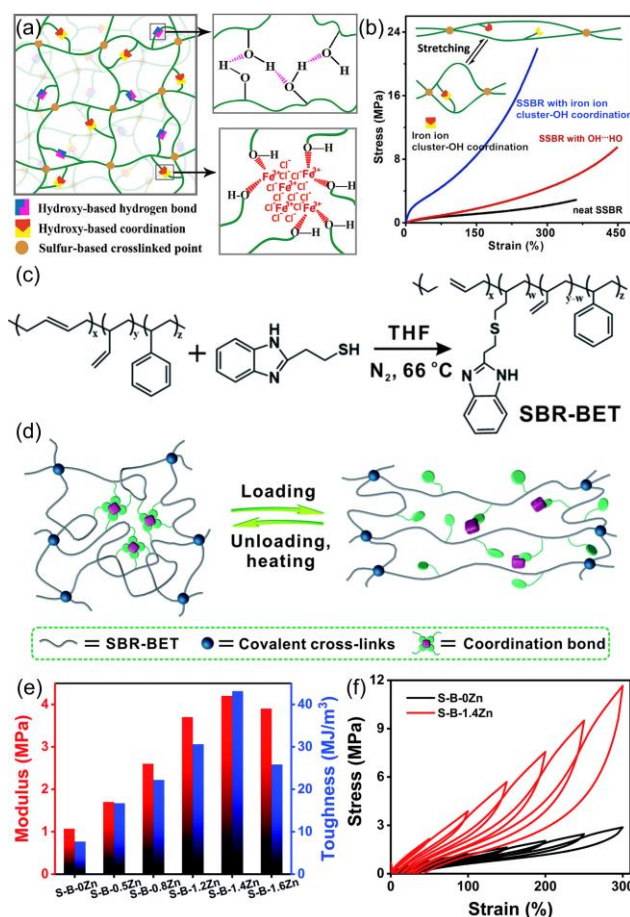


Figure 24. Modification of rubbers using metal-ligand coordination interactions as sacrificial bonds. (a) Scheme showing the formation of cluster Fe³⁺-OH coordination bonds (red) and HBs (magenta) in solution-polymerized styrene-butadiene rubber (SSBR). (b) A stress-strain graph shows much higher yield stress in the presence of metal coordination in iron cluster-based SSBR. Reprinted from ref. 114 with permission from Elsevier, copyright 2020. (c) Formation of SBR with dangling benzimidazole ligands (SBR-BET). (d) Scheme showing sacrificial nature of the Zn²⁺-imidazole coordination network under loading and unloading conditions. (e) Change in Young's modulus and tensile toughness under different concentrations of Zn²⁺ ions in SBR-BET. (f) Cyclic tensile loading tests show much higher dissipation of energy in the presence of the Zn²⁺-imidazole coordination network in SBR-BET. Figure adapted from ref. 116 with permission from the Royal Society of Chemistry, copyright 2019.

Liu and coworkers reported Zn²⁺-imidazole coordination as an alternative sacrificial network in SBR.¹¹⁶ Their study utilized SBR crosslinked with sacrificial boronic ester-based dynamic covalent bonds to impart vitrimer-like properties to the rubber, rendering it melt-processible. However, vitrimers usually lack mechanical toughness, owing to their ability to melt at higher temperatures. To address this problem, 2-(2-benzimidazolyl)ethanethiol was incorporated during the thiol-ene click reaction to functionalize the SBR vitrimer with pendent benzimidazole motifs (Figure 24c). Upon the introduction of Zn²⁺ ions, the benzimidazole motifs serve as ligands to form a sacrificial coordinate network (Figure 24d), which not only provides added mechanical toughness to the mechanically weak vitrimer but also imparts creep resistance: the authors report a maximum 4.6-fold increase in tensile toughness and a 5.4-fold increase in Young's modulus, compared to the Zn²⁺-free polymer, without

compromising the maximum elongation of the material (Figure 24e). Cyclic tensile loading-unloading experiments showed an increase in hysteresis loop area, characteristic of greater energy dissipation, in the Zn^{2+} -modified polymer versus the unmodified polymer (Figure 24f). Other groups have employed similar strategies, using pyridine as an alternative ligand in modified SBR¹¹⁷ and Zn^{2+} -carboxylate interactions to reinforce isoprene rubber.¹¹⁸ Notably, Weder and coworkers achieved an 81-fold improvement in toughness over neat SBR using pendent 2,6-bis(1'-methylbenzimidazolyl)-pyridine groups to coordinate Zn^{2+} ions.¹¹⁹ Recently, Shangguan and coworkers have demonstrated the mechanical toughening of SBR vulcanizates by incorporating sacrificial Zn^{2+} -pyridine coordination linkages.¹²⁰

As noted earlier, hydrogels are another materials class with many desirable properties, such as shape retention and ease of formation, that often have weak mechanical properties, in this case due to their high content of liquid water. Employing metal-coordination sacrificial bonds in hydrogels requires additional considerations due to the ability of water to bind competitively to metal ions, typically necessitating multidentate or cooperative ligands to successfully make mechanically tough hydrogels.^{121–124} Recently, Li and coworkers have reported a polyacrylamide hydrogel with isotropically-distributed deacetylated chitin nanowhiskers (ChNWs).¹²⁵ The monomers within these ChNWs are comprised of pyranose rings decorated with free amine and hydroxyl groups that can form a H-bonding network or, upon introduction of Zn^{2+} ions, form a multidentate metal coordination network (Figure 25a). The metal-doped material can be stretched up to 13 times its original length, and can also be bent, twisted, and even knotted to give different complex shapes (Figure 25b). Furthermore, increasing the concentration of the ChNWs and Zn^{2+} ions led to increasing tensile toughness, compared to unadulterated polyacrylamide gels or chitin-modified gels without Zn^{2+} (Figure 25c). These results validate that the energy dissipation behavior can be attributed primarily to the rupture of the sacrificial metal coordination network. The authors also noted that, at higher stress, the relative viscous character of the metal-coordinated material was higher than the unadulterated polyacrylamide gels, due to the more fluid-like behavior caused by sacrificial bond breaking.

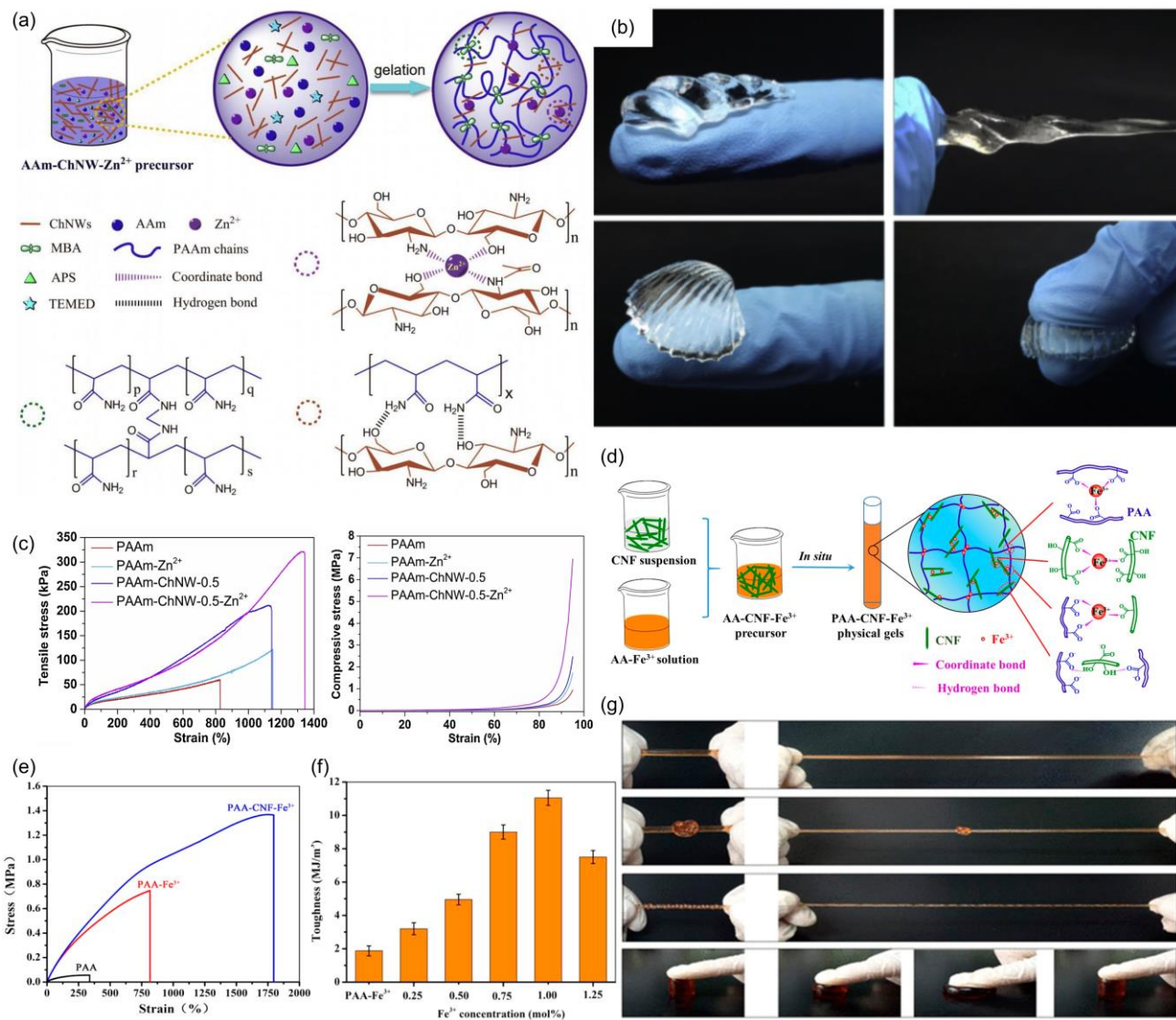


Figure 25. Mechanical properties of hydrogels enhanced by metal coordination sacrificial bonds. (a) Scheme showing polyacrylamide gels with chitin nanowhiskers and Zn²⁺ ions (PAAm-ChNW-Zn²⁺). (b) Different shapes of PAAm-ChNW-Zn²⁺ achieved by twisting and bending. (c) Tensile and compressive stress versus strain curves of PAAm-ChNW-Zn²⁺ gels show that addition of Zn²⁺ increases both tensile strength and toughness. Reprinted from ref. 125 with permission from Elsevier, copyright 2020. (d) Scheme showing the formation of a PAA matrix with cellulose nanofibrils and Fe³⁺ ions (PAA-CNF-Fe³⁺). (e) Uniaxial tensile tests reveal enhancement of mechanical properties upon addition of CNF and Fe³⁺ ions. (f) Tensile toughness varies with [Fe³⁺] in PAA-CNF-Fe³⁺. (g) PAA-CNF-Fe³⁺ gels can be (from top) stretched, knotted, twisted and compressed. Adapted with permission from ref. 126. Copyright 2017 American Chemical Society.

Shao and coworkers enhanced the mechanical properties of a PAA-based hydrogel using modified cellulose nanofibrils (CNFs).¹²⁶ Prior to their incorporation into the gel, the CNFs were oxidized using TEMPO, which converted the 5'-hydroxyl group on each pyranose ring to a carboxylate group. This carboxylate group, together with the residual acid groups from the PAA backbone, served as ligands when Fe³⁺ ions were introduced, forming a sacrificial coordinate network (Figure 25d). Unlike pristine PAA gels, which showed low tensile strength (0.035 MPa) and low maximum elongation (320%), PAA-Fe³⁺ gels with no CNFs show enhanced tensile strength (20-fold

increase) and elongation ratio (800%). This was further improved by the introduction of CNFs: PAA-CNF-Fe³⁺ showed a tensile strength of 1.37 MPa (~40 fold improvement over pristine PAA) and an elongation ratio of 1803% (Figure 25e). This proved that the enhancement in toughness of the material could be primarily attributed to the synergistic dual sacrificial metal coordinate network formed between the metal ion and both PAA and CNF matrices. Increasing [Fe³⁺] led to an increase in tensile toughness, likely due to a greater number of sacrificial metal coordination interactions. However, the authors note that material toughness and maximum elongation reach a maximum at 1 mol% Fe³⁺ (Figure 25f), allowing the material to be knotted, twisted and stretched (Figure 25g).

Hussain and coworkers used a similar strategy to make mechanically tough hydrogels by incorporating the natural polymer glycogen into a synthetic polymeric matrix composed of PVA and PAA.¹²⁷ Adding Fe³⁺ ions established a coordination network in which the hydroxyl groups of PVA and glycogen and the carboxylic acid motifs of PAA acted as ligands. Varying the concentrations of glycogen and PVA allowed the tensile strength, maximum elongation ratio, and tensile toughness of the material to be tuned, with a maximum toughness of 0.65 MJ/m³. Recently Zhou and coworkers have developed a dual network hydrogel that incorporates HB interactions in addition to ionic interactions between EDTA ligands and amine groups on the amino-functionalized polysiloxane backbone.¹²⁸ Tuning the relative amount of each motif enabled precise control over the hydrogel's tensile strength (0.07 to 1.46 MPa) and tensile toughness (0.06 to 2.17 MJ/m³).

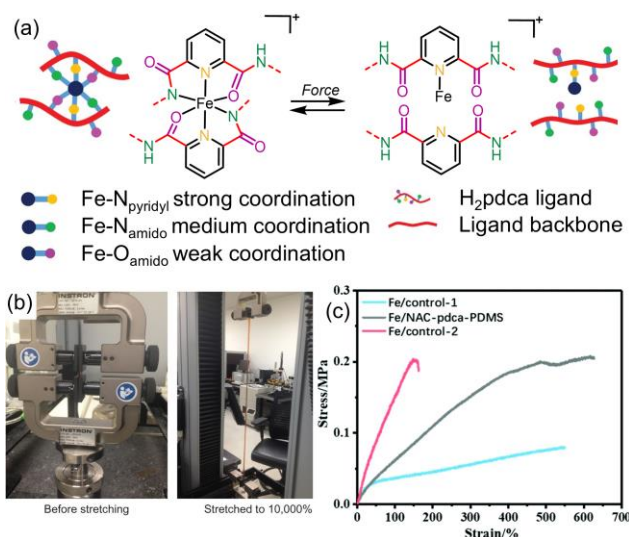


Figure 26. Modification of silicone-based elastomers using metal-ligand interactions as sacrificial bonds. (a) Scheme showing reversible formation and breaking of coordination bonds with various strengths in pdca-doped PDMS-based elastomers under application of force. (b) PDMS-pdca-Fe³⁺ can be stretched up to 10,000%. Figure adapted from ref. 129 with permission from Springer Nature, copyright 2016. (c) Stress-strain graph shows that Fe/NAC-pdca-PDMS has considerably higher toughness than coordination-free controls. Figure adapted from ref. 130 with permission from the Royal Society of Chemistry, copyright 2020.

Silicon-based elastomers are becoming increasingly important due to their applications in electronic materials. However, there is often a trade-off between the heat-dissipating characteristics and mechanical properties of these materials. To address this drawback, metal-coordination bonds have also been employed as sacrificial bonds to improve mechanical toughness of silicon-based elastomers.^{131–134} Bao and coworkers innovated this approach by positioning weakly- and strongly-coordinating binding sites next to each other in a single multidentate ligand incorporated within a polydimethylsiloxane (PDMS) elastomer.¹²⁹ To achieve this, 2,6-pyridinedicarboxamide (pdca) was selected as a tridentate Fe³⁺-binding ligand, in which the pyridine N atom coordinates most strongly and the O and N atoms of the amide form weaker, secondary coordination bonds (Figure 26a). Under stress, the authors hypothesized that the weaker coordination bonds would break to dissipate energy, but the metal ions would remain immobilized due to the stronger, Fe—N_{pyridyl} bonds, enabling rapid reformation of the weak coordination linkages, when the stress is removed. This design strategy proved successful: the pdca-modified PDMS achieved a maximum elongation of up to 10000% (Figure 26b). This result compares favorably to unadulterated PDMS, which has a maximum elongation of up to 1100%, and PDMS strengthened using other supramolecular approaches, which produced a maximum elongation of 2000%.¹³⁵ The authors attributed the enhanced mechanical properties to the ability of intrachain linkages to form, leading to the release of chain length upon fission of the weak metal-coordination bonds. Zhang and coworkers also used a similar strategy to strengthen PDMS in which Fe-pdca complexes act as weak coordination bonds which break to dissipate energy under stress, and stronger interactions between Fe³⁺ ions and carboxylate groups on *N*-acetyl-L-cysteine (Fe-NAC) held the metal ions in place.¹³⁰ This also led to enhanced tensile toughness of 0.866 MJ/m³, ~3-fold higher than controls without either ligand (Figure 26c).

6. Electrostatic Interactions

Electrostatic interactions arise from the Coulombic forces of attraction between species that have formal opposite charge. Though such attraction contributes in part to the strength of metal-ligand coordination bonds (see previous section), electrostatic interactions are far more universal and can be observed in a wide range of structural motifs. The strength of these interactions depends solely on the formal charge of interacting species and the distance between them. Consequently, these interactions have unique characteristics distinct from H-bonding and metal

coordination, such as their non-directional nature and the ability to form multiple interactions with the same charged center.

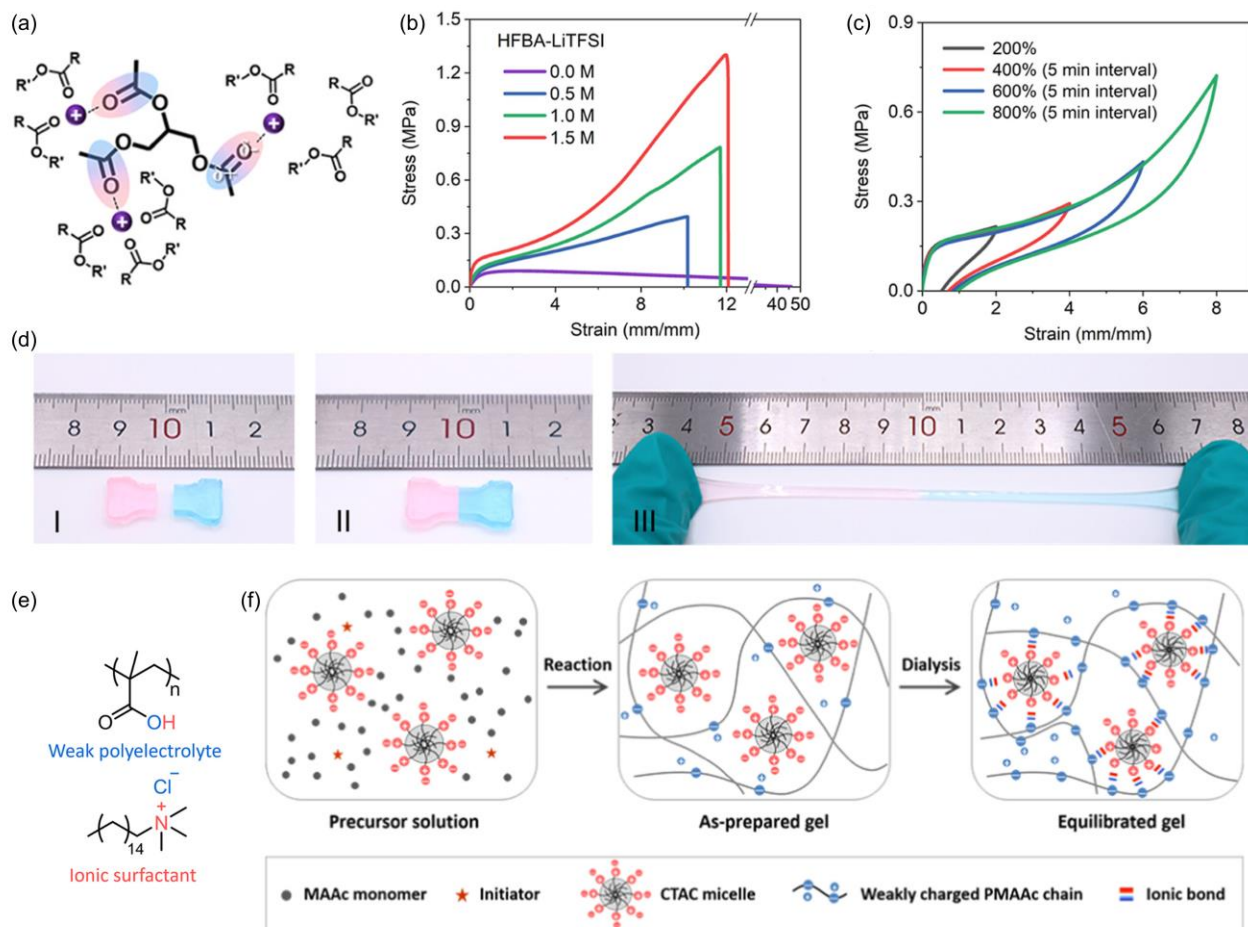


Figure 27. Ion-dipole electrostatic interactions in polymers and hydrogels. (a) Schematic representation of ion-dipole interactions between Li^+ ions and carbonyl motifs. (b) Effect of increasing $[\text{Li}^+]$ on the mechanical properties of pHFBA hybrid polymers. (c) Cyclic tensile loading experiments on pHFBA hybrid polymers reveal large hysteresis loops under different applied strains. (d) pHFBA polymeric systems show excellent stretchability even after self-healing. Adapted with permission from ref. 136. Copyright 2023 American Chemical Society. (e) Structure of polyelectrolyte and zwitterionic surfactant in methacrylic acid-based plastic-like supramolecular hydrogels. (f) Scheme showing preparation of methacrylate-based hydrogels incorporating polyelectrolyte surfactant complexes (PESCs). Adapted with permission from ref. 137. Copyright 2021 American Chemical Society.

Wang and coworkers recently reported an SPM based on an ion-dipole network generated by photopolymerizing 2,2,3,4,4,4-hexafluorobutyl acrylate (HFBA) monomers and introducing lithium bis(trifluoromethane)sulfonamide.¹³⁶ Li^+ ions interacted with the polar carbonyl bond on the acrylate species to crosslink the polymer chains (Figure 27a). The addition of these ion-dipole interactions led to increased tensile strength and tensile toughness, with maxima of 1.32 MPa and 6.67 MJ/m³, respectively, at 1.5 M Li^+ , which are 14.8- and 4.8-fold higher than the corresponding values for pure pHFBA (Figure 27b). Cyclic loading-unloading

experiments showed excellent energy dissipation (Figure 27c). The authors also noted that the high density of ionic crosslinks restricts the dynamic motion of the polymeric segments, decreasing the entropic gain upon heating and maintaining these dynamic interactions at elevated temperatures. These materials have showed excellent stretchability of over 600%, even after being cut and allowed to self-heal (Figure 27d).

Wu and coworkers have reported plastic-like supramolecular hydrogels formed by the polymerization of methacrylic acid in the presence of hexadecyltrimethylammonium chloride (CTAC) (Figure 27e).^{137,138} Hydration of the polymeric matrix induces strong electrostatic interactions between the weak polyelectrolyte and zwitterionic surfactant. These polyelectrolyte-surfactant complexes (PESCs) serve as crosslinks between polymer chains and, in conjunction with the hydrophobic effect induced by the surfactants, act as dynamic sacrificial interactions that enhance the hydrogel's mechanical properties (Figure 27f). For example, while the pure polyacrylamide gel is too weak to perform any tensile tests, increasing [CTAC] to 10 mol% leads to a tensile strength of 900 kPa. Interestingly, the authors also report that the extent of ionization of the polyelectrolyte can be affected by temperature, pH, and salt concentration providing additional handles with which to tune the mechanical properties of these materials.

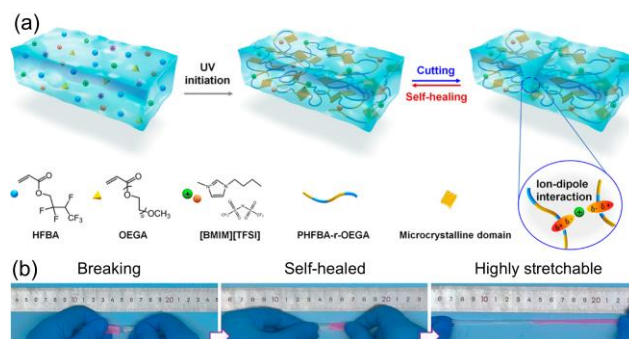


Figure 28. Electrostatic interactions in ionogels lead to enhanced mechanical properties. (a) Schematic depiction of ion-dipole interactions inside a polyethylene glycol-based matrix with embedded ionic liquid. (b) This material can be cut and self-healed and retains its high stretchability (scale in cm). Adapted with permission from ref. 139. Copyright 2021 American Chemical Society.

Ionic liquid-based polymers form an important class of polymeric gels as they have highly tunable ionic conductivity, leading to wide applications in electronics and robotics.¹⁴⁰ However, their practical applications demand high stretchability and robustness which are usually difficult to achieve in gel-like materials. To circumvent this problem, Zhang and coworkers developed a fluorinated polymeric matrix using HFBA and oligoethylene glycol monomers (OEGA) with various amounts of an embedded ionic liquid (IL), 1-butyl-3-methylimidazolium-

bis((trifluoromethylsulfonyl)imide) ([BMIM][TFSI], Figure 28a).¹³⁹ Sacrificial ion-dipole crosslinks between the imidazolium cations of the ionogel and the carbonyl groups of the acrylate component led to very high tensile strength and stretchability, reaching values of 251 KPa and 6000%, respectively, for the polymer containing 12.5 mol% IL. Furthermore, this material maintained its excellent stretchability even after being cut and allowed to self-heal (Figure 28b). Other groups have explored the effect of incorporating H-bonding alongside ion-dipole interactions on the mechanical toughness and stretchability of ionogels.^{141,142}

7. The Hydrophobic Effect

The hydrophobic effect describes the tendency of hydrophobic components to associate in aqueous solutions to minimize unfavorable enthalpic interactions with water of solvation and maximize the entropy of bulk water. This effect is a significant contributor to the formation and stability of tertiary and quaternary structures of large biomacromolecules such as proteins. In synthetic materials, the hydrophobic effect has been explored thoroughly in the context of water-soluble polymers, where hydrophilic components impart solubility whereas hydrophobic component induce polymer chain assembly.¹⁴³ Subsequent studies have explored the disruption of hydrophobic domains as a mechanism to dissipate energy,¹⁴⁴ due to their non-directional nature, which allows hydrophobic groups to re-associate quickly after force-induced dissipation. Moreover, the hydrophobic effect is compatible with a broad range of functional groups, spurring their investigation in materials alongside other non-covalent interactions to optimize mechanical properties.^{137,145}

Traditional methods to synthesize hydrogels that incorporate hydrophobic domains have been challenging due to the insolubility of the hydrophobic components in aqueous media.¹⁴⁶ These solubility issues not only frustrate materials synthesis but can also lead to a loss of structural control at the macroscopic and microscopic scales. Candau and coworkers overcame the synthetic bottleneck using a simple, robust synthesis based on a micellar copolymerization technique.¹⁴⁴ In this, the hydrophobic monomer is solubilized using micelles (formed from surfactants such as sodium dodecyl sulfate, SDS) in an aqueous environment.¹⁴⁴ Then, via a free radical mechanism, micellar hydrophobic comonomer and solubilized hydrophilic comonomer copolymerize to form the hydrogel. Due to the high local concentration of hydrophobes within the micelles, the resultant material contains blocks of randomly distributed hydrophobic components.¹⁴⁷ This method has been widely employed to incorporate hydrophobic associations as sacrificial motifs in materials.

Okay and coworkers were among the first to demonstrate that increasing hydrophobic associations, by incorporation of hydrophobic segments in a synthetic hydrogel via a copolymerization technique, led to drastically improved mechanical properties of the resultant material.¹⁴⁸ Hydrogels were constructed from acrylamide monomers and *N,N'*-methylenebis(acrylamide), which served as an interchain crosslinker. Modified acrylamides containing alkyl chains including *N*-butyl, *N*-hexyl and *N*-octyl were also incorporated, to act as the hydrophobic segment. While pure PAAm gels were too weak to conduct any mechanical experiments, increasing the amount of hydrophobic *N*-alkylacrylamide components led to increasing toughness. Indeed, PAAm hydrogels modified with 10 mol% of *N*-octylacrylamide achieved 313% elongation at a stress of 71 kPa (Figure 29a). Furthermore, these gels could also withstand knotting and extreme deformation. The authors also addressed the role of the micellar polymerization technique in generating favorable materials properties. Comparative studies showed that the creation of blocks of hydrophobic segments via the micellar copolymerization technique using SDS yielded enhanced toughness over randomly distributed hydrophobic units. However, the authors hypothesized that the presence of the SDS surfactant in the polymeric matrix might affect energy dissipation by increasing the solubility of the hydrophobic components. In a subsequent study,¹⁴⁹ the authors found that the removal of surfactant led to the loss of the dynamic mechanical properties associated with reversible hydrophobic effects (Figure 29b). This is because SDS solubilizes the hydrophobic blocks, leading to enhanced stability of the hydrophobic groups, but the removal of the surfactant forces the hydrophobic groups to form much stronger associations to avoid any interaction with the aqueous environment. Hence the presence of surfactants produces weaker gels with higher G'' values but leads to better dynamic properties. For example, hydrogels with an elongation ratio of almost 1700% in the presence of SDS achieved only a 700% elongation ratio in the absence of SDS (Figure 29b).

However, the dependence of this category of hydrogels on surfactants, which are often cytotoxic, renders them unsuitable for applications in biological systems. Hence, Meng and coworkers recently targeted the formation of hydrogels via hydrophobic associations in the absence of any synthetic surfactants.¹⁵⁰ Alginate and stearyl methacrylate (C18M) were selected as the hydrophilic and hydrophobic comonomers, respectively. Instead of using a surfactant, amphiphilic regenerated silk fibroin (RSF) allowed micelle formation by solubilizing the hydrophobic C18M via RSF's conserved hydrophobic sequences. Additionally, Ca^{2+} ions were added to induce hydrophilic ionic interactions between the metal cation and the negatively charged carboxylate ion on the alginate polymer chain

(Figure 29c). This hybrid material exhibited increasing toughness as the concentration of the hydrophobic component (C18M) was increased. Compared to an alginate-RSF gel without C18M, the stress at 50% strain increased more than 200% when [C18M] was increased to 30 mM. Additionally, the yield strength of the material was significantly increased when C18M was incorporated (Figure 29d). Moreover, strain sweep measurements revealed that the intersection of G' and G'' occurred at 109% strain for the hybrid material ([C18M]= 30 mM), which was much higher than for pure alginate gels (1.59%) or alginate-RSF gels without C18M (10%) (Figure 29e). These results clearly indicated that the enhancement in mechanical properties was primarily due to hydrophobic associations acting as sacrificial bonds and not the ionic interactions present in the material. Importantly, this study highlights a path forward to leverage the hydrophobic effect in biocompatible energy-dissipating materials.

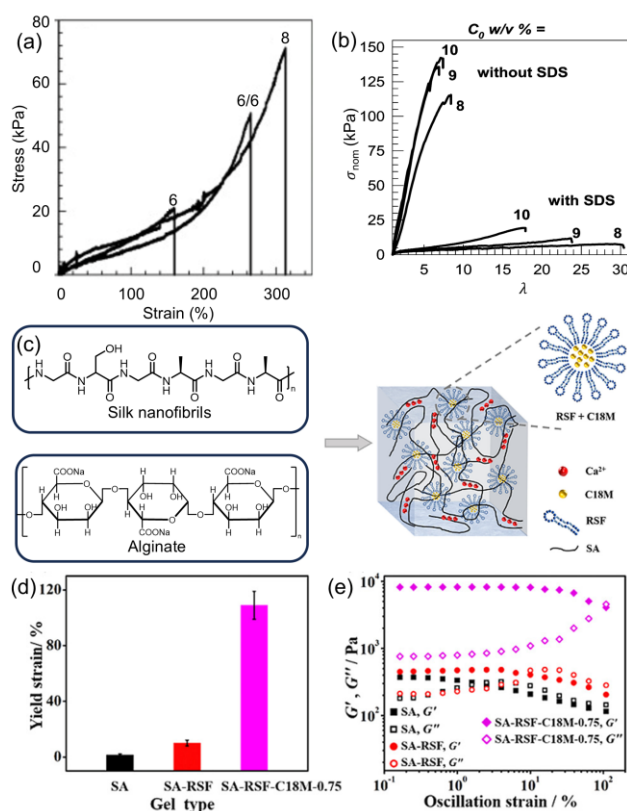


Figure 29. Hydrophobic dynamic interactions. (a) Stress-strain curves of hydrogels containing 10 mol% of various alkyl chains such as *N*-octyl (8), *N*-hexyl (6) and *N,N*-dihexyl (6/6). Reprinted from ref. 148 with permission from Elsevier, copyright 2009. (b) Without SDS, higher tensile strength is obtained but with lower elongation ratio. Adapted with permission from ref. 149. Copyright 2012 American Chemical Society. (c) Scheme showing the formation of hydrophobic interactions in water without surfactants by using natural silk fibroin and sodium alginate. (d) Strain-at-break values are much higher in the presence of hydrophobic monomers. (e) Shear rheology experiments show higher G' and G'' values in the presence of the hydrophobic component (C18M). Reprinted with permission from ref. 150. Copyright 2020 American Chemical Society.

8. Host-Guest Complexation

While previous sections have focused on individual types of non-covalent interaction, the complexation of a suitable guest by a macrocyclic host is driven by multiple effects that may include electrostatic interactions, H-bonding, the hydrophobic effect, and others.¹⁵¹ Unlike these individual interaction modes, (macrocyclic) host-guest interactions can result in a precise co-location of two components (host and guest) in 3-D space, often with a preferred relative orientation. Furthermore, both the host and guest can be independently tailored to tune the thermodynamic and kinetic favorability of association. Consequently, host-guest interactions have been explored extensively for the generation of supramolecular polymer networks, in which both polymer backbones and the crosslinks between them are scaffolded by non-covalent interactions.¹⁵¹ Furthermore, the supramolecular nature of these interactions has also been exploited as sacrificial bonds which has led to enhanced mechanical properties of the bulk polymer.

Among supramolecular hosts, cyclodextrins (CDs) have garnered particular interest due to their water solubility, synthetic accessibility, and their interior cavity that is capable of binding various hydrophobic guests including azobenzenes and adamantanes.¹⁵² Yang and coworkers leveraged the selective host-guest recognition of adamantane by β -cyclodextrin (β -CD) to modify the mechanical properties of PAAm hydrogels.¹⁵³ To synthesize these materials, a host-guest macromer (HGM) was prepared from an adamantane-functionalized hyaluronic acid and mono-acrylated β -CD, and subsequently copolymerized with acrylamide using a methylene bisacrylamide (MBA) cross-linker to synthesize MBA-HGM polymers (Figure 30a). Control samples without HGM, crosslinked solely using different concentrations of MBA, were also prepared (denoted MBA-L and MBA-H for relatively low and high concentrations, respectively). Dynamic oscillatory experiments established that increasing the concentration of MBA in the HGM-free polymers increased the storage modulus G' but did not change the loss modulus G'' , indicating that the material became stronger but also more brittle. Conversely, the MBA-HGM polymer exhibited an increase in both G' and G'' , suggesting that the addition of host-guest sacrificial bonds endowed the material with overall higher tensile toughness. Uniaxial tensile tests showed that, compared to MBA-H, MBA-HGM had much higher stretchability (750% vs 300%) and higher fracture stress (an additional 100% compared to controls, Figure 30b). The ability of MBA-HGM to dissipate energy was confirmed through cyclic loading-unloading studies, which showed that MBA-HGM polymers dissipated 22.8% of the loading energy,

compared to negligible energy dissipation by the control materials, as evidenced by negligible area inside the hysteresis loop.

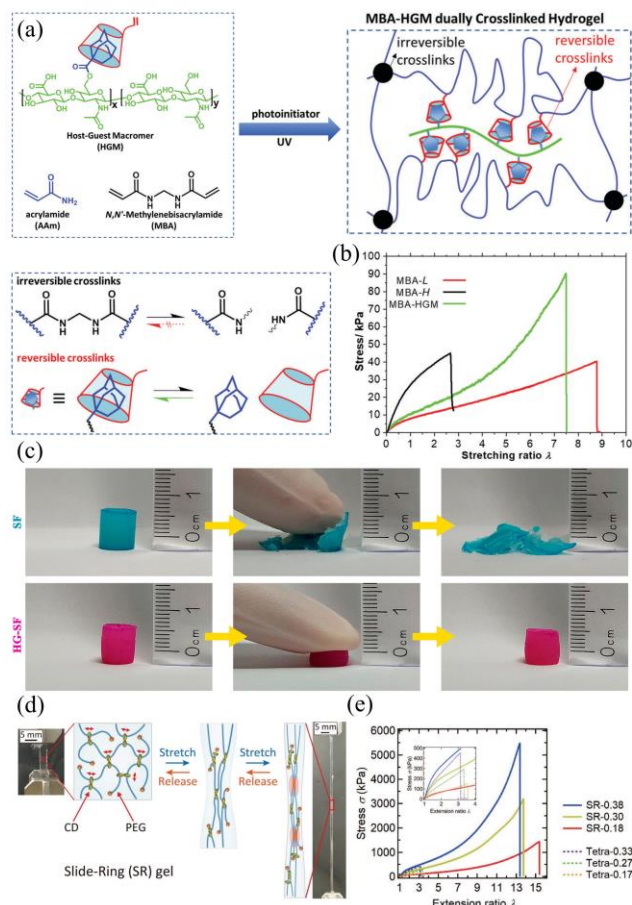


Figure 30. (a) Scheme showing the preparation of MBA-HGM polyacrylamide gels. (b) Uniaxial tensile tests showing superior performance of MBA-HGM polyacrylamide gels showing much higher overall toughness. Figure adapted from ref. 153. (c) Compression tests comparing host guest-based silk fibroin (HG-SF) and unadulterated silk fibroin (SF). Reprinted with permission from ref. 154. Copyright 2020 American Chemical Society. (d) Scheme showing energy dissipation mechanism as well as ultra-stretchability of slide-ring (SR) gels. (e) Tensile tests showing the superior performance of SR-gels (SR- x , where x denotes the volume fraction of PEG) compared to tetra-PEG (tetra- x) control gels. Figure adapted from ref. 155. Reprinted with permission from AAAS.

Bai and coworkers used a similar strategy to enhance the mechanical properties of silk fibroin (SF) based hydrogels.¹⁵⁴ To chemically modify the SF polymers, carboxylic acid functional groups were introduced to serine and tyrosine residues by oxidation and diazo linkages, respectively. These modified SFs were then coupled with either cholesterol or β -CD to give Chol-SF and CD-SF respectively. Mixing these two polymers allowed the hydrophobic cholesterol moieties to be encapsulated by β -CD, forming hydrogels (HG-SF). Tensile tests revealed a dramatic increase in both fracture stress (3.16 MPa vs 0.1 MPa) and maximum elongation (65% vs 31%) compared to a SF-only control. The combined effect of these changes resulted in a 65-fold enhancement in the tensile

toughness of HG-SF vs its counterpart without sacrificial host-guest interactions. This was also demonstrated by a compression test (Figure 30c), in which the control polymer disintegrated while HG-SF recovered its shape.

Ito and coworkers used an innovative host-guest strategy that utilizes poly(rotaxane)s to form slide-ring (SR) hydrogels (Figure 30d).¹⁵⁵ Specifically, PEG chains were first incubated with hydroxy-propyl- α -cyclodextrin (HP- α -CD), to form rotaxane-like assemblies in which PEG interpenetrated the CDs. Subsequent cross-linking of the CDs with divinyl sulfone led to hydrogels in which the CDs retained their mobility along the PEG chains. Under tensile stress, CD molecules were hypothesized to slide toward each other, leading to extended lengths of PEG chains between increasingly congested cross-linking points. Under large stress, these PEG regions were expected to become highly ordered and form a close packed structure via strain-induced crystallization, leading to further increased mechanical toughness. SR hydrogels with different volume fractions (x) of PEG (denoted SR- x) were compared to control polymers containing tetra-arm PEG as an immobile cross-linker. Uniaxial tensile tests demonstrated that SR-gels had much higher tensile toughness than their slide-free counterparts, with SR-0.38 showing a 40-fold improvement over the best performing tetra-PEG variant. The authors also noted that the toughness of the gels is reduced by using PEG chains of lower molecular weight, perhaps due to the smaller sliding distance of the cross-links as well as more defects in the PEG network. The ability of polymer chains to interpenetrate CDs was also observed by Takashima and coworkers, who unexpectedly observed favorable toughness in copolymers of ethyl acrylate and a γ -CD methylacrylamide monomer (PA γ CDAAmMe).¹⁵⁶ These copolymers exhibited a comparable fracture energy to nitrile rubber but with much higher Young's modulus.

Cucurbit[n]urils (CB[n]) are another classic barrel-like host used for molecular recognition.¹⁵⁷ Scherman and coworkers recently employed CB[8] to improve the mechanical properties of PAAm gels.¹⁵⁸ PAAm gels were prepared by copolymerizing 1-benzyl-3-vinylimidazolium ions with hydrophilic acrylamide monomers using MBA as crosslinker. Subsequent addition of CB[8] led to complexation-driven crosslinking of the gel network, in which the CB[8] motifs formed reversible 1:2 host-guest complexes with the pendent benzylimidazolyl chains on the polymer backbone (Figure 31a). This hydrogel exhibited phenomenal mechanical properties, with a tensile toughness (750 MJ/m³) comparable to cartilage, and was able to support 500 times its own weight without material failure. Control gels with either no CB[8] motifs or with only smaller CB[7] motifs, incapable of binding two benzylimidazolyl groups simultaneously, led to the loss of mechanical strength, demonstrating the essential and

specific nature of the host-guest linkages. Uniaxial tensile tests showed that the CB[8]-containing material was highly stretchable, achieving a maximum elongation of 2400%. The authors also note that, above a strain rate of 100 mm/min, there was a significant decrease in the overall toughness of the material. They hypothesize that this could arise from the inability of the dynamic host-guest components to effectively rearrange on small timescales, which could lead to decreased energy dissipation (Figure 31b). Other work from the same laboratory has focused on exploring the effect of changing the size of the guest motif.¹⁵⁹ The Scherman group has also explored CB[*n*]-based single network hydrogels without the presence of covalent cross-linking¹⁶⁰ as well as hybrid double-network hydrogels with both CB[*n*]-based and DNA-mediated cross-linking.¹⁶¹

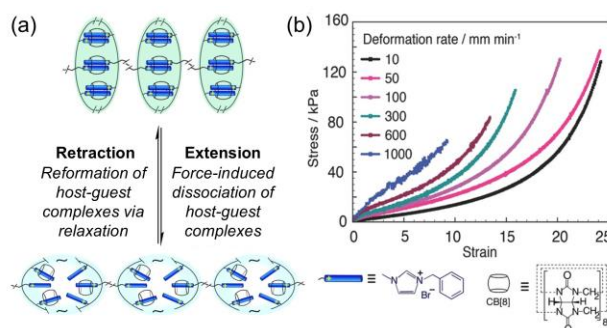


Figure 31. (a) Scheme showing cucurbit[8]uril-based gels and their energy dissipation mechanism under loading and unloading (blue rod is guest; white cylinder is host). (b) Tensile tests at different strain rates show lowered toughness at higher strain rates. Figure adapted from ref. 158. Copyright 2017 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.

Crown ethers have also found widespread utility as supramolecular hosts for sacrificial host-guest interactions. Yan and coworkers exploited the molecular recognition of a dialkylammonium salt by benzo-21-crown-7 (B21C7) molecules to prepare supramolecular polymeric networks (SPNs).¹⁶² Specifically, a covalent polymer network was synthesized by ring-opening metathesis polymerization (ROMP) of B21C7-functionalized norbornene monomers. This was then mixed with different molar ratios of a dialkylammonium crosslinker (DAAS) to form different SPNs (SPN-*x*, where the molar ratio of dialkylammonium salt to B21C7 is *x*:10; Figure 32a). The charged ammonium species interacted with the crown ether oxygens to form a reversible non-covalent interaction which could rupture to dissipate energy under mechanical stress and reform when the stress had been removed. Tensile tests established increasing the amount of the dialkylammonium-crosslinker increased both the Young's modulus and the tensile toughness of the hybrid gels. SPN-5 showed the best mechanical properties, with maximum toughness (22 MJ/m³), fracture stress (5.1 MPa) and elongation-at-break (648%) (Figure 32b,c). Subsequent work from the same laboratory compared SPNs to covalent polymeric networks (CPNs) and dual polymer networks (DPNs) that have both

supramolecular as well as covalent linkages (Figure 32d).¹⁶³ Tensile tests on the three materials revealed the SPN and DPN exhibited 10.3- and 9.1-fold improvement, respectively, of tensile toughness over CPN, further emphasizing the importance of sacrificial non-covalent host-guest interactions. Furthermore, DPNs also exhibited almost a 100% increase in tensile strength over SPNs, showing that a dual network system comprising both covalent and supramolecular network can exhibit both high strength and toughness (Figure 32e).

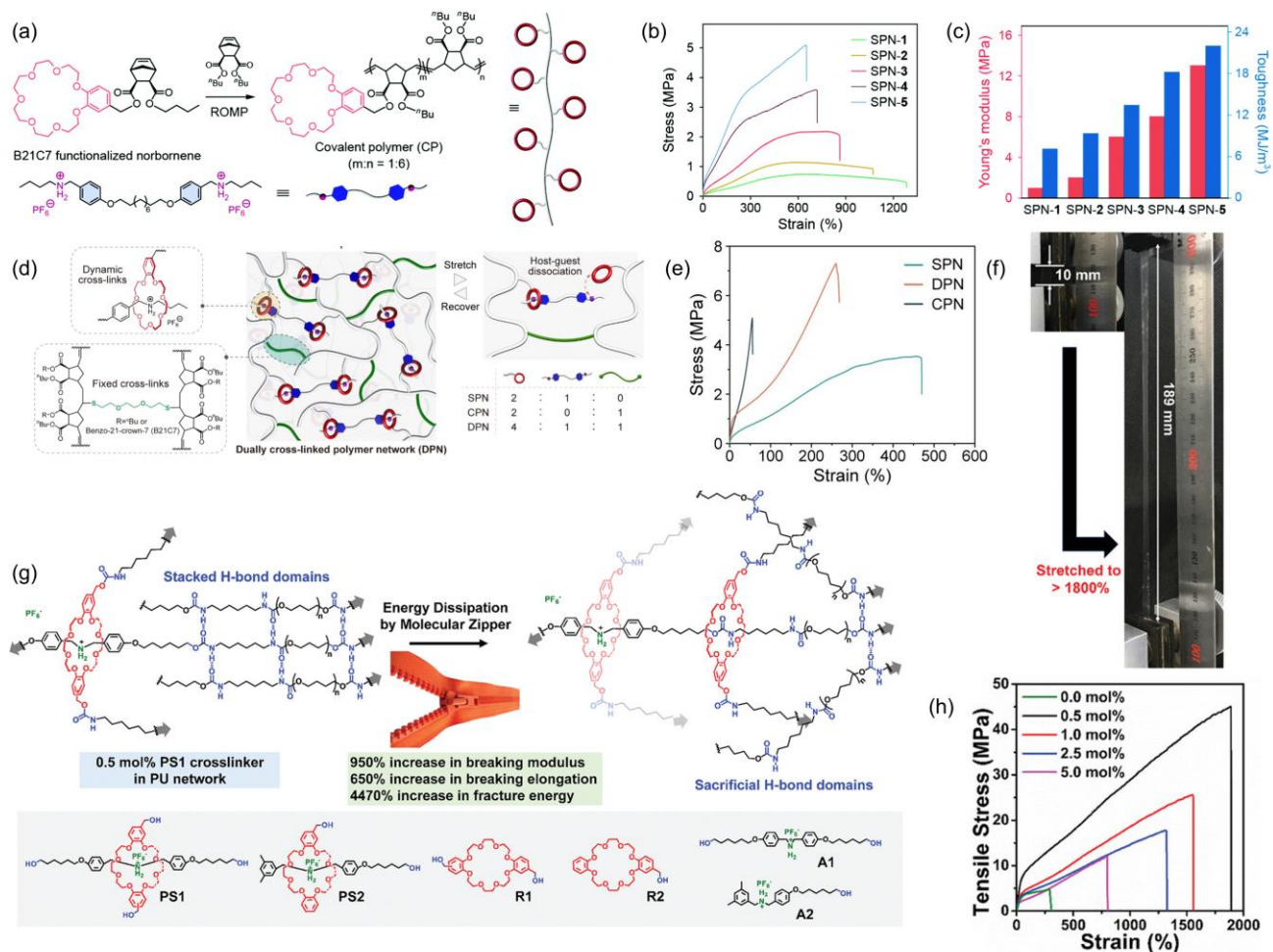


Figure 32. (a) Synthesis of supramolecular networks (SPNs) composed of B21C7-norbornene-based hosts and dialkylammonium guests. (b) Uniaxial tensile tests of SPN-*x* (molar ratio of guest:host = *x*:10) shows increasing toughness and yield strength with increasing guest concentration. (c) Comparative studies show increasing Young's modulus and tensile toughness with an increasing concentration of host-guest interactions. Figure adapted from ref. 162. (d) Dually cross-linked polymers with both supramolecular and covalent cross-links. Three polymers (SPN, CPN and DPN) were constructed with different compositions of supramolecular and covalent networks. (e) Tensile tests comparing the mechanical performance of SPN, CPN and DPN show DPN having higher overall toughness. Figure adapted from ref. 163. Copyright 2023 Wiley-VCH GmbH. (f) PPS1 can be extended to a length of more than 1800% without fracturing. (g) Scheme showing the design of zipper gels with monomers PS1 and PS2 as well as controls R1, R2, A1 and A2. (h) Mechanical performance of PPS1 gels with different concentrations of PS1. Figure adapted from ref. 164. Copyright 2020 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim.

Host-guest supramolecular networks have also been used in conjunction with other types of sacrificial bond to further improve energy dissipation. Qu and coworkers combined the host-guest complexation of a

dibenzylammonium motif by a dibenzo-crown ether with amide-mediated H-bonding to make mechanically robust polymers that undergo zipper-like ring sliding.¹⁶⁴ Central to their design was the incorporation of pseudorotaxane monomers (PS1), comprising a dibenzo-24-crown-8 motif and a dibenzylammonium-based axle, into PU networks that are inherently decorated with extensive HBs. When tensile force is applied across the resultant material (PPS1), the host-guest interaction breaks, leading leads to ring sliding of the benzo-crown ether along the PU backbone, disrupting the H-bonding between linear PU chains (Figure 32g). This phenomenon, termed ‘molecular zippering’, dissipates a large amount of energy. Tensile tests showed that PPS1 could be stretched to an impressive 1890% without fracture (Figure 32f) and had much higher tensile strength and fracture energy than pristine PU (8.5- and 45-fold improvements, respectively). On decreasing the crosslinking density (PPS2), the tensile strength, stretchability and toughness all fall significantly, further demonstrating the profound effect of the host-guest interactions. The authors also showed that a low concentration of PS1 (0.5 mol%) was optimal for mechanical properties and noted that this value was one of the lowest compared to other ring-sliding crosslinkers. Increasing the concentration of PS1 beyond this value led to decreased tensile toughness (Figure 32h). This work, together with the PEG-CD slide-ring gels by Ito and coworkers (*vide supra*),¹⁵⁵ highlights the potential for the mechanical bond¹⁶⁵ to act as a distinct sacrificial bond motif.¹⁶⁶

9. Covalent Bonds

A key feature of a sacrificial bond is its ability to re-form after mechanical force or applied load is removed. For these reasons, non-covalent interactions, as discussed in the previous sections, are natural choices for sacrificial bonds due to their inherent reversibility. Covalent bonds, in comparison, initially seem unsuitable for reversible energy dissipation. However, two major classes of covalently-bonded motifs that can spontaneously recover after fission are known: dynamic covalent bonds¹⁶⁷ and mechanophores.^{168,169}

9.1. Dynamic Covalent Bonds

As opposed to traditional, stable covalent bonds, dynamic covalent bonds readily break and subsequently reform.¹⁷⁰ A recent review of the rich chemistry of dynamic covalent bonds¹⁶⁷ emphasizes the sheer number of different reaction modes that lead to dynamic bonding, including Diels-Alder cyclization,¹⁷¹ transesterification,¹⁷² transamination,¹⁷³ photodimerization,¹⁷⁴ boronic acid exchange,¹⁷⁵ transcarbamoylation,¹⁷⁶ and imine metathesis,¹⁷⁷ among others. Consequently, the kinetics of bond cleavage and formation can be finely tuned to achieve desirable

bond lifetimes, leading to widespread study of dynamic covalent bonds in polymeric materials and even deployment in industrial processes.¹⁷⁸ However, most polymer research that utilizes dynamic covalent chemistry has been geared towards the preparation of self-healing materials, which are often flexible and have poor strength and toughness.¹⁷¹

The mechanism by which dynamic covalent bonds break and reform to achieve reversible molecular network rearrangements (MNR) can be classified as either concerted or stepwise. In a *concerted* mechanism, bond breaking and bond formation occur simultaneously (Figure 33a), such that the number of covalent bonds remains constant at all times. In contrast, *stepwise* molecular rearrangement consists of distinct bond cleavage and bond formation steps (Figure 33b). If the bond formation happens before the bond cleavage step, the total number of covalent bonds temporarily increases and the process is termed an *associative* stepwise MNR. Conversely, if the bond cleavage precedes the bond forming step, *dissociative* stepwise MNR occurs, with a momentary decrease in the total number of bonds.¹⁷⁹ Given that energy dissipation requires that there is a reduction in the extent of bonding, i.e. bonds are broken, in response to external force, dynamic covalent bonds that exchange via a dissociative stepwise mechanism are desirable as sacrificial bonds.

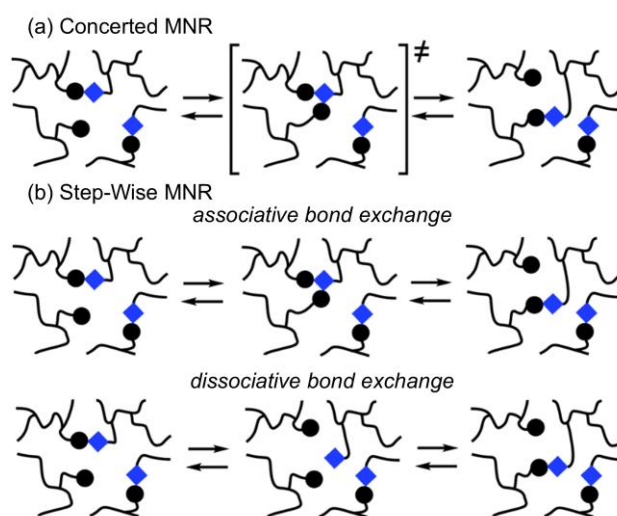


Figure 33. Mechanism of bond exchange in dynamic covalent bonds. (a) Concerted and (b) Stepwise molecular network rearrangements (MNRs). Stepwise MNRs can proceed via associative and dissociative pathways. Figure adapted from ref. 179 with permission from the Royal Society of Chemistry, copyright 2019.

In 2018, Chi and coworkers reported one of the first examples of using dynamic covalent bonds as sacrificial bonds for energy dissipation.¹⁸⁰ Amine-terminated PDMS was copolymerized with 1,3,5-triformylbenzene to generate a polymer network crosslinked with strong yet dynamic imine bonds (Figure 34a). To introduce weaker dynamic covalent bonds, 2-aminophenyl disulfide (ADPS) was incorporated as an additional crosslinker during

polymerization. Crucially, the disulfide linkages in ADPS were hypothesized to serve as sacrificial dynamic bonds which would cleave to dissipate energy. The authors reported that increasing the ADPS concentration enhanced the material's stretchability, up to a maximum elongation of 4300%. A control sample in which ADPS was replaced with an alternative crosslinker that did not contain a disulfide gave a maximum elongation of only 580%, confirming that the dynamic aromatic disulfide linkages act as sacrificial bonds. Reducing the rate of elongation allowed the material to be stretched further but resulted in a reduction in yield strength, as the slower elongation rate provides more time for the dynamic covalent bonds to reorient and reorganize (Figure 34b). Cyclic tensile loading experiments produced large hysteresis loops characteristic of energy dissipation; however, extended recovery times (48 h) were required to fully recover the material's dissipative capacity (Figure 34c). Wang and coworkers used a similar strategy to design a PDMS-based material that utilized dynamic imine and boroxine bonds to dissipate energy.¹⁸¹ Boron-oxygen linkages have also been used to modulate the mechanical properties of alginate-based materials.¹⁸²

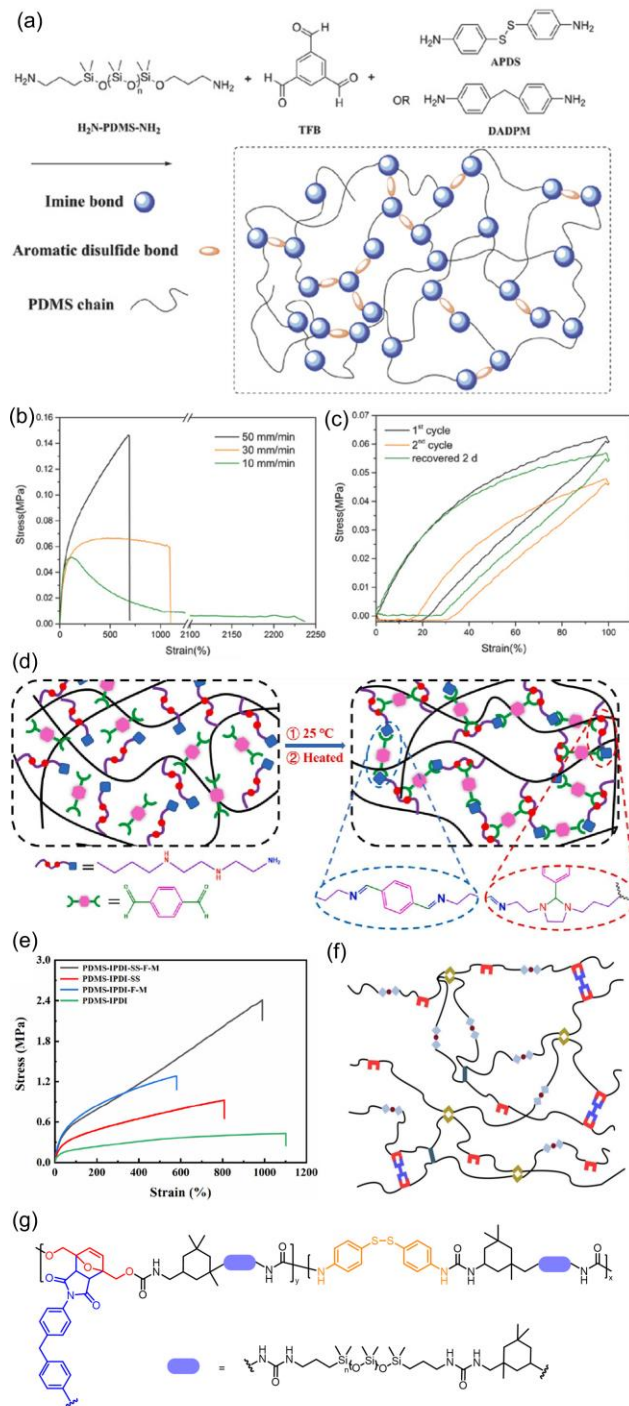


Figure 34. Siloxane-based elastomers containing dynamic covalent bonds. (a) Schematic representation of the cross-linked PDMS polymer containing imine and aromatic disulfide dynamic bonds. (b) With decreasing rate of elongation, the yield strength decreases and the maximum elongation increases. (c) Cyclic tensile tests reveal that a recovery time of 48 h is necessary to reproduce the same extent of energy dissipation. Figure adapted from ref. 180. Copyright 2018 WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim. (d) Schematic of PDETAS elastomer incorporating imine and cyclic aminal dynamic covalent bonds. Reprinted from ref. 183 with permission from Elsevier, copyright 2023. (e) PDMS elastomers showing highest mechanical properties when dynamic covalent linkages are present. (f) Schematic representation and (g) chemical structure of PDMS polymer incorporating three sacrificial bond motifs: H-bonding between urea motifs, disulfide bonds, and Diels-Alder-type dynamic covalent linkages. Figure adapted from ref. 184. Copyright 2024 Wiley Periodicals LLC.

Deng and coworkers leveraged the different amine sites in diethylenetriamine-functionalized polysiloxanes (PDETAS) to prepare two types of sacrificial dynamic covalent bond in a single polymer network.¹⁸³ Reaction of the crosslinker 1,4-diformylbenzene (DFB) with the primary amines on the PDETAS side-groups formed imine linkages whereas reaction with the secondary amines formed aminal linkages via a ring-closing mechanism (Figure 34d). The imine and the aminal bonds not only underwent dynamic exchange with the free amines but could also metathesize (exchange with) each other. This highly dynamic behavior of the system led to efficient energy dissipation, leading to a tensile strength of ~25 MPa, much higher than previously reported silicone materials. Increasing the concentration of DFB initially increased the tensile strengths due to an increase in the extent of crosslinking, but excess DFB saturated the amine sites of the polymer network, favoring single-point attachment and therefore reducing the extent of crosslinking.

As with other types of sacrificial bonds, dynamic covalent bonds can be employed synergistically with other types of sacrificial bond. Wang and coworkers polymerized amine-terminated PDMS with IPDI to generate urea motifs capable of H-bonding (Figure 34e–g).¹⁸⁴ Unreacted isocyanates were subsequently reacted with ADPS, to introduce a disulfide linkage, and 2,5-furandimethanol, which acts as a functional handle for Diels-Alder reactions with maleimide moieties. Hence the resultant material, denoted PDMS-IPDI-SS-F-M, incorporated three sacrificial networks: one based on H-bonding and two based on dynamic covalent bonds (Figure 34e). Compared to PDMS-IPDI polymers without further functionalization (i.e., polymers that could engage only in H-bonding), PDMS-IPDI-SS-F-M hybrids exhibited enhanced strength without compromising the maximum elongation of the material, suggestive of energy dissipation enabled by the sacrificial nature of the dynamic covalent bonds (Figure 34f). Moreover, eliminating either of the types of dynamic covalent bond led to a significant decrease in material toughness, indicating synergistic behavior between the two dynamic covalent sacrificial networks (Figure 34f). The authors were also able to modulate the tensile strength and toughness of the material by tuning the relative abundance of the three different sacrificial networks.

9.2. Mechanophores

Mechanophores are motifs which undergo a structural rearrangement when mechanical force is applied to them, leading to cleavage of at least one covalent bond.¹⁸⁵ Such motifs have been identified as potential energy dissipating units, as the structural rearrangement often reverts to its original state upon removal of the applied force. Recent

reviews highlight many such mechanophore moieties that have been explored as mechanochromic units or as energy dissipating units in materials.^{168,186,187} Consequently, we discuss here just two illustrative examples of mechanophore chemistry in the context of sacrificial bonds.

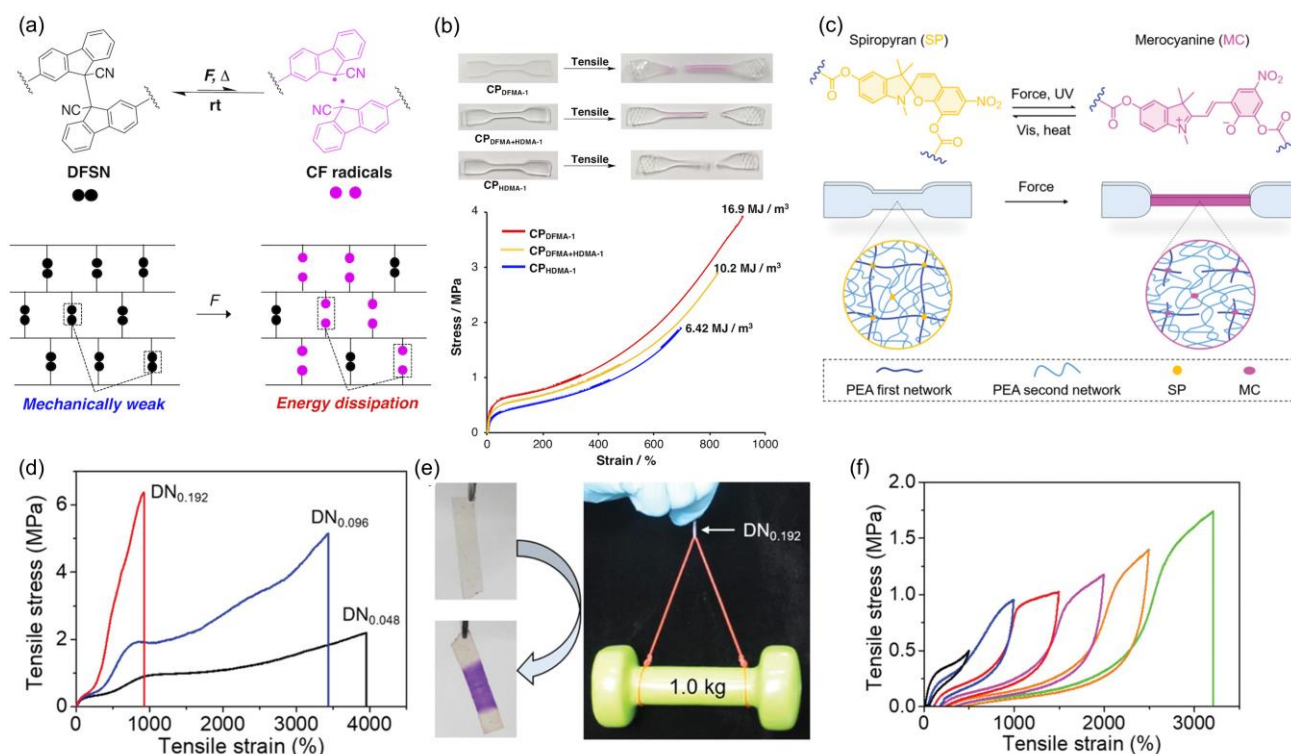


Figure 35. Mechanophores that act as sacrificial bonds to dissipate energy under high stress or strain (a) Schematic representation of how the central bond in DFSN undergoes homolytic cleavage (b) Compared to the control, the material with DFSNs show much higher yield strain and toughness evident in uniaxial tensile tests. Site of elongation turns pink due to formation of cyanofluorene radicals. Adapted with permission from ref. 188. Copyright 2020 American Chemical Society. (c) Spiroyrans in a poly(ethylacrylate) matrix undergo conversion from the SP to the MC form under stress. (d) Decreasing the density of spiropyrans (DN_x, where x is the mol% of spiropyran) leads to loss in yield strength but greater stretchability. (e) Application of lateral stress leads to macroscopic color change. (g) Cyclic tensile tests show large hysteresis loops before fracture. Figure adapted from ref. 189. Copyright 2020 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.

Otsuka and coworkers have explored difluorenylsuccinonitrile (DFSN)-based crosslinkers as mechanophores in polyacrylates.^{188,190} When force is applied to DFSN, the central symmetric C-C bond undergoes homolytic fission to form two stable cyanofluorene radicals (Figure 35a). These radicals are bright pink in color, which enables the authors to visualize the exact point in the matrix where these dynamic bonds have broken under stress (Figure 35b). The incorporation of DFSNs into a double-network polyacrylate was observed to increase fracture energies by 2.6-fold compared to a mechanophore-free control polymer (Figure 35b).¹⁸⁸ Moreover, the radical nature of the cyanofluorenes allowed the extent of sacrificial bond rupture to be monitored using electron paramagnetic resonance (EPR) spectroscopy, which revealed that increasing mechanical toughness correlated well with an

increased in the number of generated radicals. Subsequent work from the same laboratory introduced a non-symmetric mechanophore, cyanofluorenyl-arylbenzofuranone, as a crosslinker between poly(*n*-hexyl methacrylate) chains.¹⁹¹ Unlike DFSN, this mechanophore generates two different radicals that have distinct photochromic properties and hence shows mixed mechanochromism. While the tensile strength and toughness of this material was markedly improved over a mechanophore-free control, no substantial differences were observed compared to the DFSN-based materials. However, EPR studies showed that the concentration of radicals generated at 300% strain was 10 times higher for materials containing the non-symmetric mechanophore than for the symmetric DFSN-based materials.

Spiropyrans are a common mechanochromic motif that have mostly been used to visualize the stress point in polymer materials, due to their readily observable color change upon mechanically-induced cleavage from the colorless spiropyran (SP) form to the pink merocyanine (MC) form.^{192,193} Cao recently employed spiropyrans as sacrificial motifs in a double-network polymer in which both networks contained poly(ethyl acrylate) crosslinked with spiropyrans (Figure 35c).¹⁸⁹ The resultant material showed high ultimate strength and maximum elongation attributable to the strong, rigid nature of the spiropyran crosslinks. Even with a low concentration of spiropyran crosslinker (0.096 mol%), a maximum fracture energy of 8.4 kJ/m² and a maximum elongation of 3400% were achieved (Figure 35d). The rupture of the sacrificial network could be visualized by a macroscopic color change (Figure 35e), and cyclic tensile loading-unloading experiments showed large hysteresis loops consistent with energy dissipation (Figure 35f). Spiropyrans have also been explored as visualizable sacrificial toughening agents by Creton and coworkers.^{194–196}

10. Hidden Length

Most of the examples discussed so far have employed sacrificial bonds as the connection (crosslinker) between polymer chains. However, if sacrificial bonds are incorporated into a polymer chain such that the sacrificial bond bridges two atoms on the same chain, the formation of the bond produces a macrocyclic portion of the polymer chain that is necessarily not fully extended and, therefore, does not experience strain.¹⁹⁷ Consequently, cleavage of the sacrificial bond “reveals” a portion of the polymer backbone—known as *hidden length*—that can take on strain and reduce the strain experienced by other segments of the polymer chain (Figure 36a). In many cases, the stretching of this hidden length dissipates much more energy than the enabling bond cleavage event.¹⁹⁸ Unsurprisingly, many

natural systems employ the combination of sacrificial bonds and hidden length (SBHL) to dissipate energy under external stress in biomaterials, including bone and abalone shells.^{9,10,14,198} However, examples of synthetic materials that harness SBHL to withstand impact are rare.

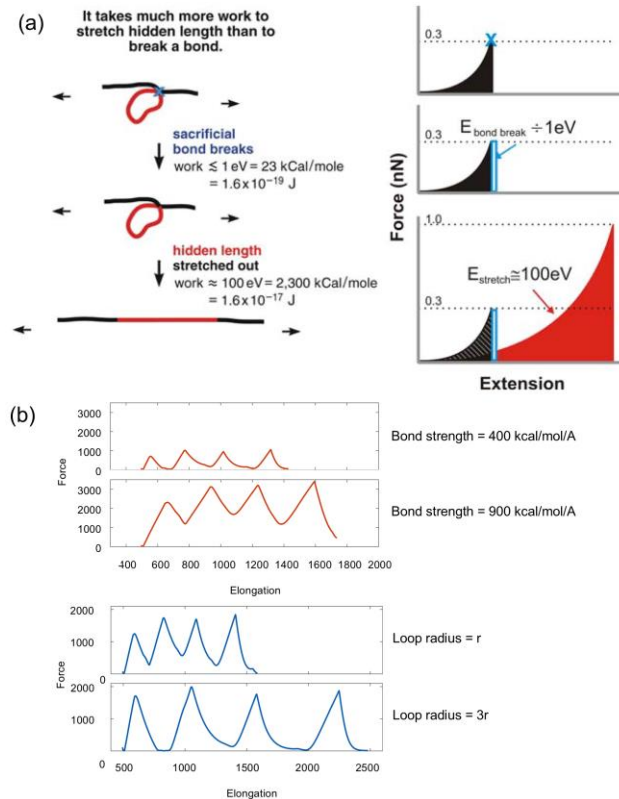


Figure 36. (a) Mechanism of energy dissipation in systems with sacrificial bonds and hidden length. Reprinted from ref. 198 with permission from Elsevier, copyright 2006. (b) Force-elongation graphs with different sacrificial bond strengths (red) and different loop radii (blue). The energy required to release the hidden length remained constant, but the elongation length increased because of the increased loop radius. The force required to release the hidden length increased with increasing bond strength. Used with permission of American Society of Mechanical Engineers (ASME), from ref. 199; permission conveyed through Copyright Clearance Center, Inc.

Theoretical insights into the design of SBHL materials were gleaned by Deng and coworkers, who employed MD studies to examine how structural parameters affected the mechanical properties of a model SBHL system.¹⁹⁹ Polymer fibers were modeled using a simple bead-spring model, due to its similarity to biological systems such as collagen, spider silk, and amyloid. Simulations were performed while varying parameters including loop radii, number of loops, crosslink interaction strength, and fiber stiffness, to establish the nature of the dependence of material toughness on these parameters (Figure 36b). For example, it was found that toughness increased linearly with the number of loops, because of the increased number of breaking and unfolding events, but toughness

exhibited a quadratic dependence on the strength of crosslinks. This study may guide the rational design of SBHL-based materials with tunable toughness and mechanical properties.

One of the few experimental examples of SBHL being deployed in a synthetic material was reported by Gault and coworkers, who realized a shock-absorbing material, based on a modified talin protein, that showed impact resistance from supersonic projectiles.²⁰⁰ Talin is a naturally-occurring mechanosensitive protein implicated in the transduction of mechanical force to the cytoskeleton. In this work, a talin mutant, modified with cysteine residues to permit crosslinking by maleimides, consisting of three rod-like domains (R1, R2, and R3) was used as a mechanosensitive module; these three rod-like domains in the wild-type protein undergo stepwise force-induced unfolding (Figure 37a).²⁰¹ To make the talin shock-absorbing material (TSAM), protein monomers were crosslinked by a trivalent maleimide crosslinker (Figure 37b–f), forming a gel. A control sample prepared by reacting talin monomers with a monovalent maleimide did not show gelation, confirming that only the trivalent crosslinker was successfully polymerizing the talin monomer (Figure 37e). TSAM and a poly(vinylpyrrolidone) (PVP) hydrogel control were subjected to impact tests with projectiles sized between 20 and 70 microns at a velocity of 1.5 km/s. Whereas PVP showed no resistance to the projectile, TSAM resisted the projectile and did not allow the projectiles to permeate the bulk material; this compared favorably to a commercial aerogel that permitted particle penetration up to 8 mm. Here, the myriad non-covalent interactions between the amino acid residues of the three domains act as sacrificial bonds, and the polypeptide backbone acts as hidden length (Figure 37f); by combining these two, protein unfolding leads to extraordinary energy dissipation in these hydrogels.²⁰⁰

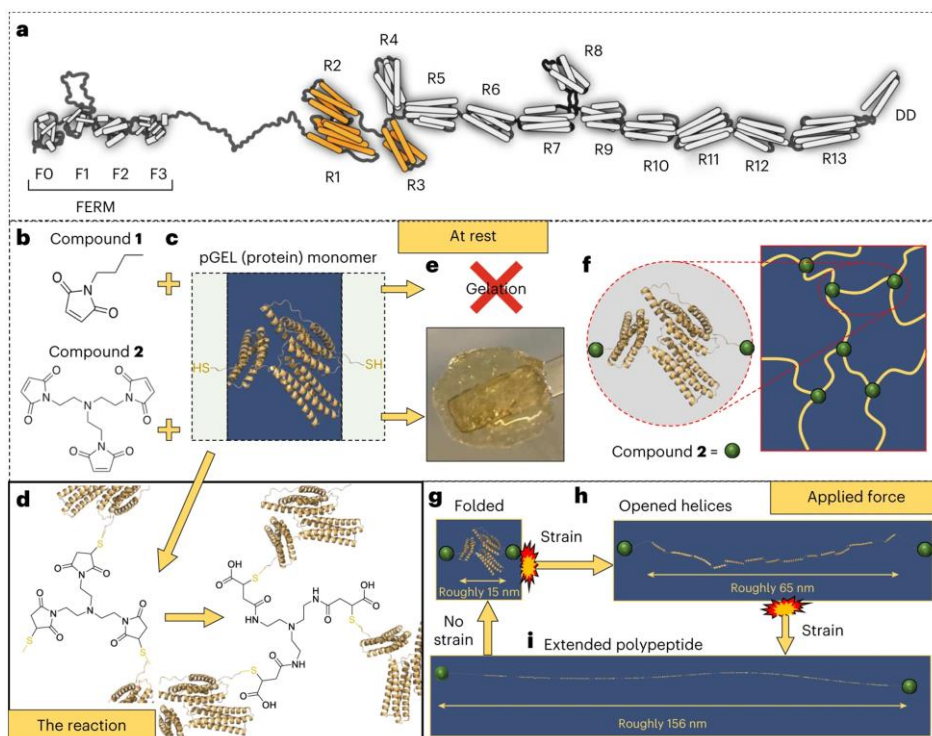


Figure 37. (a) Talin polymers undergo sequential unfolding to reveal hidden length. (b) Maleimide reagents were reacted with (c) cysteine-modified talin monomers to form (d) crosslinked polymers. (e) Trivalent maleimide crosslinkers led to the formation of gels whereas monovalent maleimides did not. (f) Crosslinked talin polymers. (g) Folded talin domains undergo (h) strain-induced unfolding under applied force to form extended polypeptides that can be (i) stretched under further strain to $\sim 10\times$ their original length. Figure reproduced from ref. 200 with permission from Springer Nature, copyright 2023.

11. Sacrificial Conformation

Recently, Wang *et al.* reported a novel strategy to dissipate energy without cleavage of a sacrificial bond or interaction, instead relying on the flexible conformation of β -cyclodextrin motif to tolerate strain.²⁰² Their cross-linked polymers (CCN1–3) were prepared by subjecting a cyclodextrin monomer functionalized with two pendent alkene motifs (CM) to photo-induced thiol-ene click chemistry with different amounts of a linear chain monomer (LM) (Figure 38a). A control material composed entirely of the linear monomers (LCN) was also prepared. The authors found that increasing the CM content increased the Young's modulus of the material but also decreased its maximum elongation. As a result, CCN2 (CD:LM = 1:3) showed the highest toughness of the samples tested, 60-fold higher than that of the control LM (Figure 38b). CCN2 also exhibited the maximum energy dissipation (Figure 38c). These results indicate that the conformational flexibility of β -cyclodextrins can be harnessed as an energy dissipation mechanism without breaking any bonds, covalent or non-covalent, signifying a unique *sacrificial conformation* approach to engineering tough materials.

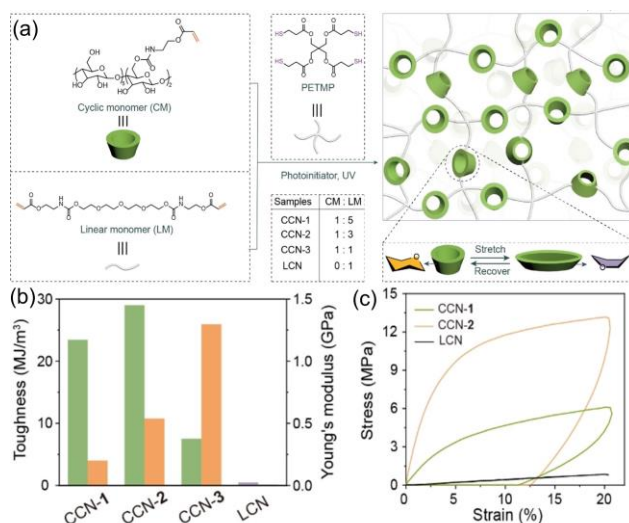


Figure 38. (a) Scheme showing the formation of cross-linked CCN polymers from cyclic monomers (CM) containing β -cyclodextrins and linear monomers (LM). (b) Comparing toughness (green) and Young's modulus (orange) of different polymers with different CM:LM compositions reveals superior mechanical performance by CCN2. (c) Cyclic tensile tests show maximum energy dissipation by CCN2. Figures adapted from ref. 202. Copyright 2024 Wiley-VCH GmbH.

12. Conclusions and Outlook

The incorporation of sacrificial bonds into synthetic materials is a powerful strategy to enhance material toughness and strength via the dissipation of energy under applied load. Progress over the past several years has advanced multiple chemical motifs that can serve as sacrificial interactions, ranging from non-covalent interactions such as H-bonding, metal coordination, and the hydrophobic effect to covalent bonds such as dynamic linkages and mechanophores. In concert with experimental studies, the development of new theories to describe energy dissipation in network materials and more widespread simulation of these theories using molecular dynamics are providing experimentalists with a clearer understanding of how to program energy-dissipating materials with a desired set of mechanical properties, balancing tensile toughness, maximum elongation, and dissipative capacity. The result is a host of new materials with mechanical properties approaching or even surpassing those of natural biomaterials.

Since the last comprehensive review on this topic almost a decade ago,¹⁷ progress in the field has continued apace and led to several enabling innovations, three of which we highlight here. First, advances in computational and theoretical approaches can now employ fully atomistic MD simulations to reproduce and rationalize experimental observations. Second, the incorporation of multiple sacrificial bonding motifs, whether using a single type of interaction (e.g., HBs using ureas and alcohols) or distinct bonding modalities (e.g., H-bonding and metal

coordination), is enabling materials with more sophisticated responses to applied force. Third, the continued expansion of the toolkit of sacrificial bonding motifs, including noncanonical energy dissipation modes such as mechanophores and conformationally-flexible cyclodextrin units, provides ever more chemical handles for the design of new, class-leading materials.

However, as we opined in the Introduction, the most substantial unmet challenges in the field are those that relate to the design of materials with specific properties. Table 1 summarizes some key mechanical properties measured for materials discussed in this review. While the absolute values of many of these metrics is remarkable, the large ranges of strength, strain, and toughness values suggest that structure-property relationships are underdeveloped and warrant rigorous attention. The routine comparison of new materials against well-defined standard reference materials, such as commercial samples of poly(urethane)s and Kevlar, could ease comparison between different studies and help to parse differences arising from sacrificial bond design.

Table 1. Summary of selected mechanical properties reported for materials using sacrificial bonds.

Sacrificial bond strategy	Ultimate strength	Maximum strain	Tensile toughness	References
Hydrogen bonding (UPy)	4.5 – 87.2 MPa	3% – 1285%	1 – 386 MJ/m ³	44, 52, 56, 64, 67–70
Hydrogen bonding (amides)	0.35 – 98.5 MPa	17% – 2160%	1.6 – 550 MJ/m ³	42, 45, 74, 78–85
Hydrogen bonding (alcohols)	0.57 – 83.5 MPa	2% – 1876%	0.55 – 1896 MJ/m ³	89–91, 95, 96
Metal coordination	0.20 – 25.7 MPa	120% – 10000%	0.0025 – 96 MJ/m ³	106, 108–110, 114, 116, 125–127, 129
Host-guest complexation	0.09 – 45.1 MPa	65% – 2400%	9.63 – 22 MJ/m ³	153–155, 158, 162–164
Electrostatic interactions	0.25 – 1.32 MPa	1100% – 6007%	6.67 – 8.7 MJ/m ³	136–139
Dynamic covalent bonds	2.4 – 24.7 MPa	988% – 4300%	13.6 MJ/m ³	180, 183, 184
Mechanophores	0.48 – 3.9 MPa	921% – 3200%	0.008 – 16.9 MJ/m ³	188, 189
Sacrificial conformation	538 MPa ^a	110%	30 MJ/m ³	202

^a Reported value denotes stiffness.

Developing comprehensive structure-property relationships will also facilitate the design of next-generation impact-resistant architectures. In the remainder of this section, we highlight several opportunities for continued innovation in this area as the community continues its pursuit of biomimetic load-bearing materials, along with potential strategies to address each opportunity.

Programmable sacrificial bond dynamics. Arguably the most salient challenge is the tendency of synthetic materials to require long relaxation times, on the order of minutes to hours, to fully recover their capacity for energy dissipation. These long relaxation times, which result from the slow re-formation of broken sacrificial bonds, hinder the development of practical materials that tolerate the accumulated damage of frequent, repetitive loads. Incorporating extrinsic sacrificial crosslinkers (cf. Figure 2c) with mobilities that approach the diffusion limit may be a viable strategy to address this limitation.

Directionality as a design parameter. Most sacrificial bond motifs reported to date rely on directional interactions, such as H-bonding and metal coordination. However, non-directional interactions such as electrostatics or the hydrophobic effect provide substantial advantages, especially with respect to recovery time, and warrant further study. Indeed, the directionality of a sacrificial bond could be viewed as an important design parameter in addition to canonical considerations such as bond strength.

Accumulated damage to covalent networks. Although the presence of sacrificial bonds greatly increases the resistance of materials to strain-induced deformation, prolonged exposure to applied loads will degrade typical soft materials due to damage to the underlying covalent polymer network. Supramolecular polymer networks offer distinct advantages in this regard, as discussed earlier, but the mechanical properties of the pristine materials are often unsatisfactory due to the reliance of the network structure on weaker, non-covalent interactions. Establishing methods to prepare strong and tough structures scaffolded entirely by non-covalent bonds between building blocks may usher in new classes of resilient soft materials.

Synergistic, hierarchical sacrificial networks. More exploration of systems with multiple sacrificial bonds is needed. Though many of the types of sacrificial bond discussed in this review have not been explored for their ability to endow impact resistance, even fewer have been combined to form synergistic sacrificial networks; those materials that do leverage multiple sacrificial bonds, especially those that do so in a hierarchical manner, already show great promise. Achieving such hierarchical bond breakage will require precise control over the number, strength, and spatial distribution of sacrificial bonds, the kinetics of their breaking and re-forming, and the dynamic nature of the covalent polymer network. Chemical strategies to modulate these parameters include the use of electron-withdrawing or -donating groups on bonding motifs to tune the strength of individual sacrificial bonds, employing molecular strain to affect the energetics of bond scission, and leveraging iterative polymerization methods to better control the distribution of sacrificial bonding motifs along covalent polymer backbones. Rational design of such networks, which requires renewed attention from both experimentalists and theoreticians, should produce materials with hitherto unprecedented properties.

Designing heterogeneity across length scales. One of the largest opportunities in materials design, not just for materials with sacrificial bonds, is the ability to design heterogeneity into materials architecture, particularly on the macroscale. This is especially pertinent for impact-resistant materials: consider the crumple zones of a modern car,

which direct energy around the passenger cabin and protect the human inhabitants from the most severe damage. How might similar macroscale heterogeneity be programmed into polymer networks? One potential solution emerges from biological systems, which routinely employ hierarchical assembly as an information-efficient means to program structural sophistication over multiple length scales. Designing hierarchically-architected sacrificially-bonded structures may provide similarly heterogeneous materials from synthetic components.

13. Abbreviations

2DPA	2D polyamide
ADPS	2-aminophenyl disulfide
APBIA	2-(4-aminophenyl)-1 <i>H</i> -benzimidazol-5-amine
ASC	acylsemicarbazide
BF	bamboo fiber
B21C7	benzo-21-crown-7
[BMIM][TFSI]	1-butyl-3-methylimidazolium-bis((trifluoromethylsulfonyl)imide)
BPE	biobased polyester elastomer
C18M	stearyl methacrylate
CD	cyclodextrin
ChNW	chitin nanowhisker
Chol	cholesterol
CNC	cellulose nanocrystal
CCN	covalently-crosslinked network
CNF	cellulose nanofibril
CNT	carbon nanotube
CTAC	hexadecyltrimethylammonium chloride
DABA	4,4-diaminobenzoyl triphenylamine
DDM	diamino-diphenylmethane
DFB	1,2-diformylbenzene
DFSN	difluorenylsuccinonitrile
DGEBA	diglycidyl ether-bisphenol A
DPA	3-hydroxy-2-(hydroxymethyl)-2-methylpropanoic acid
ENR	epoxidized natural rubber
EPR	electron paramagnetic resonance
HAPAA	hydrophobic-associating poly(acrylic) acid
HB	hydrogen bond
HDI	hexamethylene diisocyanate

HEMA	2-hydroxyethyl methacrylate
HFBA	2,2,3,4,4,4-hexafluorobutyl acrylate
HGM	host-guest macromer
HMDI	cyclohexylmethane-4,4'-diisocyanate
iGWLC	inelastic glassy worm-like chain
IHP	impact-hardening polymer
IL	ionic liquid
Im	imidazole
IPDI	isophorone diisocyanate
IU	imidazolidinyl urea
LCN	linearly-crosslinked network
MA	methacrylate
MBA	methylene bis-acrylamide
MD	molecular dynamics
MWCNT	multiwall carbon nanotube
NAC	<i>N</i> -acetyl-L-cysteine
NAGA	<i>N</i> -acryloyl glycinamide
PAA	poly(acrylic acid)
PAAm	poly(acrylamide)
PANI	poly(aniline)
PBSA	poly(butylene succinate- <i>co</i> -butylene adipate)
PCL	poly(caprolactone)
pdca	2,6-pyridinedicarboxamide
PDETAS	diethylenetriamine-functionalized polysiloxanes
PDMS	poly(dimethylsiloxane)
PEG	poly(ethylene glycol)
PEG-DE	poly(ethylene glycol) diglycidyl ether
PESC	polyelectrolyte-surfactant complexes
PPG	poly(propylene glycol)
PTA	poly(tannic acid)
PTMEG	poly(tetramethylene ether glycol)
PU	polyurethane
PVA	poly(vinyl alcohol)
PVP	poly(vinylpyrrolidone)
ROMP	ring-opening metathesis polymerization
RSF	regenerated silk fibroin

SBHL	sacrificial bond-hidden length
SDS	sodium dodecyl sulfate
SF	silk fibroin
SPM	supramolecular polymeric material
SPN	supramolecular polymeric network
(S)SBR	(solution-polymerized) styrene-butadiene rubber
SSG	shear-stiffening gel
STF	shear-thickening fluid
TA	tannic acid
TPY	terpyridine
TSAM	talin shock-absorbing material
TST	transition state theory
WLC	worm-like chain
UPy	ureidopyrimidinone

14. Conflicts of Interest

The authors declare no competing financial interests.

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Data Availability Statement

No primary research results, software or code have been included and no new data were generated or analyzed as part of this submission.