Soft Materials by Design: Unconventional Polymer Networks Give Extreme Properties

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Acronym

APS: ammonium persulfate CB[n]: cucurbit[n]uril CD: cyclodextrin CNT: carbon nanotubes DOPA: 3,4-dihydroxyphenyl-L-alanine EDC: 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide Fmoc: fluorenylmethyloxycarbonyl G: α -L-guluronic acid GO: graphene oxide M: β -D-mannuronic MBA: *N*,*N*'-Methylenebisacrylamide NdFeB: neodymium iron boron NHS: N-hydroxysuccinimide NVP: N-vinylpyrrolidone PAA: poly(acrylic acid) PAAm: polyacrylamide PCL: poly(caprolactone) PDLLA: poly(DL-lactic acid) PDMA: poly(*N*,*N*-Dimethylacrylamide) PEDOT:PSS: poly(3,4-ethylenedioxythiophene):poly (styrene sulfonate) PEG: poly(ethylene glycol) PEO: poly(ethylene oxide) PHEMA: poly(2-hydroxethyl methacrylate) PLA: polylactide PLGA: poly(DL-lactic acid-co-glycolic acid) PMA: polymethacrylic acid PNIPAm: poly(*N*-isopropylacrylamide) PPG: poly(propylene glycol) PPS: poly(propylene sulfide) PVA: poly(vinyl alcohol) RGD: Arg-Gly-Asp TEGDMA: trimethylene glycol dimethacrylate TEMED: *N*,*N*,*N'N'*-tetramethylene-diamine UPN: unconventional polymer network UPy: ureido-pyrimidinone

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Abstract

Hydrogels are polymer networks infiltrated with water. Many biological hydrogels in animal bodies such as muscles, heart valves, cartilages, and tendons possess extreme mechanical properties including extremely tough, strong, resilient, adhesive, and fatigue-resistant. These mechanical properties are also critical for hydrogels' diverse applications ranging from drug delivery, tissue engineering, medical implants, wound dressing, and contact lenses to sensors, actuators, electronic devices, optical devices, batteries, water harvesters, and soft robots. Whereas numerous hydrogels have been developed over the last few decades, a set of general principles that can rationally guide the design of hydrogels using different materials and fabrication methods for various applications remain a central need in the field of soft materials. This review is aimed to synergistically report: i). general design principles for hydrogels to achieve extreme mechanical and physical properties, ii). implementation strategies for the design principles using unconventional polymer networks, and iii). future directions for the orthogonal design of hydrogels to achieve multiple combined mechanical, physical, chemical, and biological properties. Because these design principles and implementation strategies are based on generic polymer networks, they are also applicable to other soft materials including elastomers and organogels. Overall, the review will not only provide a comprehensive discussion on the rational design of soft materials, but also provoke interdisciplinary discussions on a fundamental question: why does nature select soft materials with unconventional polymer networks to constitute the major parts of animal bodies?

Soft Materials by Design: Unconventional Polymer Networks Give Extreme Properties

1. Introduction

As hydrophilic polymer networks infiltrated with water¹, hydrogels are the major components of animal bodies, constituting most of their cells, extracellular matrices, tissues, and organs (**Figure 1** and **Figure 2**). Not surprisingly, hydrogels have been widely used in biological and biomedical applications such as vehicles for drug delivery²⁻⁵, scaffolds for tissue engineering⁶⁻⁸, models for biological studies⁹⁻¹³, medical implants^{14,15}, wound dressing¹⁶⁻¹⁸, and contact lenses^{15,19}. More recently, intensive efforts have been devoted to exploring hydrogels' emerging applications in devices and machines²⁰ such as hydrogel sensors²¹⁻²⁴, actuators²⁵⁻²⁸, soft robots^{26,29,30}, electronic devices³¹⁻³³, batteries^{34,35}, supercapacitors³⁶, iontronic devices^{27,37}, magnetic devices^{28,38-40}, optical devices⁴¹⁻⁴³, acoustic devices^{44,45}, living devices^{24,46}, underwater adhesives⁴⁷⁻⁴⁹, bioadhesive devices^{47,50}, and coatings^{51,52} (**Figure 3**).

Mechanical properties of hydrogels are crucial to the survival and wellbeing of animals, and greatly affect the abovementioned applications of hydrogels. The pioneering works in the field of polymers and soft materials have laid the foundation for understanding the elasticity, swelling, poroelasticity, viscoelasticity, fracture, and fatigue of hydrogels (e.g., Ref.⁵³⁻⁶⁶ and the references in them). However, the inverse question – how to design hydrogels that possess certain mechanical properties or certain properties in general – still poses a grand challenge in the field of polymers and soft materials⁶⁶⁻⁶⁹. This challenge becomes even more daunting, when one targets at hydrogels' extreme mechanical properties, such as extremely high values of fracture toughness⁷⁰, strength^{71,72}, resilience^{73,74}, interfacial toughness⁴⁹, fatigue threshold⁷⁵⁻⁷⁷, and interfacial fatigue threshold⁷⁸.

Despite the abovementioned grand challenge, the design of hydrogels with extreme mechanical properties is of both fundamental and practical importance. From the fundamental aspect, many biological hydrogels have achieved extreme mechanical properties necessary for their survival and well-being through evolution (Figure 1). For example, cartilage is a tough connective tissue that covers the surfaces of joints to provide reduced friction⁷⁹. The human knee joint cartilage (i.e., articular cartilage) typically needs to sustain compressive stresses of 4-9 MPa for 1 million cycles per year, while maintaining high fracture toughness around 1,000 Jm⁻² (Figure $(2a)^{80}$. The high fracture toughness of articular cartilage is mainly attributed to its abundant strong collagen fibers interpenetrated with proteoglycan macromolecules. This structure of articular cartilage provides both viscoelasticity and poroelasticity for mechanical dissipation^{81,82}. The viscoelasticity of articular cartilage is mainly associated with local rearrangement of aggrecan, adhesive interactions of aggrecan, and reconfiguration of collagen⁸²; the poroelasticity of articular cartilage is governed by the interstitial fluid movement through the porous extracellular matrix⁸¹. Tendon is a strong connective tissue that connects muscle to bone and muscle to muscle. The human patellar tendon can sustain a high tensile strength over 50 MPa^{83,84}, owing to its unique hierarchical fibrous structure that enables the simultaneous stiffening of bundles of collagen fibers before their tensile failure (Figure 2b)^{85,86}. Heart valves generally possess both high resilience

above 80% and high fracture toughness around 1,200 Jm⁻² ^{87,88}, which are two seemingly contradictory properties (**Figure 2c**). The elastin and crimped collagen fibers in the heart valve are elastic and non-dissipative under moderate deformation, giving the heart valve the high resilience ⁸⁹. However, under large deformation, the stiffening and fracture of the collagen fibers dissipate substantial mechanical energy, making the heart valve tough as well⁹⁰. The adhesion of soft connective tissues on bones can be extremely fatigue-resistant. For example, the cartilage-bone interface in the human knee joint can sustain compressive stresses of 1 MPa along with an interfacial toughness around 800 Jm⁻² over 1 million cycles of loads per year (**Figure 2d**)^{80,91,92}. The fatigue-resistant adhesion of soft tissues (e.g., tendons, ligaments, and cartilages) to rigid bones is commonly achieved through nanostructured interfaces composed of aligned collagen nano-fibrils and ordered hydroxyapatite nano-crystals⁹³⁻⁹⁵. What are nature's design principles for various biological hydrogels besides the abovementioned ones (**Figure 2**) to achieve extreme mechanical properties? This is still a largely unanswered question, even in light of the pioneering works in the field of polymers and soft materials (e.g., Ref.⁵³⁻⁶⁵ and the references in them).

From the practical aspect, applications of hydrogels generally require the hydrogels to possess a set of specific properties. For example, hydrogels designed with different moduli and viscoelastic properties have been used to regulate stem cell fate and activity (**Figure 3**)^{10,12,93,96}. The applications of hydrogels as artificial cartilages and spinal discs require the hydrogels to be fatigue-resistant under cyclic mechanical loading^{75,76,97,98}. The mesh size of hydrogels' polymer networks is critical to their applications in controlled drug delivery (**Figure 3**)^{42,99,100}. More recent applications of hydrogels as various devices and machines require the hydrogels to possess specific properties, for instance, stimuli-sensitivity for hydrogel sensors and actuators^{30,101-103}, strong adhesion for hydrogel coatings⁴⁹, optical transparency for hydrogel water harvesters¹⁰⁴ (**Figure 3**).



Figure 1. Biological hydrogels in the human body can possess extreme mechanical properties. Aorta with tensile strength of 0.2-3.7 MPa⁸⁰; heart valve with resilience above 80% and fracture toughness around 1,000 J m^{-2 87,88}; tendon with tensile strength of 10-100 MPa¹⁰⁵, fracture toughness of 20-30 kJ m⁻² and fatigue threshold of 1,000 J m^{-2 106}; skeletal muscle with fracture toughness around 2,490 J m⁻² and fatigue threshold around 1,000 J m^{-2 80}; articular cartilage with fracture toughness of 800-1,800 J m^{-2 107}; and tendon/cartilage/ligament-bone interfaces with interfacial fatigue threshold around 800 J m^{-2 78,91}. These biological hydrogels can provide inspirations for the design of synthetic hydrogels that possess extreme mechanical properties.



Figure 2. Design principles and implementation strategies for various biological hydrogels to achieve extreme mechanical properties: a. high toughness of cartilage due to viscoelastic and poroelastic dissipation of the polymer networks^{82,108,109}, **b.** high tensile strength of tendon due to simultaneous stiffening of multiple polymers in the fibrous hierarchical structure^{85,105}, **c.** high resilience and facture toughness of heart valve due to delayed mechanical dissipation^{110,111}, **d.** high interfacial fatigue threshold of cartilage-/ligament-/tendon-bone interfaces due to intrinsically high-energy phases including nano-crystals and nano-fibers strongly bonded on the interfaces⁷⁸. **a** is reproduced with permission from Ref¹⁰⁸. Copyright 2006 American Chemical Society. **b** is reproduced with permission from is adopted from Ref⁸⁵. Copyright 2013 Elsevier. **c** is reproduced with permission from Ref^{110,111}. Copyright 2014 PLOS and 2018 IntechOpen. **d** is reproduced with permission from Ref⁷⁸. Copyright 2020 Springer Nature.



Figure 3. Hydrogels have been widely used in applications including a. vehicles for drug delivery, b. scaffolds for tissue engineering, c. models for biological studies, d. medical implants, e. wound dressing, f. contact lenses, g. sensors, h. actuators, i. soft robots, j. electronic devices, k. batteries and supercapacitors, I. magnetic devices, m. optical devices, n. acoustic devices, o. water harvesters, p. living devices, q. bioadhesive devices, and r. coatings. a is reproduced with permission from Ref¹¹². Copyright 2015 Springer Nature. **b** is reproduced with permission from Ref¹¹³. Copyright 2019 American Association for the Advancement of Science. c is reproduced with permission from Ref¹⁰. Copyright 2006 Elsevier. d is reproduced with permission from Ref¹¹⁴. Copyright 2015 The Korean Society of Magnetic Resonance in Medicine. g is reproduced with permission from Ref^{21} . Copyright 2006 Springer Nature. h is reproduced with permission from is adopted from Ref¹¹⁵. Copyright 2019 Springer Nature. i is reproduced with permission from Ref²⁶. Copyright 2017 Springer Nature. j is reproduced with permission from Ref¹¹⁶. Copyright 2020 Springer Nature. k is reproduced with permission from Ref³⁴. Copyright 2017 Springer Nature. I is reproduced with permission from Ref²⁸. Copyright 2011 National Academy of Sciences. **m** is reproduced with permission from Ref⁴³. Copyright 2019 Wiley. **n** is reproduced with permission from Ref⁴⁵. Copyright 2020 American Association for the Advancement of Science. o is reproduced with permission from Ref¹⁰⁴. Copyright 2018 Springer Nature. **p** is reproduced with permission from Ref⁴⁶. Copyright 2018 Wiley. \mathbf{q} is reproduced with permission from Ref⁴⁷. Copyright 2020 Springer Nature. \mathbf{r} is reproduced with permission from Ref⁵¹. Copyright 2019 Wiley.

Over the last few decades, intensive efforts have led to the development of a plethora of hydrogels that possess extreme mechanical properties using diverse material candidates, including various natural and synthetic polymers, nano-/micro-/macro-fillers, and nano-/micro-/macro-fibers. Whereas the properties of these hydrogels are remarkable, their design often follows the Edisonian approach – trial and error with specific material candidates. As the field rapidly evolves, emerging applications of hydrogels in biomedicine and machines (**Figure 3**) pose escalating demands on the rationally-guided design of hydrogels beyond the Edisonian approach, so that one can select from diverse material candidates and fabrication methods to design the hydrogels achieving multiple combined extreme properties. However, a set of general principles capable of rationally guiding the design of hydrogels using different materials and fabrication methods for various applications remain a central need in the field of soft materials. In this review, we aim to provide:

i). A set of general principles for the rational design of hydrogels to achieve extreme mechanical properties, including extremely high fracture toughness, tensile strength, resilience, interfacial toughness, fatigue threshold, and interfacial fatigue threshold; and extreme physical properties, including high electrical conductivity, patterned magnetization, high refractive index and transparency, tunable acoustic impedance, and self-healing. The design principles are generally based on fundamental mechanics and

physics (beyond polymers) and/or inspired by biological hydrogels (e.g., muscles, cartilages, tendons, and heart valves) (Figure 4).

- ii). A set of general strategies to implement the design principles discussed in i) with various materials and fabrication methods using *unconventional polymer networks* (UPNs). The UPNs can be broadly categorized into: UPN architectures including ideal polymer networks, polymer networks with slidable crosslinks, interpenetrating and semi-interpenetrating polymer networks, polymer networks with high-functionality crosslinks, and nano-/micro-fibrous polymer networks; and UPN interactions including strong physical crosslinks, weak physical crosslinks, and dynamic covalent crosslinks (**Figure 4**).
- iii). A set of orthogonal design principles and synergistic implementation strategies for the design and fabrication of future hydrogels to achieve multiple combined mechanical, physical, chemical, and biological properties (Figure 4).



Figure 4. This review systematically discusses the design principles and implementation strategies for soft materials including hydrogels, elastomers and organogels to achieve extreme properties.

Notably, because the aforementioned design principles and implementation strategies for hydrogels are based on generic polymer networks, they are also applicable to other soft materials comprised of polymer networks, including elastomers and organogels (i.e., polymer networks infiltrated with organic solvents)¹¹⁷⁻¹¹⁹. In fact, many extreme mechanical and physical properties were first achieved in other soft materials than hydrogels. For example, high values of fracture toughness, tensile strength, resilience and interfacial toughness were realized in elastomers long before in hydrogels; ferromagnetic domains in soft materials were first programmed and 3D printed with elastomeric inks as well.

The review is organized as the following. Section 2 will discuss a variety of natural polymers, synthetic polymers, and permanent covalent crosslinks commonly used for the design and fabrication of hydrogels. Section 3 will introduce conventional polymer networks, and then show that a number of mechanical properties of conventional polymer networks are coupled. Section 4 will define a set of unconventional polymer networks (UPNs), including both UPN architectures and UPN interactions, and then discuss that UPNs can provide decoupled mechanical properties. Thereafter, Sections 5 will systematically discuss the design principles for various extreme mechanical properties of hydrogels and the implement strategies for the design principles using UPNs. Sections 6 will briefly discuss the design principles and implementation strategies for hydrogels to achieve a set of extreme physical properties. In Section 7, we will conclude the review by proposing the orthogonal design principles and synergistic implementation strategies to design future hydrogels that can achieve multiple combined mechanical, physical, chemical and biological properties.

2. Common polymers and crosslinks for hydrogels

A rich library of polymers and crosslinks have been used for the design and fabrication of various hydrogels. These polymers can be broadly categorized into natural polymers and synthetic polymers. In this section, we will briefly discuss the commonly-used natural polymers, synthetic polymers, and permanent covalent crosslinks for hydrogels. We will discuss other types of crosslinks for hydrogels in **Section 4**.

a Examples of common natural polymers for hydrogels



Figure 5. Chemical structures and schematics of typical examples of a. common natural polymers, **b.** common synthetic polymers, and **c.** permanent covalent crosslinks for hydrogels. R represents an organyl substituent or hydrogen.

2.1. Natural polymers for hydrogels

Naturally derived polymers have been widely used to compose the polymer networks of hydrogels (**Figure 5a**). Hydrogels based on natural polymers usually possess properties compatible with biological tissues due to the similarity in their compositions. In addition, the natural polymer networks can often degrade in and be absorbed by the body through metabolism and tissue remodeling processes. Furthermore, the majority of natural polymers have reactive sites amenable to crosslinking and modification, which can endow the corresponding hydrogels with tailored biological and/or mechanical properties. In this subsection, we will briefly discuss a few natural polymers commonly-used for hydrogels. For more detailed discussions, a few classical reviews are recommended^{4-6,68,120}.

Alginate. Alginate is a polysaccharide usually obtained from brown-algae cell walls and two kinds of bacteria, *Azotobacter* and *Pseudomonas*¹²¹. Alginate is known to be a family of linear copolymers containing blocks of β -(1→4)-linked D-mannuronic (M) and α -(1→4)-linked Lguluronic acid (G) residues. The blocks are composed of consecutive G residues (GGGGGGG), consecutive M residues (MMMMM), and alternating M and G residues (GMGMGM)¹²². Alginate hydrogels can be formed with various covalent and physical crosslinks. In particular, the ionic crosslinks have been widely used for alginate hydrogels, because the G blocks¹²³ (and GM blocks¹²⁴) in alginate can be readily bound with one another by divalent cations such as Ca²⁺, Mg²⁺, Ba²⁺, and Sr^{2+ 125-127}. The mechanical properties of alginate hydrogels can be tuned to match those of various biological tissues by changing different parameters, such as the molecular weight, polymer concentration, chemical modification, G/M ratio, and type or density of crosslinks^{123,128}. Alginate hydrogels have been widely used as scaffolds in tissue engineering, such as in intervertebral disk regeneration¹²⁹, adipose tissue regeneration¹³⁰, cardiac regeneration¹³¹, and liver regeneration¹³², since alginate allows the formation of hydrogels under physiological conditions and thus enables easy cell and drug encapsulation.

Hyaluronic acid. Hyaluronic acid (also known as hyaluronan or hyaluronate) is a linear polymer of disaccharides, which are composed of D-glucuronic acid and *N*-acetyl-D-glucosamine, linked together via alternating β -(1 \rightarrow 4)-linked and β -(1 \rightarrow 3)-linked glycosidic bonds^{133,134}. Hyaluronic acid is present in all mammals, especially in various soft connective tissues, acting as a space filler, lubricant, and/or osmotic buffer¹³⁵. Hyaluronic acid can be covalently crosslinked into hydrogels by various hydrazide derivatives^{136,137}. The abundant carboxyl and hydroxyl groups on the polysaccharide structure of hyaluronic acid also offer many active sites for chemical modifications¹³⁸. For example, hyaluronic acid can be modified with thiol^{139,140}, haloacetate¹⁴¹, dihydrazide^{136,142}, aldehyde^{143,144} and tyramine¹⁴⁵ groups, which can react with corresponding covalent crosslinkers through addition or condensation reactions¹⁴⁶. As another example, hyaluronic acid can also be modified by methacrylic anhydride or glycidyl methacrylate to possess reactive methacrylic groups, which can be polymerized by radical polymerization¹⁴⁷⁻¹⁴⁹. Owing to the naturally-derived, nonimmunogenic, biodegradable, and non-adhesive properties¹⁵⁰⁻¹⁵², hyaluronic acid hydrogels have been widely used as scaffolds in cell therapy and tissue engineering,

such as in cell delivery¹⁵³, molecule delivery^{154,155}, stem cell therapy^{156,157}, cartilage engineering^{154,158}, cardiac repair¹⁵⁹, and valvular engineering¹⁶⁰.

Collagen. Collagen is one of the major proteins in animal bodies. There are approximately 29 types of collagens discovered so far¹⁶¹. The structures of collagens can be defined at different levels, including primary structure (amino acid triplet), secondary structure (α -helix), tertiary structure (triple helix), and quaternary structure (fibril) ^{162,163}. The primary structure of collagen is the tripeptide sequence of $-(Gly-X-Y-)_n-$, where Gly is glycine, X and Y are other amino acids than Gly. The sequence of the amino acids governs how the peptide folds into a secondary structure, mainly left-handed α -helix, which is stabilized by the hydrogen bonds between amino acid residues¹⁶⁴. Three left-handed α -helix polypeptide chains then form a tertiary structure via the aldol condensation crosslinking, aldehyde amine condensation crosslinking, and aldol histidine crosslinking¹⁶⁵. The triple strands can further self-assemble into a collagen fiber as the quaternary structure¹⁶⁶.

Acid-solubilized collagens can self-assemble to form physically-crosslinked hydrogels when the collagen solutions are neutralized and heated. Since the physically-crosslinked collagen hydrogels are usually mechanically weak and thermally unstable^{167,168}, they have been strengthened and stabilized with chemical crosslinks such as glutaraldehyde, genipin, carbodiimides, and diphenylphosphoryl azide¹⁶⁹⁻¹⁷¹. Collagens can be biodegraded by collagenases and metalloproteases; the crosslinked collagens usually have slower degradation rates than the uncrosslinked collagens¹⁷². Because collagens usually have low antigenicity, low inflammatory response, good biocompatibility, and natural cell-adhesive motifs¹⁷³⁻¹⁷⁵, collagen hydrogels have been widely used as scaffolds for drug and protein delivery^{176,177} and for reconstructions of liver¹⁷⁸, skin¹⁷⁹, blood vessel¹⁸⁰, small intestine¹⁸¹, cartilage¹⁸², vocal cord¹⁸³, and spinal cord¹⁸⁴.

Gelatin. Gelatins are naturally derived polymers obtained through breaking the triple-helix structures of collagens into single-strand molecules. There are two types of gelatins, type A and type B, which are obtained with acid and alkaline treatments of collagens, respectively¹⁸⁵. Gelatins can be physically crosslinked by simply reducing the temperature of aqueous solutions of gelatins below a certain temperature^{186,187}. The physically-crosslinked gelatins are usually unstable for long-term biomedical applications under physiological conditions. To further stabilize the physically-crosslinked gelatin hydrogels, covalent crosslinkers¹⁸⁸ such as aldehydes (e.g., formaldehyde, glutaraldehyde, and glyceraldehyde)^{189,190}, polyepoxides¹⁹¹, and isocyanates¹⁹² have been widely used to react with and bridge the free amine groups (from lysine and hydroxylysine) and free carboxylic acids (from glutamic and aspartic acid) on the gelatin molecules. Besides the introduction of covalent crosslinkers, the gelatin backbones can also be modified by methacrylates to form covalently-crosslinked gelatin methacryloyl hydrogels¹⁹³. In addition, synthetic polymers can also be coupled on gelatin chains through the grafting-from¹⁹⁴, grafting-to¹⁹⁵ and grafting-through¹⁹⁶ methods to enhance the mechanical properties of gelatin hydrogels. Furthermore, the gelatin molecules tend to form physical interactions with various dopants, such as carbon nanotubes¹⁹⁷, graphene oxide¹⁹⁸, inorganic nanoparticles, and minerals^{199,200}. The aforementioned covalent crosslinks, modifications, and interactions can significantly improve the mechanical properties of gelatin hydrogels^{189,201}. The easy gelation process and the excellent biocompatibility of gelatin hydrogels make them attractive for biomedical applications, such as drug delivery²⁰² and tissue engineering^{203,204}.

Fibrin. Fibrin is a naturally-derived polymer obtained from thrombin-treated fibrinogen²⁰⁵. Fibrin is involved in the natural wound healing process by forming extensive fibrous networks. Fibrin can form clots or hydrogels when mixing fibrinogen and thrombin solutions at the room temperature²⁰⁶. The resultant fibrin hydrogels usually have weak mechanical properties due to the nature of physical crosslinks. To improve the mechanical properties of fibrin hydrogels, chemical crosslinkers such as genipin can be introduced to crosslink the amine residues on fibrin proteins, which subsequently form stable covalently-crosslinked networks²⁰⁷. In addition, fibrin hydrogels can also be combined with synthetic polymers such as polyurethane²⁰⁸, polycaprolactone²⁰⁹, β -tricalciumphosphate²¹⁰, and polyethylene glycol²¹¹ to enhance the mechanical strength of the hydrogels. Fibrin hydrogels have been widely used as tissue sealants and adhesives to control bleeding in surgery²¹², and as scaffolds for cardiac tissue engineering²¹³, neurological regeneration²¹⁴, ocular therapy²¹⁵, cartilage and bone reparation^{216,217}, muscle tissue engineering²¹⁸, and exogenous delivery in wound healing²¹⁹. In particular, fibrin hydrogels can be produced autologously from a patient's own blood, thereby reducing the risk of foreign-body reactions²²⁰.

Agarose. Agarose is a neutral polysaccharide composed of β -D-galactopyranosyl and 3,6anhydro- α -L-galactopyranosyl, mainly extracted from red algae (Rhodophyceae)²²¹. As a thermoresponsive polymer, agarose can be heated to dissolve in water, and then cooled down to form a hydrogel. During this gelation process, the agarose structure changes from a random-coil configuration to bundles of associated double helices with multiple-chain aggregation in the junction zone^{222,223}. The gelling temperature and mechanical properties of agarose hydrogels can be tuned by changing the concentration, molecular weight, and structure of the agarose in the hydrogels^{224,225}. Agarose hydrogels have been used as scaffolds for cell encapsulation²²⁶, cartilage reparation²²⁷, and nerve regeneration²²⁸, due to its low immunoreaction in human bodies²²⁹. Notably, since the native agarose does not possess cell adhesion motifs, cell adhesion peptides have been covalently conjugated to the agarose backbone to enhance the interactions between cells and agarose hydrogels²³⁰.

Chitosan. Chitosan is a linear polysaccharide composed of β -(1→4)-linked D-glucosamine and *N*-acetyl-D-glucosamine. Chitosan is mainly prepared by partial deacetylation of chitin (obtained from crab and shrimp shells) to less than 40% of *N*-acetyl-D glucosamine residues^{231,232}. The physical, chemical and biological properties of chitosan materials are highly related to the molecular weight and the degree of deacetylation of chitosan^{233,234}. Chitosan can form physicallycrosslinked hydrogels by hydrophobic interaction, hydrogen bonding^{235,236}, metal coordination (with metal ions such as Pt(II), Pd(II), Mo(VI)^{237,238}), and electrostatic interaction (with multivalent anions such as sulfate, citrate and phosphate ions^{239,240}; with anionic polyelectrolytes²³¹ such as polysaccharides^{241,242}, proteins^{243,244} and synthetic polymers²⁴⁵). These physically-crosslinked chitosan hydrogels usually have weak mechanical properties and short lifetime, which are also highly influenced by pH, temperature and environments^{232,246}. To enhance the mechanical properties and stability of chitosan hydrogels, covalent crosslinkers have been introduced into the hydrogels. The commonly-used covalent crosslinkers include dialdehydes^{247,248}, formaldehyde²⁴⁹, di-glycidyl ether²⁵⁰, and genipin^{251,252}, which can react with the residual functional groups (such as OH, COOH, and NH₂) on chitosan backbones to form the amide bonds, ester bonds, and Schiff base linkages^{235,253,254}. In addition, chitosan can also be modified with methacrylate or aryl azide groups to form photo-crosslinkable macromers²⁵⁵. The gelation degree and mechanical properties of these photo-crosslinkable chitosan hydrogels can be controlled by UV irradiation time and intensity²⁵⁶⁻²⁵⁸. Furthermore, chitosan hydrogels can be modified with biofunctional ligands such as Arg-Gly-Asp (RGD) peptides to facilitate cell adhesion and proliferation^{259,260}. Chitosan hydrogels have been widely used in biomedical applications such as drug delivery²⁶¹, cell encapsulation²⁶², neural tissue engineering²⁶³, and bone regeneration²⁶², owing to their excellent biocompatibility and biodegradability²⁶⁴.

Cellulose. Cellulose is the most abundant natural polysaccharide, and the main constituent of plants and natural fibers such as cotton and linen²⁶⁵⁻²⁶⁷. Some bacteria such as acetobacter xylinum are also able to produce cellulose²⁶⁸. Cellulose consists of β -(1 \rightarrow 4)-linked D-glucose units, which result in cellulose's high crystallinity (over 40%) and difficulty in dissolving in water and other common solvents²⁶⁹. Solvents such as *N*-methylmorpholine-*N*-oxide^{270,271}, ionic liquids^{272,273}, and alkali/urea (or thiourea) aqueous systems^{274,275} have been developed to dissolve native cellulose. Cellulose can also be modified through partial esterification or etherification of the hydroxyl groups on the backbone²⁶⁵. These cellulose derivatives, including methyl cellulose²⁷⁶, hydroxypropyl cellulose²⁷⁷, hydroxypropylmethyl cellulose^{278,279}, and carboxymethyl cellulose²⁸⁰ are easier to dissolve and process compared to the native cellulose.

Cellulose and its derivatives can be chemically crosslinked to form stable threedimensional networks. Bifunctional and multifunctional molecules, such as 1,2,3,4butanetetracarboxylic dianhydride²⁸¹, succinic anhydride²⁸², citric acid²⁸³, epichlorohydrin²⁸⁴, ethylene glycol diglycidyl ether²⁸⁵, and divinyl sulfone²⁸⁶ can form covalent ester or ether bonds between cellulose chains. Cellulose chains can also be covalently crosslinked by the irradiation of electron beams and gamma rays^{287,288}, which avoids the usage of toxic crosslinkers and allows the simultaneous sterilization of the resultant hydrogels. Cellulose and its derivatives can also be blended with natural polymers, such as chitosan²⁸⁹, starch²⁹⁰, alginates²⁹¹ and hyaluronic acid²⁹², or synthetic polymers such as polyethylene glycol²⁹³, polyvinyl alcohol²⁹⁴ and poly(*N*,*N*dimethylacrylamide)²⁹⁵ to form interpenetrating polymer networks with excellent mechanical properties. Notably, bacterial cellulose produced from certain bacterial species such as acetobacter xylinum can directly form cellulose hydrogels with high purity and tensile strength^{296,297}. Since cellulose-based hydrogels are proven to have superior hydrophilicity, biodegradability, biocompatibility, and transparency, they have been widely used in drug delivery²⁹⁸, tissue engineering²⁹⁹, blood purification³⁰⁰, strain sensor³⁰¹ as well as water purification³⁰².

2.2. Synthetic polymers for hydrogels

In addition to natural polymers, synthetic polymers have been widely used for the design and fabrication of hydrogels (**Figure 5b**). The synthetic polymer networks of hydrogels are commonly formed by copolymerization of monomers for the polymer backbones and crosslinkers, or by reactions of synthetic polymers, macromers and/or crosslinkers.

Poly(acrylic acid). Poly(acrylic acid) (PAA) is a linear polymer prepared by radical polymerization of acrylic acid monomers. The backbone of PAA contains a large number of carboxyl groups. PAA can form hydrogels through covalent and physical crosslinking. Covalently-crosslinked PAA hydrogels are usually formed by copolymerization of di-/multi-vinyl crosslinkers together with acrylic acid monomers³⁰³. In addition, the carboxyl groups of PAA can form physical interactions with various doping agents such as clay³⁰⁴, graphene oxide³⁰⁵ and cations³⁰⁶, which can act as physical crosslinks for PAA hydrogels; the carboxyl groups can also form hydrogen bonds between PAA chains and introduce self-healing or self-adhesive properties to PAA hydrogels³⁰⁷. Furthermore, the abundant carboxyl groups on PAA can associate with water molecules to facilitate the absorption of water by PAA hydrogels¹¹⁵. Since the carboxyl groups are sensitive to pH and ionic strength, the equilibrium swelling ratio of PAA hydrogels is affected by the pH and ionic strength of the solutions for the hydrogels^{308,309}. PAA hydrogels can also incorporate other linear polymers, such as biological polymers, to form various adhesives and hydrogels for biomedical applications^{47,310}.

Poly(2-hydroxyethyl methacrylate). Poly(2-hydroxyethyl methacrylate) (PHEMA) hydrogels can be prepared by free-radical polymerization of 2-hydroxyethyl methacrylate (HEMA) monomers with covalent crosslinkers such as trimethylene glycol dimethacrylate (TEGDMA), initiators such as sodium pyrosulfite (SMBS) and ammonium persulfate (APS). The HEMA monomers can also be copolymerized with acrylic or acrylamide monomers to control the swelling and mechanical properties of the resultant hydrogels³¹¹. PHEMA hydrogels are optically transparent and mechanically stable in the physiological environments. Pure PHEMA hydrogels are also resistant to cell adhesion and difficult to degrade in the physiological environments; however, various biofunctional and bioactive motifs can be coupled onto the hydrogels to improve their cell interactions and degradability^{312,313}. PHEMA hydrogels are famous for their ophthalmic applications such as contact lens³¹⁴ and artificial cornea³¹⁵.

Poly(vinyl alcohol). Poly(vinyl alcohol) (PVA) is mainly obtained from the partial hydrolysis of poly(vinyl acetate)³¹⁶. PVA can form stable and elastic hydrogels through either physical or covalent crosslinking^{317,318}. The physically-crosslinked PVA hydrogels are commonly obtained by repeated freezing and thawing of PVA solutions³¹⁹, which gives elastic, tough, strong and fatigue-resistant PVA hydrogels^{76,78,320}. PVA can also be covalently crosslinked through the use of difunctional crosslinkers such as glutaraldehyde, epichlorohydrin, boric acid and dialdehyde^{321,322}. Electron-beam and gamma irradiation can also crosslink PVA to avoid residual covalent crosslinkers in the hydrogels³²³. Pure PVA hydrogels are non-adhesive to cells, but

several oligopeptide sequences can be conjugated onto the backbones of PVA hydrogels to enhance their cellular interactions³²⁴. PVA hydrogels have been extensively studied and used in biomedical applications^{325,326}, such as articular cartilage replacement and regeneration^{327,328}.

Poly(ethylene glycol) or poly(ethylene oxide). Poly(ethylene glycol) (PEG) is usually obtained from the anionic or cationic polymerization of ethylene oxide. When the PEG has a molecular weight more than 10 kDa, it is also named poly(ethylene oxide) (PEO) since the end groups are negligible³²⁹. There are various methods to crosslink PEG polymers into hydrogels. The ends of PEG chains can be modified with unsaturated groups, such as acrylate or methacrylate ends, and then be used as macro-crosslinkers to form hydrogels with other unsaturated monomers by the photo-/UV-induced radical polymerization^{330,331}. PEG can also form hydrogels by electron beam irradiation via radiation-induced free radical processes³³². Furthermore, the end groups of the PEG chain can be modified with various reactive pairs, such as *N*-hydroxysuccinimide/NH₂³³³, maleimide/thiol³³⁴, and acetylene/azide³³⁵. Since these functional chain-end motifs usually have high reaction efficiency and fast reaction kinetics, the obtained hydrogels by the coupling reactions of these groups can give relatively well-defined network architectures⁷⁴.

PEG polymers can also form physically-crosslinked networks. Similar to the chemical crosslinking method, the ends of PEG chains can be modified with various motifs for physical crosslinking. For example, nucleobase pairs of adenines and thymines³³⁶, ureido-pyrimidinone (UPy) units³³⁷, or host-guest molecules³³⁸ can be introduced onto the chain-ends of PEG molecules to prepare physically-crosslinked PEG hydrogels. These physically-crosslinked PEG hydrogels can exhibit switchable, self-healable or stimuli-responsive properties and high mechanical strength³³⁹. Besides the modification and utilization of chain-end groups, physically-crosslinked PEG hydrogel can also be prepared by using PEG block copolymers³⁴⁰. PEG-b-PPG (poly(propylene glycol)) is one of the most widely used PEG-derived block copolymers to prepare thermo-responsive physical hydrogels³⁴¹. These physical hydrogels are formed by the hydrophobic interaction of the PPG blocks. The phase transition behavior of these hydrogels can be optimized by balancing the hydrophobic PPG blocks and the hydrophilic PEG blocks. Based on the same gelation mechanism, PEG block copolymers with poly(DL-lactic acid) (PDLLA)³⁴², poly(dl-lactic acid-co-glycolic acid) (PLGA)^{343,344}, polylactide (PLA)³⁴⁵, poly(caprolactone)(PCL)³⁴⁶ and poly(propylene sulfide)(PPS)³⁴⁷ can also form physically-crosslinked hydrogels with injectable or stimuli-responsive properties. PEG, as well as its derivatives, are widely used in biomedical applications due to their non-toxic and non-immunogenic properties³⁴⁸. While the inert biological property of PEG hydrogels can prevent undesired interactions between native PEG hydrogels and cells^{349,350}, PEG hydrogels can also be modified with various bioactive conjugations such as growth factors ³⁵¹ and cell-adhesive peptides³⁵² through Michael-type addition^{353,354}or click chemistry³⁵⁰. PEG hydrogels with these bioactive molecules can facilitate their biomedical applications³⁵⁵ such as drug or cell delivery^{356,357} and tissue engineering³⁵⁸.

Poly (N-isopropylacrylamide). Acrylamide and its derivatives have been widely used to prepare hydrogels by radical copolymerization with crosslinkers. One interesting hydrogel based

on acrylamide and its derivatives is the poly(*N*-isopropylacrylamide) (PNIPAm) hydrogel. Uncrosslinked linear PNIPAm exhibits a coil-to-globule phase transition in aqueous solutions when the temperature is raised above a critical temperature^{359,360}. The PNIPAm can be covalently crosslinked by crosslinkers such as bis-acrylamide derivatives through the radical polymerization process. The crosslinked PNIPAm hydrogels also possess the reversible thermo-responsive behavior with a critical temperature of around 34 °C ³⁶¹, above which the hydrogel structure will collapse and exude water^{362,363}. While the thermo-responsive behavior of PNIPAm hydrogels is usually slow, many studies have improved the phase-transition speed of PNIPAm hydrogels by incorporating porous structures during the hydrogel formation^{364,365}. The thermo-responsive PNIPAm hydrogels can be used as actuators for soft robotics³⁶⁶, injectable scaffolds for tissue engineering³⁶⁷, and thermo-responsive substrates for on-demand detachment of cell sheets^{368,369}.

Silicone. Silicone hydrogels are hydrogels that contain silicone polymers as one of its polymer components³⁷⁰. Silicone polymers are commonly hydrophobic³⁷¹. In order to form silicone hydrogels, hydrophilic monomers and/or polymers have been introduced into the silicone matrix by blending or copolymerization to improve the hydrophilicity of silicone hydrogels^{14,372,373}. For example, hydrophilic polymers such as PHEMA can be blended directly into the silicone polymer matrix, forming a hydrophilic interpenetrating polymer network³⁷⁴. Hydrophilic monomers such as *N*-vinylpyrrolidone (NVP) can be copolymerized with silicon-macromers to form hydrophilic silicone hydrogels³⁷⁵. Hydrophilic polymer segments such as PEG³⁷⁶ can also be copolymerized onto silicone segments to form block-modified^{377,378} or graft-modified³⁷⁹ hydrophilic silicone hydrogels. Since these hydrophilic silicone hydrogels usually have excellent gas permeability as well good biocompatibility, they have been used in biomedical applications such as contact lenses^{14,380}, histological engineering materials^{381,382}, and drug-delivery carriers^{383,384}.

2.3. Permanent covalent crosslinks for hydrogels

In this section, we will discuss permanent covalent crosslinks that are commonly used in hydrogels (**Figure 5c**). We will discuss other types of crosslinks in **Section 4**. The energy of permanent covalent crosslinks ranges from 220 kJ mol⁻¹ to 570 kJ mol⁻¹ (**Figure 6**)³⁸⁵⁻³⁸⁷.



Figure 6. Bond energies of various types of permanent covalent crosslinks³⁸⁵⁻³⁸⁷, weak physical crosslinks³⁸⁸⁻³⁹¹, and dynamic covalent crosslinks^{387,392-396}.

Carbon-carbon bonds. The energy of the carbon-carbon bond is around 300 kJ mol⁻¹ to 450 kJ mol^{-1 385-387}. Hydrogels covalently crosslinked by carbon-carbon bonds are usually formed by radical copolymerization of monomers and di-/multi-vinyl crosslinkers. The crosslinkers can be small molecules with two double bonds such as N,N'-Methylenebisacrylamide (MBA) or macromolecules with several acrylate groups^{6,397}. These crosslinkers are compatible with various initiation and polymerization systems^{4,254,398}. For example, photo-radical initiators can be added the pre-polymerization solution together with monomers and di-/multi-vinyl into crosslinkers^{318,399,400}. Once the initiator is irradiated by UV light, radicals will be generated to initiate the polymerization of the double bonds on monomers as well as crosslinkers^{401,402}. As a result, hydrogels can be formed in-situ and with patterned structures or biology functions^{399,403}. The polymerization of vinyl monomers and crosslinkers can also be carried out with a system composed of peroxydisulfate and N, N, N'N'-tetramethylene-diamine (TEMED), where TEMED can accelerate the decomposition of peroxydisulfate to generate a large number of radicals⁴⁰⁴. This initiation and polymerization system can effectively and rapidly form various hydrogels under room temperature.

The carbon-carbon crosslinks of hydrogels can also be formed by high-energy irradiation (e.g., gamma and electron beams). Similar to UV light, high-energy radiation can be used to polymerize unsaturated compounds such as monomers and crosslinkers with vinyl groups or acrylate groups^{405,406}. High-energy radiation can also crosslink polymers without unsaturated bonds⁴⁰⁷, because radicals can be generated from the homolytic scission of the polymer chains under high-energy radiation. The radiolysis of water molecules in the solvent can also generate hydroxyl radicals that attack polymer chains to form macroradicals⁴⁰⁸. These radicals can then

undergo recombination and termination to form covalent polymer networks crosslinked by carboncarbon bonds.

Carbon-nitrogen bonds. The energy of the carbon-nitrogen bond is around 300 kJ mol⁻¹ to 430 kJ mol^{-1 385-387}. Hydrogels covalently crosslinked by carbon-nitrogen bonds are usually formed by highly effective chemical reactions of complementary groups. For example, the amide bonds have been widely used as the covalent crosslinks for hydrogels by the condensation reactions between amines with carboxylic acids and derivatives⁴⁰⁹. N-hydroxysuccinimide (NHS) and N,N-(3-dimethylaminopropyl)-N-ethyl carbodiimide (EDC) are widely used to facilitate the condensation reaction of amines with carboxylic acids⁷⁴. The addition of NHS and EDC will also suppress possible side-reactions and give better control of the crosslink density in the hydrogels⁴¹⁰. The carbon-nitrogen bonds can also be formed through the addition reactions of amines with electrophiles such as adipic acid dihydrazide and diisocyanate crosslinkers^{6,120,411}. These difunctional crosslinkers have been widely used to crosslink natural macromolecules due to the high reaction efficiency. The mechanical properties of the resultant hydrogels can be controlled by tuning the concentration and ratio of the polymers and the crosslinking agents. Another category of reactions that can form carbon-nitrogen crosslinks for hydrogels is the azide-alkyne cycloaddition reaction, which is one typical click reaction to connect alkyne and azide into triazole. The click reaction has high efficiency without side reactions⁴¹². Furthermore, the azide-alkyne cycloaddition can be conducted in the absence of metal catalysis⁴¹³, expanding the applicability of the azide-alkyne cycloaddition for preparing biocompatible hydrogels.

Carbon-oxygen bonds. The energy of the carbon-oxygen bond is around 280 kJ mol⁻¹ to 370 kJ mol⁻¹ ³⁸⁵⁻³⁸⁷. The most common carbon-oxygen bond is the ester bond formed by the reactions between hydroxyl groups and carboxylic acids or derivatives⁴¹⁴. Such ester crosslinks can be hydrolyzed easily and make the hydrogels degradable under ambient temperature and physiological conditions. Besides the ester crosslinks, the carbon-oxygen bonds are also present in ether groups and urethane groups, which can become crosslinks due to the reaction between side groups on polymers (such as hydroxyl groups on polysaccharides and PVA) and reactive crosslinkers (such as glutaraldehyde^{415,416}, divinyl sulfone⁴¹⁷, dibromide⁴¹⁸ and diisocyanate⁴¹⁹).

Carbon-sulfide bonds. The energy of the carbon-sulfide bond is around 220 kJ mol⁻¹ to 310 kJ mol^{-1 387}. The covalent crosslinking of hydrogels through carbon-sulfide bonds are mainly formed by the thiol-click reactions^{420,421}. The inherent electron density of the sulfide atom makes thiols prone to react with many functional groups through a radical or catalyzed process^{422,423}. Thiol groups can be easily converted into nucleophilic thiolates or electrophilic thiyl radicals, which then proceed with nucleophilic reactions or radical chain processes to achieve the thiol-click reactions⁴²⁴. Specifically, for the radical thiol-click reactions, the thiol group can be activated by heat and/or UV light to generate radicals that initiate the radical-mediated thiol-ene or thiol-yne reactions⁴²⁵. For the nucleophilic thiol-click reactions initiated by strong bases, the thiol groups can readily react with electron-poor ene-functional compounds through the Michael addition, with isocyanates derivatives through carbonyl addition, with halide through S_N2 nucleophilic

substitution, and with epoxy motifs through S_N2 ring-opening reactions⁴²⁶⁻⁴²⁹. The thiol-click reactions commonly have high efficiency and high conversion rate without any side products, even in the presence of water, ions, and oxygen. The thiol-click reactions have been extensively used to prepare hydrogels for various biomedical applications^{430,431}.

Silicon-oxygen bonds. The energy of the silicon-oxygen bond is around 420 kJ mol⁻¹ to 570 kJ mol^{-1 385-387}. The silicon-oxygen bonds are mainly used in the formation of silicone-based hydrogels^{376,432,433}, and can usually enhance the mechanical properties of the silicone-based hydrogels⁴³⁴. In addition, silicon-oxygen bonds have been widely used to form strong bonding between hydrogels and diverse engineering materials with modified surfaces such as salinized surfaces⁴⁹.

3. Conventional polymer networks

As illustrated in **Figure 7**, a conventional polymer network is defined as polymer chains crosslinked via permanent covalent bonds into a network, in which entanglements, physical crosslinks and reversible crosslinks of the polymer chains are negligible^{54,57,62}. Conventional polymer networks have provided the basic models for the development of unentangled rubber elasticity, including the affine network model and the phantom network model^{54,57,62}. Conventional polymer networks have also been widely adopted in synthetic hydrogels, although biological hydrogels (**Figures 1** and **2**) generally reply on unconventional polymer networks which will be discussed in **Section 4**.



Figure 7. Schematics of a conventional polymer network a. in the dry state and b. covalently bonded on a substrate, and c. in the swollen state and d. covalently bonded on a substrate.

3.1. Conventional polymer networks in the dry state

In the dry state (Figure 7a), a conventional polymer network contains *n* polymer chains per unit volume, where a polymer chain is defined as the segment of polymer between two successive covalent crosslinks. Each polymer chain contains *N* Kuhn monomers, and the length of each Kuhn monomer is *b*. The end-to-end distances of a polymer chain at the relaxed and fully stretched states are \sqrt{Nb} and *Nb*, respectively. Therefore, the stretch limit of the polymer chains λ_{im} in the dry polymer network can be calculated as^{54,57,62}

$$\lambda_{\rm lim} = \frac{Nb}{\sqrt{Nb}} = N^{1/2} \tag{1}$$

The stretch limit of the bulk polymer network scales with the chain stretch limit λ_{lim} , and the prefactor of the scaling relation depends on the polymer network architecture⁶¹.

Assuming that the dry polymer network follows the affine network model, the shear modulus of the network under initial deformation can be expressed as^{54,57,62},

$$G = nkT \tag{2}$$

where k is the Boltzmann constant and T is the absolute temperature.

Following the Lake-Thomas model⁵⁶, the fracture toughness of the dry polymer network is its intrinsic fracture energy Γ_0 , which is the energy required to fracture a single layer of polymer chains per unit area,

$$\Gamma_0 = n\sqrt{Nb} \cdot NU_f = nbN^{3/2}U_f \tag{3}$$

where $n\sqrt{N}b$ is the number of polymer chains per unit area, NU_f is the energy required to fracture a polymer chain, and U_f is the energy required to fracture a single Kuhn monomer.

Also based on the Lake-Thomas model^{55,56}, the fatigue threshold of the dry polymer network is the intrinsic fracture energy Γ_0 . If the dry polymer network is covalently bonded on a substrate (**Figure 7b**), both the interfacial toughness and the interfacial fatigue threshold of the adhesion are on the level of Γ_0 as well^{49,78,435}.

By substituting the typical values of b, N, n, kT and U_f into **Eqs. 1-3**, we can estimate that the shear modulus G can be on the order of kilopascals to megapascals, the chain stretch limit λ_{lim} can reach up to a few tens (without entanglement), and the intrinsic fracture energy Γ_0 can reach up to a few hundreds of joule per meter squared⁶².

The mechanical properties of the dry polymer network are coupled with one another. It is commonly assumed that the polymer chains occupy the major volume of the polymer network in the dry state, and therefore the volume conservation of the polymer network gives

$$Nnv = 1$$
 (4)

where *v* is the volume of a Kuhn monomer.

By substituting Eq. 4 into Eqs. 1-3, we can express the chain stretch limit λ_{lim} , shear modulus *G* and intrinsic fracture energy Γ_0 of a conventional polymer network in the dry state as functions of its chain length *N*,

$$\lambda_{\rm lim} = N^{1/2}, \ G = N^{-1} v^{-1} kT, \ \Gamma_0 = N^{1/2} v^{-1} b U_f$$
(5)

From Eq. 5, it is evident that enhancing the chain length N increases the chain stretch limit λ_{lim} and the intrinsic fracture energy Γ_0 but decreases the shear modulus G of the

conventional polymer network in the dry state. These mechanical properties of the conventional polymer network in the dry state are coupled through the following relation,

$$\lambda_{\rm lim} \sim \Gamma_0 \sim G^{-1/2} \tag{6}$$

3.2. Conventional polymer networks in the swollen state

A dry conventional polymer network with the parameters discussed in Section 3.1 can imbibe water and swell into a hydrogel composed of the conventional polymer network and water (Figure 7c). The swelling of the dry polymer network stretches polymer chains in the network by a ratio of λ_s , named the chain stretch of swelling.

Since the swelling of the dry polymer network stretches its polymer chains by a ratio of λ_s , the end-to-end distance of a polymer chain in the hydrogel at the relaxed and fully stretched states are $\lambda_s \sqrt{Nb}$ and Nb, respectively. Therefore, the stretch limit of polymer chains λ_{lim} in the hydrogel can be calculated as,

$$\lambda_{\rm lim} = \frac{Nb}{\lambda_s \sqrt{Nb}} = N^{1/2} \lambda_s^{-1} \tag{7}$$

The stretch limit of the bulk hydrogel scales with the chain stretch limit λ_{lim} , and the pre-factor of the scaling relation depends on the polymer network architecture⁶¹.

The swelling of the dry polymer network reduces its shear modulus by a ratio of λ_s ⁵⁴. Therefore, the shear modulus of the hydrogel under initial deformation can be expressed as,

$$G = nkT\lambda_s^{-1} \tag{8}$$

Note that n in Eq. 8 is the number of polymer chains per unit volume of the dry polymer network.

The swelling of the dry polymer network reduces the number of polymer chains per unit area by a ratio of λ_s^2 but does not significantly change the energy required for fracturing a polymer chain in the network. Therefore, the intrinsic fracture energy Γ_0 of the hydrogel can be calculated as,

$$\Gamma_0 = \frac{n\sqrt{N}b}{\lambda_s^2} \cdot NU_f = nbN^{3/2}U_f \lambda_s^{-2}$$
(9)

The fracture toughness and fatigue threshold of a hydrogel with the conventional polymer network are the hydrogel's intrinsic fracture energy $\Gamma_0^{55,56}$. If the hydrogel's polymer network is covalently bonded on a substrate (**Figure 7d**), both the interfacial toughness and the interfacial fatigue threshold of the adhesion are on the order of the hydrogel's intrinsic fracture energy Γ_0 as well^{49,78,435}.

By comparing Eqs. 1-3 and Eqs. 7-9, we can see that swelling the dry polymer network into the hydrogel reduces the chain stretch limit λ_{lim} , shear modulus G and intrinsic fracture

energy Γ_0 of the dry network by factors of λ_s , λ_s and λ_s^2 , respectively^{54,57,62}. By substituting the typical values of λ_s , b, N, n, kT and U_f into **Eqs. 7-9**, we estimate that the shear modulus G of the hydrogel with the conventional polymer network can be on the order of pascals to megapascals, the chain stretch limit λ_{lim} can reach up to a few times (without entanglement), and the intrinsic fracture energy Γ_0 can reach a few tens of joule per meter squared.

By substituting **Eq. 4** into **Eqs. 7-9**, we can express the chain stretch limit λ_{lim} , shear modulus *G* and intrinsic fracture energy Γ_0 of the hydrogel with the conventional polymer network as functions of its chain length *N*,

$$\lambda_{\rm lim} = N^{1/2} \lambda_s^{-1}, \ G = N^{-1} v^{-1} k T \lambda_s^{-1}, \ \Gamma_0 = N^{1/2} v^{-1} b U_f \lambda_s^{-2}$$
(10)

From Eq. 10, it is evident that enhancing the chain length N increases the chain stretch limit λ_{lim} and the intrinsic fracture energy Γ_0 but decreases the shear modulus G of the hydrogel with the conventional polymer network. These mechanical properties of the hydrogel are coupled through the following relation,

$$\lambda_{\rm lim} \sim \Gamma_0 \sim G^{-1/2} \tag{11}$$

Notably, the chain stretches due to equilibrium swelling of a conventional polymer network can be calculated. Without loss of generality, let consider a dry conventional polymer network with a cubic shape. When the polymer network reaches the equilibrium state in water, one side of the cube increases its length from the dry state by a ratio of λ_{eq} . At the equilibrium state, the Helmholtz free energy for stretching polymer chains $W_{stretch}$ and for mixing polymers and water W_{mix} per unit volume of the dry polymer network can be expressed as^{54,436},

$$W_{stretch} = \frac{1}{2} nkT \left(3\lambda_{eq}^2 - 3 - 2\log \lambda_{eq}^3 \right)$$
(12a)

$$W_{mix} = -\frac{kT}{v_s} \left[\left(1 - \lambda_{eq}^3 \right) \log \left(1 - \lambda_{eq}^{-3} \right) + \chi \lambda_{eq}^{-3} \right]$$
(12b)

where χ is the Flory polymer-solvent interaction parameter and v_s is the volume of a solvent molecule. Subsequently, the Helmholtz free energy per unit volume of the dry polymer network can be expressed as^{54,436},

$$W = W_{stretch} + W_{mix} \tag{13}$$

When the polymer network reaches the equilibrium state in water, λ_{eq} minimizes the Helmholtz free energy^{54,436}, giving

$$\frac{\partial W}{\partial \lambda_{eq}} = 0 \tag{14}$$

By solving Eq. 14, one can obtain λ_{eq} of the hydrogel at the equilibrium swollen state. The stretch of polymer chains λ_s in the hydrogel scales with λ_{eq} , and the pre-factor of the scaling relation depends on the polymer network architecture⁶¹. While Eqs. 12-14 assume that the polymer network of the hydrogel is uncharged, the effect of charges on the equilibrium swelling of hydrogels can be accounted for by introducing additional terms to the Helmholtz free energy function, Eq. 12^{54,437}. It should be noted that hydrogels are not necessary to reach the equilibrium sweller or do not have sufficient time to equilibrate with water.

4. Unconventional polymer networks

Section 3 has established that elastomers and hydrogels with conventional polymer networks have intrinsically coupled mechanical properties, including shear modulus, stretch limit, fracture toughness, fatigue threshold, interfacial toughness of adhesion, and interfacial fatigue threshold of adhesion (Eqs. 6 and 11). This section will discuss unconventional polymer networks (UPNs), which constitute most biological hydrogels (Figures 1 and 2) and have been widely used in synthetic hydrogels to achieve extreme mechanical properties.

The UPNs are defined as polymer networks that are different from the conventional polymer networks in terms of the architectures of the polymer networks and/or the interactions among polymer chains in the networks^{70,438-446}. Therefore, the UPNs can be broadly classified into two categories: the UPN architectures and the UPN interactions.

4.1. Unconventional polymer network architectures

As illustrated in **Figure 8**, the UPN architectures are distinctly different from the architecture of the conventional polymer networks composed of randomly-crosslinked polymer chains with covalent bonds (**Figure 5**). Almost all biological tissues (**Figures 1** and **2**) possess certain types of UPN architectures. Over the last few decades, multiple UPN architectures have been designed and synthesized for soft materials including elastomers, hydrogels and organogels to achieve extreme properties. Based on their topologies, the typical UPN architectures can be classified into a number of categories, including ideal polymer networks, polymer networks with slidable crosslinks, interpenetrating polymer networks, semi-interpenetrating polymer networks, and bottlebrush polymer networks (**Figure 8** and **Table 1**).

4.1.1. Ideal polymer networks

Ideal polymer networks are polymer networks that have uniform chain length, uniform functionality, and no defect (**Figure 8**)³³³. Following the pioneer work by Sakai et al.^{333,447-449}, the ideal polymer networks have been commonly fabricated using multi-arm macromers, where the arms of adjacent macromers are crosslinked into polymer chains. Because the lengths of the

macromer arms are uniform and the reaction efficiency of the crosslinking process is high, various ideal polymer networks with uniform chain length, uniform functionality, and almost no defect have been achieved^{74,333,450-460}. The tetra-arm PEG^{333,461,462} is among the most frequently used macromers for the fabrication of hydrogels with ideal polymer networks. The ends of the PEG macromers are commonly modified with pairs of reaction groups such as *N*-hydroxysuccinimide and amine^{74,333,463}, tetrabenzaldehyde and tetrabenzaacylhydrazide⁴⁶⁴, maleimide and thiol⁴⁶⁵, or boronic acid and diol^{450,454,461}. Due to the almost defect-free nature, the ideal polymer networks have been made highly stretchable and resilient⁷⁴. It should be noted that, although the conventional polymer networks usually have non-uniform chain lengths and topological defects, their mechanical properties are commonly calculated based on the models of ideal polymer networks such as the affine network model and the phantom network model (**Section 3**). Therefore, the ideal polymer networks by themselves still have coupled mechanical properties.

4.1.2. Polymer networks with slidable crosslinks

A slidable crosslink, commonly in the form of two covalently-crosslinked polymer rings, can interconnect two polymer chains that thread through and slide inside the rings (**Figure 8**)⁴⁶⁶. Polymer networks with slidable crosslinks are both mechanically stable and reconfigurable due to the permanent and slidable nature of the crosslinks, respectively. Under mechanical loads, the slidable crosslinks tend to reconfigure the polymer network in a way that the polymer chains in the network sustain the same level of forces, so that the reconfigured polymer network approximates an ideal polymer network.

The polymer networks with slidable crosslinks are mainly synthesized from cyclodextrinbased polyrotaxanes⁴⁶⁷⁻⁴⁶⁹. Cyclodextrins are a series of cyclic oligosaccharides with 6, 7, or 8 glucose units (named α -, β -, or γ - cyclodextrin, respectively). Cyclodextrin-based polyrotaxanes are inclusion complexes composed of linear polymer chains that are threaded through the cyclodextrin molecules and then capped by bulky groups at the chain ends⁴⁶⁹⁻⁴⁷¹. The formation of cyclodextrin-based polyrotaxanes mainly depends on the size matching between the interior cavities of the cyclodextrins and the cross-section of the polymer chains⁴⁷². Many polymer chains have been investigated to form cyclodextrin-based polyrotaxanes including linear homopolymers, linear block copolymers as well as branched polymers⁴⁷². The α -cyclodextrin has the smallest cavity size and can form inclusion complexes with PEG or PCL, but not with PPO chains^{473,474}. The β -cyclodextrin can form complexes with PCL or PPO, but not with PEG^{473,475,476}. The γ cyclodextrin, which has the largest cavity size, can thread through a PPO chain or two chains of PEG or PCL⁴⁷⁷. The cyclodextrins can be crosslinked with each other to interconnect the threaded polymer chains and form the polymer networks with slidable crosslinks^{67,372}. Because a polymer network with slidable crosslinks under mechanical loads approximates an ideal polymer network, the mechanical properties of the polymer network with slidable crosslinks are usually coupled with one another as discussed in Section $3^{466,478-480}$.

4.1.3. Interpenetrating and semi-interpenetrating polymer networks

An interpenetrating polymer network is comprised of two or more interpenetrated polymer networks, which are individually crosslinked but not joint together (Figure 8); a semiinterpenetrating polymer network is comprised of two or more interpenetrated polymer networks, in which at least one network is uncrosslinked and others are individually crosslinked but not joint together (Figure 8) $^{372,398,481-488}$. The interpenetrating and semi-interpenetrating polymer networks are entangled or interlocked in a way that they cannot be pulled apart unless the networks are broken⁴⁸⁵⁻⁴⁸⁸. Hydrogels based on the interpenetrating and semi-interpenetrating polymer networks are commonly prepared following the sequential or simultaneous method. In the sequential method, one polymer network is first prepared and then immersed into a solution of monomers, initiators and/or crosslinkers for another polymer network. Thereafter, the interpenetrating or semiinterpenetrating polymer network is formed by polymerizing the second polymer network within the first network. As a remarkable example, Gong et al. have adopted the sequential method to fabricate the double-network hydrogels with high fracture toughness⁷⁰. In the simultaneous method, a mixture of the polymers, monomers, initiators and crosslinkers for all polymer networks form the interpenetrating or semi-interpenetrating polymer networks in one step or one pot⁴⁸⁹. This onestep or one-pot fabrication process is a desirable feature of the simultaneous method compared to the sequential method. One remarkable example of the simultaneous method is the simple fabrication of the polyacrylamide-alginate hydrogel with high stretchability and fracture toughness⁴⁹⁰. A wide range of material candidates including both natural and synthetic polymers as discussed in Section 2^{67,372,398,482-484,490-492} have been used to synthesize hydrogels with interpenetrating and semi-interpenetrating polymer networks via various crosslinking strategies^{254,398,484,493}. As will be discussed in Section 4.1.7 and Section 5, the interpenetrating and semi-interpenetrating polymer networks can provide decoupled and extreme mechanical properties for hydrogels, such as extremely high stretchability and fracture toughness^{70,490,494-496}.

4.1.4. Polymer networks with high-functionality crosslinks

The functionality of a crosslink refers to the number of polymer chains interconnected at the crosslink. Common covalent crosslinks as discussed in Section 2.3 usually have relatively low functionality (e.g., less than 10), and there is usually a single polymer chain bridging between two adjacent covalent crosslinks. To dramatically enhance the functionality of a polymer network, various types of high-functionality crosslinks can be introduced into the polymer networks, domains^{232,316,318,497}. nodules498,499, including crystalline glassy nano-/microparticles^{372,398,492,497,500-502}, and micro-phase separations⁵⁰³⁻⁵⁰⁵ (Figure 8). For example, poly(vinyl alcohol) can form nano-crystalline domains to crosslink the polymer networks through the freezethaw method^{318,506}; poly(methyl methacrylate) can form glassy spheres to crosslink poly(methyl methacrylate)-based block copolymers into networks⁵⁰⁷; exfoliated particles, such as nano-clays⁵⁰⁸, graphene oxide⁵⁰⁹ and stratified lamellar bilayers⁵¹⁰, can crosslink polyacrylamide into moldable

and self-healable hydrogels; mixtures of styrene, butyl acrylate and acrylic acid can form microspheres to crosslink the residual polymer chains into microsphere-composite hydrogels⁵¹¹.

Multiple polymer chains (e.g., over 10) can be interconnected at each high-functionality crosslink (**Figure 8**). In addition, there can be multiple polymer chains bridging between two neighboring high-functionality crosslinks, where the lengths of the polymer chains can be highly non-uniform (**Figure 8**)^{67,451}. As will be discussed in **Section 4.1.7** and **Section 5**, the polymer networks with high-functionality crosslinks can provide decoupled and extreme mechanical properties for hydrogels, such as extremely high fracture toughness, resilience, tensile strength, and fatigue resistance.

4.1.5. Nano-/micro-fibrous polymer networks

Both synthetic and natural polymers can assemble into fibers (or fibrils referring to short fibers) with diameters ranging from nanometers to micrometers via covalent or physical bonds. The nano-/micro-fibers can further entangle, aggregate, and crosslink into percolated polymer networks^{6,71,512-517} (**Figure 8**). In biological organisms, cells can secrete proteins (e.g., collagens) and polysaccharides (e.g., celluloses), which then assemble into nano-/micro-fibrous polymer networks^{120,267,518-521}. These naturally derived fibers and fibrous networks have been widely harnessed for the fabrication of hydrogels with nano-/micro-fibrous polymer networks^{397,522-525}. In addition, a wide range of natural and synthetic polymers have been fabricated into nano-/micro-fibrous polymer networks with the spinning techniques⁵²⁶⁻⁵²⁸, among which the electrospinning is most popular due to its simplicity, low cost, and wide applicability⁵²⁹. In particular, the diameter, alignment and density of the fibers can be readily controlled by tuning the parameters of the electrospinning process⁵³⁰⁻⁵³⁴. As will be discussed in **Section 4.1.7** and **Section 5**, the nano-/micro-fibrous polymer networks can provide decoupled and extreme mechanical properties for hydrogels, such as extremely high fracture toughness, tensile strength, resilience, and fatigue resistance^{535,536}.

4.1.6. Other unconventional polymer network architectures

Many other types of UPN architectures can provide extraordinary mechanical properties as well. For example, the bottlebrush polymer networks (**Figure 8**) have shown extremely low shear moduli and tissue-like stress-strain relations in the solvent-free state^{537,538}. Although these UPN architectures have not been widely used in hydrogels, they can be exploited for the design of hydrogels in future⁵³⁹. Furthermore, it is also expected that new UPN architectures will be invented together with the development of polymers and soft materials.



Figure 8. Schematics of unconventional polymer network architectures, including ideal polymer networks, polymer networks with slidable crosslinks, interpenetrating polymer networks, semi-interpenetrating polymer networks, polymer networks with high-functionality crosslinks, nano-/micro-fibrous polymer networks, and bottlebrush polymer networks.

UPN architectures	Examples	References
Ideal polymer	Covalently-crosslinked 4-arm end-	74,333,372,450-465
networks	functionalized PEG;	
	Reversibly-crosslinked 4-arm end-	
	functionalized PEG	
Polymer	PEG with polyrotaxanes as slidable	67,372,466,469-473,475-480
networks with	crosslinks;	
slidable	PCL with polyrotaxanes as slidable	
crosslinks	crosslinks;	
	PPO with polyrotaxanes as slidable	
	crosslinks	
Interpenetrating	Covalently-crosslinked PAMPS	67,70,372,398,482-485,491
polymer	interpenetrated with covalently-	
networks	crosslinked PAAm;	

Table 1. Examples of unconventional polymer network architectures.

	Covalently-crosslinked polysaccharide	
	arosslinked PHEMA DEG DAAm	
	DNIDAm and DDMA	
	FNIFAIII, and FDMA	
Semi-	Covalently-crosslinked PAAm	67,70,372,398,481,483,484,490,492
interpenetrating	interpenetrated with reversibly-	
polymer	crosslinked or uncrosslinked PVA,	
networks	alginate, chitosan, and hyaluronan;	
	Covalently-crosslinked gelatin	
	interpenetrated with reversibly-	
	crosslinked or uncrosslinked alginate;	
	Covalently-crosslinked PEG-DA,	
	gelatin interpenetrated with reversibly-	
	crosslinked or uncrosslinked alginate,	
	chitosan, and hyaluronan;	
	Covalently-crosslinked PAAm	
	interpenetrated with reversibly-	
	crosslinked or uncrosslinked PVA	
Polymer	PVA with crystalline domains as high-	6/,232,316,318,372,398,451,492,493,497,500,503-
networks with	functionality crosslinks;	511
high-	Poly(methyl methacrylate) with glassy	
functionality	spheres as high-functionality	
crosslinks	crosslinks;	
	Polyacrylamide with exfoliated	
	particles as high-functionality	
	crosslinks;	
	Mixtures of polystyrene, poly(butyl	
	acrylate), and poly(acrylic acid) with	
	microsphere composites as high-	
	functionality crosslinks	
		6 120 397 522-536
Nano-/micro-	Cellulose, cellulose-derivative,	
fibrous polymer	collagen, gelatin, fibrin and elastin	
networks	nano-/micro-fibrous networks	

Other UPN	Poly(dimethylsiloxane) bottlebrush;	537-539
architectures	Poly(n-butyl acrylate) bottlebrush;	
	Poly polylactic acid-b-polyethylene	
	glycol- <i>b</i> -poly(<i>N</i> -isopropylacrylamide)	
	triblock bottlebrush	

4.1.7. Decoupled mechanical properties due to unconventional polymer network architectures

Polymer chains in the ideal polymer networks have uniform chain lengths (i.e., the same N), and the polymer networks with slidable crosslinks also tend to give relatively uniform chain lengths under mechanical loads. These polymer networks with relatively uniform chain lengths are named unimodal polymer networks (**Figure 9a**)^{60,63,443}. Because the shear moduli, stretch limits and intrinsic fracture energy of conventional polymer networks have been derived based on the unimodal polymer networks, these mechanical properties are still coupled in hydrogels with the ideal polymer networks and the polymer networks with slidable crosslinks (**Section 3**).

The interpenetrating polymer networks, semi-interpenetrating polymer networks and polymer networks with high-functionality crosslinks can integrate polymer chains with varying chain lengths (i.e., different N) into the same polymer networks, which are often named multimodal polymer networks (**Figure 9b**)^{60,63,443}. Let's classify the polymer chains in a multimodal polymer network into different types based on their chain lengths (**Figure 9b**). For the *i*th type of polymer chains, the number of the polymer chains per unit volume of the polymer network in the dry state, the number of Kuhn monomers per polymer chain, and the volume of the Kuhn monomer are denoted as n_i , N_i and v_i , respectively.

The multimodal polymer network (**Figure 9b**) can be designed such that the corresponding hydrogel can sustain its integrity up to the stretch limit of the longest polymer chains λ_{lim} , which can be expressed as^{70,490},

$$\lambda_{\rm lim} = \sqrt{N_{\rm max}} \lambda_s^{-1} \tag{15}$$

where N_{max} is the number of Kuhn monomers on the longest polymer chain, and λ_s^{-1} accounts for the effect of swelling on the chain stretch limit λ_{lim} . The stretch limit of the bulk hydrogel scales with λ_{lim} , and the pre-factor of the scaling relation depends on the polymer network architecture⁶¹.

Based on the affine network model, the shear modulus of the hydrogel with the multimodal polymer network (Figure 9b) can be expressed as

$$G = \sum n_i k T \lambda_s^{-1} \tag{16}$$

where n_i and λ_s^{-1} account for the effects of the *i*th type of polymer chains and swelling on the initial shear modulus of the hydrogel.

Following the Lake-Thomas model, the intrinsic fracture energy of the hydrogel with the multimodal polymer network (Figure 9b) can be calculated as,

$$\Gamma_0 = \sum n_i b_i N_i^{3/2} U_i \lambda_s^{-2} \tag{17}$$

where b_i and U_i are the length and the fracture energy of a Kuhn monomer on the *i*th type of polymer chain, respectively, and λ_s^{-2} accounts for the effect of swelling on the intrinsic fracture energy. It should be noted that the fracture toughness and interfacial fracture toughness of the multimodal polymer network (**Figure 9b**) can be much higher than the intrinsic fracture energy, which will be discussed in Section 5.

It is commonly assumed that the polymer chains occupy the major volume of the polymer network in the dry state, and therefore the volume conservation of the multimodal polymer network (**Figure 9b**) gives

$$\sum N_i n_i v_i = 1 \tag{18}$$

Despite the relation of **Eq. 18**, the stretch limit, shear modulus and intrinsic fracture energy of a hydrogel with the multimodal polymer network (**Figure 9b**) can still be decoupled and independently designed. Without loss of generality, let's consider a hydrogel with bimodal distribution of chain lengths as an example (**Figure 9b**). A high density of the short polymer chains can give a high initial shear modulus of the hydrogel. While these short chains will be fractured when the hydrogel is highly stretched, the long polymer chains can still maintain the integrity and high stretch limit of the hydrogel^{70,490}. Similarly, the long polymer chains can give a relatively high intrinsic fracture energy of the hydrogel⁵⁴⁰.

The mechanical properties of the nano-/micro-fibrous polymer networks are determined by their fibers, interactions of the fibers (e.g., crosslinks between fibers), and topologies of the fibrous polymer networks. Therefore, the stretch limit, shear modulus and intrinsic fracture energy of nano-/micro-fibrous hydrogels do not follow the coupling relations for the conventional polymer networks (**Eqs. 13** and **14**), and therefore they can be independently designed.



Figure 9. Decoupled mechanical properties of hydrogels due to UPN architectures: a. unimodal polymer networks such as ideal polymer networks and polymer networks with slidable crosslinks give coupled mechanical properties; b. multimodal polymer networks such as interpenetrating polymer networks, semi-interpenetrating polymer networks and polym

4.2. Unconventional polymer network interactions

As illustrated in **Figure 10**, the UPN interactions are defined as inter-polymer and intra-polymer interactions that are different from those in the conventional polymer networks (i.e., permanent covalent crosslinks, excluded volumes, and osmotic interactions) (**Figure 10**). The UPN interactions are vastly abundant in biological organisms⁵⁴¹, and the UPN interactions have been intensively studied for the design of soft materials such as elastomers, hydrogels and organogels to achieve extreme mechanical properties among many other purposes²⁵⁴. Based on the nature of the UPN interactions, they can be broadly classified into three categories⁶²: strong physical crosslinks, weak physical crosslinks, and dynamic covalent crosslinks (**Figure 10** and **Table 2**).

4.2.1. Strong physical crosslinks

In addition to the permanent covalent crosslinks discussed in **Section 2.3**, various types of strong physical bonds can act as effectively permanent crosslinks in polymer networks. Typical examples
of strong physical crosslinks include crystalline domains, glassy nodules, and helical structures. The energy of strong physical crosslinks is similar to those of permanent covalent crosslinks.

Crystalline domain. A specific subset of synthetic and natural polymers can form crystalline domains under appropriate conditions. A crystalline domain, with the size from nanometers to micrometers, can serve as a strong physical crosslink for multiple amorphous polymer chains connected to the crystalline domain (Figure 10). As an example in synthetic polymers, PVA can form crystalline domains by repeated freeze-thaw cycles or by annealing at temperatures above its glass transition temperature^{316,318,497}. The formation of PVA crystalline domains is mainly due to the hydrogen-bonding interactions of the hydroxyl groups on PVA chains⁴. As an example in natural polymers, chitin and chitosan can form semi-crystalline polymer networks with crystalline domains crosslinking amorphous chains by treating the chitin and chitosan with strongly acidic or basic solutions to overcome the inter-chain electrostatic repulsions^{232,235}. As another example in natural polymers, cellulose can also form highly crystalized nanofibers due to the strong interaction between glucose units⁵⁴². These cellulose nanofibers can further aggregate and form a stable network by alkaline treatments^{512,543}. It should be noted that heating the abovementioned semi-crystalline polymer networks above their melting temperatures can destroy the crystalline domains in the networks, although most crystalline domains are stable at the room and body temperatures.

Because a crystalline domain usually interconnects multiple polymer chains, they often act as high-functionality crosslinks in the polymer networks as discussed in **Section 4.1.4**. In addition, the energy required to pull a polymer chain out of a crystalline domain is much higher than that required to fracture the same polymer chain⁷⁸; therefore, crystalline domains can also introduce intrinsically high-energy phases into the polymer networks. These attributes of crystalline domains have endowed the hydrogels containing crystalline domains with extreme mechanical properties, such as tough, strong, resilient and fatigue-resistant, which will be discussed in **Section 5**.

Glassy nodule. Glassy nodules are formed by the reversible liquid-glass transition of amorphous polymers when the temperature is decreased below their glass-transition temperatures⁵⁴⁴. In order to harness glassy nodules as strong physical crosslinks, block copolymers that contain at least one segment with a high glass-transition temperature have been commonly used. As the temperature reduces to room or body temperature, the segments with the high glass-transition temperature form glassy nodules that effectively crosslink the adjacent amorphous polymer chains (**Figure 10**)⁵⁴⁵. For example, the polystyrene segments in the polystyrene-*b*-poly(*N*-isopropylacrylamide)-*b*-polystyrene copolymers can form glassy nodules at room temperature to crosslink the block copolymer chains into a polymer network⁵⁴⁶. As another example, poly(methyl methacrylate) has a glass transition temperature around 115 °C⁵⁴⁷; therefore, the poly(methyl methacrylate) segments in the poly(methyl methacrylate) segments in the poly(methyl methacrylate)-*b*-poly(n-butyl acrylate) copolymers can form glassy spheres at room temperature to crosslink the poly(methyl methacrylate) segments in the poly(methyl methacrylate)-*b*-poly(n-butyl acrylate) copolymers can form glassy spheres at room temperature to crosslink the poly(methyl methacrylate) segments in the poly(methyl methacrylate)-*b*-poly(n-butyl acrylate) copolymers can form glassy spheres at room temperature to crosslink the polymer network⁵⁰⁷.

intrinsically high-energy phases in polymer networks to give the corresponding hydrogels extreme mechanical properties, which will be discussed in **Section 5**.

Helical association. Many natural polymers, due to their precisely-controlled structures, can assemble into nanometer-scale helical fibers (or fibrils), which then can aggregate or entangle to form a crosslinked network (**Figure 10**)^{120,522,524,525}. For example, the well-known triple-helix structure of type I collagen is formed by the self-assembly of three peptide strands. These collagen triple helices can pack together to form collagen nanofibers which further self-assemble into an interconnected hydrogel network^{548,549}. As another example, the linear agarose chains are disordered coils in aqueous solutions at high temperatures, and can form double-helix strings⁵⁵⁰ or simple helical chains⁵⁵¹ when the temperature is decreased to the room or body temperature. These stings or chains can associate to form agarose fibers through hydrogen bonding and further be entangled to form the interconnected hydrogel network⁵⁵².

4.2.2. Weak physical crosslinks

Compared to the strong physical crosslinks, many other physical crosslinks in polymer networks are relatively weak, transient, and reversible. Typical examples of weak physical crosslinks include hydrogen bond, electrostatic interaction, metal coordination, guest-host interaction, hydrophobic association, and π - π stacking (**Figure 10**). The energy of weak physical crosslinks is usually lower than those of strong physical crosslinks and permeant covalent crosslinks.

Hydrogen bond. The energy of a single hydrogen bond ranges from 0.8 kJ mol⁻¹ to 167 kJ mol⁻¹ (**Figure 6**) ^{388,389}. Many natural polymers can form hydrogels by the intermolecular hydrogen bonds. For example, gelatin can form polymer networks of helical structures crosslinked by hydrogen bonds¹⁸⁷; certain types of polysaccharides, such as agarose, amylose, amylopectin and carrageenan, can also form helical structures in solutions and be crosslinked into hydrogels by hydrogen bonds⁵⁵³. A number of synthetic polymers are also capable of forming physical hydrogels via hydrogen bonds. For example, PVA hydrogels can be obtained by forming hydrogen bonds between polymer chains through repeated freezing and thawing of PVA solutions³¹⁹. Polymethacrylic acid (PMA) or PAA can form complexes with PEG by hydrogen bonds between the oxygen groups of PEG and the carboxyl groups of PMA⁵⁵⁴ or PAA⁵⁵⁵.

Despite the abundance of hydrogen-bond groups (-OH, -NH, -C=O, -C-O) in natural and synthetic polymers, the hydrogen-bond interactions in hydrogels are usually screened due to the water molecules in hydrogels. To enable effective hydrogen-bond crosslinks, hydrophobic moieties with multiple self-complementary hydrogen-bond groups have been introduced onto the polymers^{493,556,557}. For instance, functionalizing PEG, PHMEA and PNIPAM with amine triazine or diamino triazine groups enables the formation of triple hydrogen bonds per crosslink^{493,505,556,558}. Similarly, the introduction of ureidopyrinimidone (UPy) groups onto PEG, PHMEA, PNIPAM, PAA and PDMAA chains gives quadruple hydrogen bonds per crosslink^{493,556,557}. Complementary DNA base pairs (A-T, C-G) can also serve as hydrogen-bond motifs when the DNA pairs are attached to polymer chains⁵⁵⁹.

Electrostatic interaction. The energy of electrostatic interactions ranges from 5 kJ mol⁻¹ to 200 kJ mol⁻¹ (Figure 6)³⁹⁰. Natural and synthetic polymers with fixed charges, named polyelectrolytes, can be physically crosslinked by electrostatic interactions^{122,490,560,561}. As a typical example of the anionic polyelectrolytes, alginate has been physically crosslinked with a wide range of divalent cations such as Ca^{2+} , Ba^{2+} and Mg^{2+} . Although the energy of a single ionic bond in alginate is relatively low, multiple (e.g., over 20) adjacent ionic crosslinks on the alginate chains can form a densely-crosslinked region following the "eggbox" model^{6,120,122,562}, giving relatively stable alginate hydrogels. As a typical example of the cationic polyelectrolytes, chitosan has been crosslinked by multivalent anions such as citrate and tripolyphosphate⁵⁶³⁻⁵⁶⁵. Electrostatic interactions of oppositely charged polyelectrolytes can also give physically-crosslinked hydrogels. For example, anionic poly(L-glutamic acid) and cationic poly(L-lysine) can form an injectable hydrogel by simply mixing them in phosphate buffered saline solutions⁵⁶⁰. As another example, poly(3-(methacryloylamino) propyl-trimethylammonium chloride) and poly(sodium pstyrenesulfonate) can form polyion complexes and give a series of tough and self-healing hydrogels by the stepwise polymerization of the oppositely charged monomers⁵⁶⁶. It should be noted that the formation of ionic crosslinks usually requires low ionic strength of the solvents for the hydrogels to avoid charge shielding.

Coordination complex. A coordination complex consists of a central metal ion, especially transition metal ion, and a surrounding array of organic ligands^{567,568}. The energy of coordination complexes ranges from 100 kJ mol⁻¹ to 300 kJ mol⁻¹ (Figure 6) ³⁹⁰. Coordination bonds provide structural support in many living tissues, such as human bone⁵⁶⁹, insect mandible⁵⁷⁰ as well as mussel byssal thread⁵⁷¹. Hydrogels crosslinked by coordination complexes are primarily achieved by functionalizing polymer backbones with chelating ligands, which then form coordination complexes with metal ions. Bisphosphonate⁵⁷²⁻⁵⁷⁴, catechol⁵⁷⁵⁻⁵⁷⁷, histidine⁵⁷⁸⁻⁵⁸⁰, thiolate^{581,582}, carboxylate^{583,584}, pyridine⁵⁸⁵, bipyridine⁵⁸⁶, and iminodiacetate^{580,587} have been widely used as the chelating ligands; Cu²⁺, Zn²⁺, Fe³⁺, Co²⁺ and Ni²⁺ are the commonly-used metal ions. Bisphosphonate ligands can be modified onto hyaluronan⁵⁸⁸, gelatin⁵⁸⁹ and PEG⁵⁷³ to form coordination complex with Ca²⁺, Mg²⁺ and Ag⁺. Besides bisphosphonate, catechol ligands are also widely used to functionalize various polymers such as PEG^{590,591}, gelatin⁵⁹², hyaluronic acid⁵⁹³, chitosan⁵⁹⁴, polyacrylamide⁵⁷⁶, and PAA⁵⁷⁷. As a typical example, PEG-modified with 3,4dihydroxyphenyl-L-alanine (DOPA) residues can form coordination complexes with metal ions (Cu²⁺, Zn²⁺ and Fe³⁺ ions) when the pH is above 8^{493,591}. In natural proteins, the histidine amino acid can give an imidazole ligand residue⁵⁹, which is one of the most important chelators in the human body⁵⁹⁵. PEG-modified with histidine can form coordination complexes with metal ions (Cu²⁺, Co²⁺ and Ni²⁺ ions) to achieve physical crosslinking of the PEG hydrogels^{493,596}. The mechanical properties of the hydrogels crosslinked by coordination complexes can be tuned by varying the metal ions and/or the chelating ligands^{578,597}.

Host-guest interactions. Host-gust interactions refer to two or more molecules or ions that are held together in unique structural relationships by forces other than those of covalent

bonds^{476,598,599}. The two most common host moieties are cyclodextrins and curcubit[n]urils. Cyclodextrins (CDs) are cyclic oligosaccharides which compose of 6 to 8 D-glucose repeating units linked by α -(1→4)-linked glycosidic bonds^{600,601}. Commonly-used CDs include α -, β -, and γ -CDs which are composed of 6, 7, and 8 D-glucose repeating units, respectively. These CDs have a truncated cone shape with the secondary and primary hydroxyl groups on the smaller cone rim exposed to the solvent⁵⁹⁹, which makes the CDs show a relatively hydrophobic inner cavity and a relatively hydrophilic outer surface. Therefore, these CDs can act as the host molecules for various hydrophobic guest molecules with appropriate molecular sizes through hydrophobic and van der Waals interactions^{398,472,476}. For example, common guests for α -CD include azobenzene⁶⁰² and ferrocene⁶⁰³; common guests for β -CD include adamantane⁶⁰⁴, benzimidazole⁶⁰⁵, 3-(trimethylsilyl)propionic acid⁶⁰⁶, azobenzene⁶⁰², ferrocene⁶⁰³. Among various guest molecules, adamantane has one of the greatest affinities due to its complementary size for β -CD and its high hydrophobicity⁶¹¹. In addition, the complexation of azobenzene or ferrocene to CDs is responsive to light⁶¹² or redox conditions^{613,614}, respectively.

The cucurbit[n]urils (CB[n, n=5-8]) are pumpkin-shaped macrocyclic oligomers made from the condensation reactions of formaldehyde and glycoluril^{599,615}. The CB[n]s usually have a structure of a rigid hydrophobic cavity with two identical hydrophilic polar carbonyl groups surrounding the portals. The cavity size ranges from 4.4 to 8.8 Å (for CB[n], n = 5–8) and the portal diameter ranges from 2.4 to 6.9 Å ⁶¹⁶. The binding affinities of CBs are often greater than that of other cavitands⁶¹⁷, mainly due to the formation of the strong charge–dipole, hydrogen bonding, and hydrophobic/hydrophilic interactions by the rigid inner cavities and the negative portals of CB[n]s⁶¹⁸⁻⁶²⁰. CB[8] also displays remarkable binding affinities towards positively charged and relatively large guests such as amantadine derivatives. Furthermore, the cavity of CB[8] is large enough to accommodate two organic guests simultaneously, thus forming highly stable ternary complexes. For example, CB[8] can form stable complexes with two 2,6-bis(4,5dihydro-1H-imidazol-2-yl)naphthalene molecules⁶²¹, or one viologen (paraquat) and one 2,6dihydroxynaphthalene together⁶²².

Hydrogels crossslinked by the host-guest interactions are usually constructed with polymer networks modified by the guest molecules and/or host molecules. For example, monomers, host molecules, and guest molecules can be copolymerized into polymer networks crossslinked by the host-guest interactions⁶²³. The host/gust molecules can also be attached to the backbones or ends of polymers such as PEG, PDMAA, hyaluronic acid and PAA, and then the addition of the corresponding difunctional gust/host crosslinkers will crosslink the polymer network⁵⁹⁹. Alternatively, guest-functionalized and host-functionalized polymers can also be synthesized separately, and a mixture of the two types of polymers gives hydrogels crosslinked by the host-guest interactions^{624,625}. Supramolecular hydrogels with these host-guest interactions have been extensively utilized to fabricate responsive materials⁶²⁶ and other dynamically assembling systems^{472,627}.

Hydrophobic association. The physical crosslink of hydrophobic association relies on the microphase separation and aggregation of hydrophobic domains of the polymer chains^{493,505}. The energy of hydrophobic association ranges from 0.1 kJ mol⁻¹ to 20 kJ mol⁻¹ (**Figure 6**) ³⁹⁰. The hydrophobic domains can be introduced by post-polymerization modification (e.g., via the grafting-to approach) or by copolymerizing hydrophobic monomers within the polymer chains, either randomly or as blocks⁶²⁸. These modifications usually require the usage of non-aqueous solvents, mixed solvents, or micellar systems^{503,629}. As a typical example of introducing the hydrophobic domains is the synthesis of multiblock copolymers with hydrophobic n-alkyl acrylate end blocks and a large middle block of PEG, PAAm, PAA or PHEMA polymers⁶³⁰⁻⁶³². Notably, because one hydrophobic association can interconnect multiple polymer chains, the hydrophobic association has also been used as high-functionality crosslinks in hydrogels^{307,633}, although the energy of hydrophobic association is usually lower than those of crystalline domains and glassy nodules.

π-π stacking. The π-π stacking interaction is a type of noncovalent interaction which refers specifically to the attractive interactions between π electrons in the aromatic groups⁶³⁴. The π-πinteractions can be divided into the edge-to-face (T-shaped), offset and face-to-surface stacking structures based on the geometry of the aromatic interactions⁶³⁵. The energy of π-π stacking ranges from 1 kJ mol⁻¹ to 50 kJ mol⁻¹ (**Figure 6**)³⁹¹. Natural amino acids with aromatic rings, such as phenylalanine, tyrosine and tryptophan, and other compounds with conjugated structures such as fluorenylmethyloxycarbonyl (Fmoc), 1-pyrenebutyric acid, 2-naphthalene acetic acid and nitrophenyl methacrylate can be used to design and prepare polymers with aromatic moieties for gelation by the π-π stacking interactions^{514,636,637}. For example, aromatic moiety containing short peptides and *N*-terminal Fmoc-amino acids can self-assemble into robust supramolecular architectures⁵¹⁴. In addition, carbon nanotubes^{638,639}, polythiophene^{32,116}, and graphene-based nanomaterials^{640,641} (including single-layer graphene, multilayer graphene, graphene oxide, and reduced graphene oxide) are also capable to form π-π interactions, which are useful for preparing electrically conductive hydrogels^{642,643}.

4.2.3. Dynamic covalent crosslinks

In addition to the weak physical bonds, dynamic covalent bonds can also act as reversible crosslinks that are cleavable by external stimuli. The energy of dynamic covalent bonds is usually similar to or lower than those of permanent covalent bonds⁶⁴⁴, and higher than those of weak physical bonds (**Figure 6**). Typical examples of dynamic covalent crosslinks in hydrogels include imine bond, boronate ester bond, disulfide bond, cyclohexenes hydrazone bond, oxime bond, and reversible Diels-Alder reaction (**Figure 10**).

Imine bond. An imine is a carbon-nitrogen double bond commonly formed by reactions between amine and aldehyde or ketone⁶⁴⁵. In particular, the imine crosslinks in hydrogels are usually formed through the Schiff base reactions, which give aliphatic Schiff bases or aromatic

Schiff bases⁶⁴⁶⁻⁶⁴⁹. The reversible nature of the imine crosslinks endows the resultant hydrogels with properties such as mechanical dissipation, self-healing and stimuli responses⁶⁵⁰. The energy of the Schiff bases ranges from 67 kJ mol⁻¹ to 477 kJ mol⁻¹ (**Figure 6**)^{387,392}. The aromatic Schiff bases usually have higher energy and stability than the aliphatic Schiff bases^{651,652}.

The imide bonds are particularly useful for preparing biopolymer-based hydrogels, because most biopolymers such as proteins contain amine groups. These amines can form imide bonds with various aldehydic crosslinkers at mild conditions^{650,653}. The obtained hydrogels with imine bonds are usually sensitive to various chemical and biological stimuli, including pH, free amine, and free aldehydes⁶⁵². These hydrogels can be used as self-healing materials and injectable scaffolds in biomedical applications^{440,654,655}.

Boronate ester bond. The dynamic boronate ester bonds are formed by the reaction of diols and boronic $\operatorname{acid}^{656-658}$. The energy of boronate ester bonds ranges from 27.2 kJ mol⁻¹ to 93.3 kJ mol⁻¹ (**Figure 6**) ³⁹³, highly dependent on pH and temperature⁶⁵⁹⁻⁶⁶¹. The boronic acid can be introduced into hydrogels by polymerizing boronic acid-containing monomers together with other monomers, such as acrylamide (AAm)⁶⁶² and *N*-isopropylacrylamide (NIPAM)⁶⁶³. Alternatively, boronic acid functional groups can also be grafted onto pre-formed polymer chains through the carbodiimide chemistry^{664,665}.

The boronic acid-containing polymers can react with polymers containing diol functional groups. For example, polymers modified with boronic acid can form the dynamic boronate ester crosslinks with salicyl hydroxamic acid groups in acidic environment or catechol groups in alkaline environment^{590,660,666-668}. As another example, polyhydroxy polymers such as PVA^{669,670}, alginate^{671,672}, and cellulose⁶⁷³ can also be crosslinked into dynamic hydrogels by mixing the polyhydroxy polymers with boronic acid-containing polymers in aqueous solutions. The transient boronate ester networks usually can dynamically restructure after fracture, making the resultant hydrogels injectable and self-healable^{674,675}. In addition, the boronate ester crosslinked hydrogels are also glucose-sensitive, because glucose can compete with diol groups to form boronate–glucose complexes and therefore de-crosslink the hydrogels⁶⁶⁹. These glucose-sensitivity hydrogels, based on the boronate ester bonds, have been used for self-regulated insulin release and glucose sensing^{398,664,669,676}.

Disulfide bond. Disulfide bonds are dynamic covalent bonds based on thiol-thiol interactions at slightly alkaline environments or at mild oxidative conditions^{677,678}. The energy of disulfide bonds is around 425 kJ mol⁻¹ (**Figure 6**)^{387,394}. Many natural polymers have disulfide bonds to stabilize their structures such as fibrinogen⁶⁷⁹ and collagen⁶⁸⁰. The disulfide bond can also be introduced into polymers by using disulfide bond-containing crosslinkers such as 3,3'-dithiobis(propanoic dihydrazide)^{681,682} and *N*,*N'*-cystamine-bis-acrylamide⁶⁸³⁻⁶⁸⁵. The thiol-thiol reaction has relatively fast kinetics and can be used to prepare dynamic hydrogels^{686,687}. Hydrogels crosslinked by disulfide bonds can be used to encapsulate various types of cells, due to the mild reaction conditions^{688,689}. In addition, the disulfide bonds can be cleaved by reducing agents such as tris(2-carboxyethyl)phosphine⁶⁹⁰, 1,4-dithiothreitol⁶⁹¹, and glutathione^{686,692}.

Hydrazone bond. Hydrazone bonds are formed by the reaction of aldehyde and hydrazide groups⁶⁹³. Polymers with hydroxyl groups, such as PEG⁶⁹⁴, cellulose⁶⁹⁵ and polysaccharide⁶⁴⁷, can be easily modified with aldehydes and hydrazide (or acylhydrazine) motifs. The reversible hydrazone bonds can be formed by simply mixing the aldehyde- and hydrazide-containing polymers under physiological conditions⁶⁹⁶⁻⁶⁹⁸.

Hydrogels crosslinked by the hydrazone bonds can exhibit reversible sol-gel transition properties by changing the pH^{694,699-701}. Hydrogels crosslinked by hydrazone bonds can be used for in situ cell encapsulation due to the cytocompatibility and fast gelation kinetics of the aldehyde and hydrazide coupling^{702,703}. The mechanical properties of these hydrogels can be easily tuned, which facilitates the study of the relationships between cell behaviors and mechanics (such as stress-relaxation kinetics) of the hydrogels⁷⁰⁴. Hydrazone bonds can also be used to prepare self-healing and injectable hydrogels based on the reversibility of hydrazone bonds at the mildly acid environment (pH 4.0–6.0)^{696,698,700}.

Oxime bond. Oxime bonds are formed by the reaction between hydroxylamine and aldehyde or ketone with high efficiency under mild conditions⁷⁰⁵. The reactive aldehyde or ketone groups can be modified onto polymers through radical polymerization⁷⁰⁶ or oxidation^{707,708}, while the hydroxylamine motifs are mainly modified onto hydroxyl-containing polymers through a sequential *N*-hydroxyphthalimide induced Mitsunobu reaction and hydrazine reduction⁷⁰⁹. Then, the oxime bonds can be formed by mixing the aldehyde- or ketone-containing polymers with the hydroxylamine-containing polymers in a neutral or slightly acid aqueous solution⁷¹⁰. This reaction is biocompatible without cytotoxic side products and can be used to crosslink biopolymers into hydrogels^{709,711}. Due to the dynamic nature, oxime bonds have been used for building self-healing and injectable hydrogels which show higher hydrolytic stability than the hydrogels crosslinked by imines and hydrazones^{710,712}.

Reversible Diels–Alder reaction. Diels-Alder reaction is a click reaction between diene and dienophile groups^{713,714}. The energy of Diels-Alder bonds ranges from 37.6 kJ mol⁻¹ to 130 kJ mol⁻¹ (**Figure 6**)^{395,396}. To harness the dynamic Diels-Alder reaction as reversible crosslinks for hydrogels, natural polymers (such as hyaluronic acid⁷¹⁵, cellulose⁷¹⁶ and other polysaccharides⁷¹⁷) and synthetic polymers (such as PNIPAM⁷¹⁸ and PEG⁷¹⁹) can be modified with diene (such as furan) functional groups and dienophile (such as maleimide) functional groups on their backbones or chain ends. The equilibrium of the Diels-Alder linkage is thermally responsive. For example, the adducted Diels-Alder linkage can reform maleimide and furan moieties when increasing the temperature^{720,721}. The Diels-Alder reaction can be performed in aqueous media at physiologically compatible conditions without any side reactions or byproducts⁷²²⁻⁷²⁵. Therefore, the Diels-Alder reaction has been used for preparing self-healing or adaptable hydrogels for biological applications in drug delivery and tissue engineering^{715,725,726}.



Figure 10. Schematics of unconventional polymer network interactions including **a.** strong physical crosslinks, **b.** weak physical crosslinks, and **c.** dynamic covalent crosslinks.

UPN interactions		Examples	References
Strong physical crosslinks	Crystalline domain	PVA treated by freeze-thawing or annealing	4,316,318,497
		Chitin and chitosan treated by acidic or basic solutions	232,235
		Cellulose treated by alkalines	512,542,543
	Glassy nodule	Polystyrene- <i>b</i> -poly(<i>N</i> -isopropylacrylamide)- <i>b</i> -polystyrene	546
		Poly(methyl methacrylate)- <i>b</i> -poly(n-butyl acrylate)	507
	Helical association	Self-assemble of agarose or gelatin	6,120,550,551
		Self-assemble of collagen or fibrinogen	6,120,548,549

 Table 2. Examples of unconventional polymer network interactions.

		Self-assemble of elastin-like polypeptides	522,525
Weak physical crosslinks	Hydrogen bond	PAA or polymethacrylic acid (PMA) with PEG	493,554-556
		PEG, PHEMA, PNIPAM with self- complementary hydrogen-bond groups (triazine moieties or ureido pyrinimidone moieties)	493,505,556-558
		Alginate with Ca ²⁺ , Ba ²⁺ , Mg ²⁺ , Zn ²⁺	6,120,122,562
	Electrostatic interaction	Chitosan with tripolyphosphate, citrate ions	563-565
		Cationic polyelectrolytes with anionic polyelectrolytes	560,566
		Bisphosphonate-containing polymers with metal ions (Ca ²⁺ , Mg ²⁺ or Ag ⁺)	573,588,589
	Coordination complex	Catechol-containing polymers with metal ions $(Cu^{2+}, Zn^{2+} \text{ and } Fe^{3+})$	493,590-593,596 576,577,594
		Histidine-containing polymers with metal ions (Cu ²⁺ , Co ²⁺ and Ni ²⁺)	493,595,59659
	Host-guest	Polymers containing β-CD moieties with azobenzene group, adamantyl group, ferrocene group, t-butyl group, cyclohexyl(ester) group, cyclododecyl(amide) group, benzyl group, 2-naphthylmethyl group, 1-pyrenylmethyl group	398,472,493,556,602,604- 610
	interaction	Polymers containing α -CD moieties with n- butyl group, adamantyl group, benzyl group, trans-azobenzene group	398,493,556,602,603,
		Polymers containing curcubit[n]uril moieties with spermine, diaminohexane, viologens, naphtalenes	398,556,621,622
	Hydrophobic association	PEG, PAAm, PNIPAM, PDMAA, PVA containing hydrophobic moieties (octylphenol-PEG acrylate, stearyl acrylate, lauryl acrylate)	493,503,505 503,628,629
		Triblock amphiphilic copolymers with PEG, PAAm, PVA, PHEMA middle blocks and n- alkyl acrylate end blocks	630-632

	π - π stacking	Polymers modified with aromatic moieties or	514,636,637
		conjugated structures	
		Hydrogels containing carbon nanotubes,	
		polythiophene, and graphene-based	32,116,638-641
		nanomaterials	
Dynamic covalent crosslinks	Imine bond	Polymers containing amine and aldehyde (or	398,440,646-650,652-655
		ketone) functional groups	
	Boronate	Polymers containing boronic acid and diol	398,658,664,669,670,676
	ester bond	functional groups	
	Disulfide	Polymers containing disulfide functional	398,631,682,688,689,693
	bond	groups	
	Hydrazone	Polymers containing hydrazide and aldehyde	693,694,696-700,702-704
	bond	(or ketone) functional groups	
	Oxime bond	Polymers containing hydroxylamine and	709-712,720
		aldehyde (or ketone) functional groups	
	Reversible	Polymers containing diene and dienophile functional groups	715,725,726
	Diels–Alder		
	reaction		

4.2.4. Decoupled mechanical properties due to unconventional polymer network interactions. The crystalline domains and glassy nodules have been widely used as the high-functionality crosslinks in UPNs, whose stretch limit, shear moduli and intrinsic fracture energy are decoupled as discussed in **Section 4.1.7** (Figure 9b).

The weak physical crosslinks and dynamic covalent crosslinks can act as reversible crosslinks in polymer networks, leading to decoupled mechanical properties of the resultant hydrogels (Figure 11). Without loss of generality, let's consider a conventional polymer network with long polymer chains (i.e., polymer network sparsely crosslinked by permanent covalent bonds that gives a high N value), whose stretch limit, shear modulus and intrinsic fracture energy are given by Eqs. 7-9, respectively. We next introduce reversible crosslinks such as weak physical crosslinks and dynamic covalent crosslinks into the polymer network. When the polymer network undergoes initial small deformation, the reversible crosslinks act as additional crosslinks⁴⁵⁰, increasing the effective chain density of the polymer network to n_{eff} . Therefore, the shear modulus of the hydrogel with the reversible crosslinks under initial deformation increases to

$$G = n_{eff} k T \lambda_s^{-1} \tag{19}$$

As the hydrogel is highly stretched, the reversible crosslinks can be de-crosslinked. However, the covalently-crosslinked long polymer chains (i.e., polymers with high N) still endow the hydrogel with a high stretch limit and high intrinsic fracture energy according to **Eq. 13**. Notably, hydrogels with only reversible crosslinks will display permanent plastic deformation when the reversible crosslinks are de-crosslinked under mechanical loads, because there is no permanent covalent crosslink to maintain the shape or elasticity of the hydrogels. Therefore, hybrid reversible and permanent crosslinks, instead of solely reversible crosslinks, are critical to achieving decoupled and extreme mechanical properties of hydrogels.



Figure 11. Decoupled mechanical properties of hydrogels due to UPN interactions. The reversible crosslinks give an effectively high density of short chains for high modulus, and the sparse covalent crosslinks give long chains for high stretchability and intrinsic fracture energy.

4.3. Synergy of unconventional polymer network architectures and interactions

It is not uncommon for an unconventional polymer network to simultaneously possess both UPN architectures and UPN interactions. In some cases, the UPN architectures and interactions are interdependent. The formation of certain UPN architectures requires certain UPN interactions, or certain UPN interactions naturally lead to the self-assembly of polymers into certain UPN architectures. For example, the UPN interactions of strong physical crosslinks such as crystalline domains and glassy nodules mostly have high functionalities, giving rise to UPN architectures with high-functionality crosslinks. As another example, the self-assembly of polymer chains into nano-/micro-fibers usually requires the UPN interactions such as weak physical crosslinks.

In other cases, specific types of UPN architectures and UPN interactions can be separately designed and then integrated into the same UPN. For example, weak physical crosslinks and dynamic covalent crosslinks have been introduced into various UPN architectures in order to design tough hydrogels⁴⁹⁰, because the dissociation and reformation of these reversible crosslinks can dissipate mechanical energy to toughen the hydrogels.

5. Design of hydrogels with extreme mechanical properties

While numerous UPN architectures and UPN interactions have been developed over the last few decades, the design of hydrogels that possess extreme mechanical properties has largely followed an Edisonian approach – trial and error with specific polymers. The rational design of hydrogels using different material candidates and fabrication methods for various applications remains a central need in the field of soft materials. In this section, we will summarize a set of general design principles for hydrogels to achieve the corresponding extreme mechanical properties, including extremely high fracture toughness, tensile strength, resilience, interfacial toughness, fatigue-threshold, interfacial toughness, and interfacial fatigue threshold (**Table 3** and **Table 4**). Then, we will discuss the implementations strategies for these design principles using the UPN architectures and/or the UPN interactions.

5.1. Tough: build dissipation into stretchy polymer networks

Fracture toughness. Fracture toughness has been widely used to characterize a material's capability to resist fracture under mechanical loads. One common definition for the fracture toughness of a material is the energy required to propagate a crack in the material over a unit area measured in the undeformed state (**Figure 12a**) which can be quantitatively expressed as,

$$\Gamma = G_c = -\frac{dU}{dA} \tag{20}$$

where U is the total potential energy of the system, A is the crack area measured in the undeformed state, and G_c is the critical energy release rate that drives crack propagation. According to **Eq. 20**, the unit for the fracture toughness is joule per meter squared (i.e., J m⁻²).

The fracture toughness of soft materials such as elastomers and hydrogels has been measured with many experimental methods such as the pure-shear test and the single-notch test, which have been summarized in a few recent review papers^{67,727,728}. For example, in the pure-shear test, two identical pieces of a hydrogel are fabricated with the same thickness *T*, width *W* and height *H*, where W >> H >> T (Figure 12a). Both pieces of the samples are clamped along their long edges (i.e., along the width direction) with rigid plates. A notch with a length of ~0.5*W* is introduced into the first sample, which is then gradually pulled to a stretch of λ_c times of its undeformed height until a crack begins to propagate from the notch (Figure 12a). The second sample without notch is uniformly stretched above the critical stretch λ_c to measure the nominal stress *s* vs stretch λ relation (Figure 12a). Thereafter, the fracture toughness of the hydrogel can be calculated as $\Gamma = H \int_{1}^{\lambda_c} sd\lambda$, based on the measured λ_c and $s - \lambda$ relation in the pure-shear tests.

As discussed in **Section 3**, the fracture toughness of a conventional polymer network is its intrinsic fracture energy Γ_0 , which is the energy required to fracture a layer of polymer chains over a unit area (**Figure 12b**). Evaluated with typical parameters of conventional polymer networks, the fracture toughness of the corresponding hydrogels is commonly limited to a few tens of joule per meter squared. In addition, the fracture toughness of hydrogels with conventional polymer networks is also coupled with their stretch limits and shear moduli (**Eqs. 11**). For example, in order

to increase the fracture toughness of a conventional polymer network, the chain length (i.e., N) and thus the stretch limit of the polymer network need to be increased. Consequently, the chain density (i.e., n) and thus the shear modulus of the polymer network will be decreased.

Design principle for tough hydrogels. The design principle for tough hydrogels is the same as the principle for toughening various engineering materials (such as metals⁷²⁹, ceramics ⁷³⁰, composites⁷³¹ and polymers⁶⁴) and various biological tissues (such as tendons, cartilages, muscles and blood vessels⁷³²). That is to integrate both ductility and mechanical dissipation in the same material, so that a process zone with substantial mechanical dissipation develops around the crack tip prior to crack propagation (**Figure 12c-e**). The mechanical dissipation of a material manifests as the hysteresis loop on its stress-stretch curve under a loading-unloading cycle (**Figure 12c-e**). The ductility of hydrogels generally relies on the high stretchability (or the high stretch limit) of their polymer networks (**Figure 12c-e**). Overall, the design principle for tough hydrogels is *to build dissipation into stretchy polymer networks*^{67,733}. Quantitatively, the total fracture toughness of a hydrogel with the capability of mechanical dissipation can be expressed as ^{67,734}

$$\Gamma = \Gamma_0 + \Gamma_D \tag{21}$$

where Γ , Γ_0 , and Γ_D are the total fracture toughness, the intrinsic fracture energy, and the contribution of mechanical dissipation in the process zone to the total fracture toughness, respectively. While the intrinsic fracture energy for hydrogels is usually limited to a few tens of joule per meter squared, the contribution of the process-zone dissipation can be extremely high because both the dissipated energy per volume of the process zone and the size of the process zone can be large values (**Figure 12c-e**). Indeed, the fracture toughness of tough hydrogels has exceeded 10,000 Jm⁻², orders of magnitude higher than that of hydrogels with conventional polymer networks⁶⁷.



Figure 12. Design principle for tough hydrogels: build dissipation into stretchy polymer networks. a. definition of fracture toughness, and the pure-shear test to measure the fracture toughness. When a notched sample with height *H* at the undeformed state is stretched by a critical ratio of λ_c under the pure-shear deformation, the crack begins to propagate (top). The relation of the nominal stress *s* and the stretch λ is measured for an un-notched sample (otherwise the same as the notched sample) under the pure-shear deformation (bottom). The fracture toughness can be calculated as $\Gamma = H \int_{1}^{\lambda_c} s d\lambda$ based on the measured λ_c and $s - \lambda$ relation in the pure-shear tests. **b.** the intrinsic fracture energy Γ_0 from fracturing a layer of polymer chains. **c.** the mechanical dissipation in the process zone around the crack tip contributes to the total fracture toughness by Γ_D . The mechanical dissipation manifests as a hysteresis loop on the stress-stretch curve. The total

fracture toughness of the tough hydrogel is $\Gamma = \Gamma_0 + \Gamma_D$. **d.** microscope image of the process zone around the crack in a PAMPS-PAAm hydrogel⁷³⁵. **e.** nominal stress *s* vs stretch λ relations for a PAAm-alginate hydrogel under loading and unloading cycles⁴⁹⁰. **d** is reproduced with permission from Ref⁷³⁵. Copyright 2009 American Chemical Society. **e** is reproduced with permission from Ref⁴⁹⁰. Copyright 2012 Springer Nature.

Implementation strategies for tough hydrogels. The design principle for tough hydrogels requires: i). at least one polymer network in the hydrogel maintains a high stretch limit and therefore the polymer chains in that polymer network need to have a high N value according to Eq. 7; and ii). at least one component in the hydrogel dissipates substantial mechanical energy under the deformation typically experienced in the process zone. The design principle for tough hydrogels has been implemented using various types of UPN architectures and UPN interactions. We will discuss a few examples in the following paragraphs.

The interpenetrating polymer networks and semi-interpenetrating polymer networks have been widely used for the design of tough hydrogels since the pioneer work of double-network hydrogels by Gong et al. in 2003 (Figure 13a)⁷⁰. A typical double-network hydrogel interpenetrates a long-chain network (with high *N*) and a short-chain network (with low *N*)⁷⁰. As the double-network hydrogel deforms, the short-chain network fractures and dissipates substantial mechanical energy, while the long-chain network maintains the integrity of the hydrogel even under high stretches, implementing the design principle for tough hydrogels (Figure 12c)^{70,733,735}. Gong et al first demonstrated that the fracture toughness of double-network hydrogels can exceed 1,000 Jm^{2 70}. Other interpenetrating and semi-interpenetrating polymer networks such as the triple-network architecture have also been developed for tough hydrogels⁷³⁶ and elastomers⁷³⁷, implementing the design principle. Notably, since the fracture of the short-chain network is usually irreversible, these hydrogels' capability of mechanical dissipation may be substantially reduced after a few cycles of large deformation⁷³⁴.

The polymer networks with high-functionality crosslinks have given tough hydrogels based on various types of polymers and high-functionality crosslinks. There are multiple polymer chains (e.g., over 10) bridging between two adjacent high-functionality crosslinks, and the lengths of these polymer chains are usually non-uniform (**Figure 13b**). As the hydrogel deforms, the relatively short polymer chains fracture or detach from the high-functionality crosslinks, while the relatively long polymer chains maintain the integrity and high stretchability of the hydrogel, implementing the design principle for tough hydrogels. The bonds between the polymer chains and the high-functionality crosslinks can be permanent covalent crosslinks⁷³⁸, strong physical crosslinks^{741,742}, and dynamic covalent crosslinks^{743,744}, or a combination of them⁵⁰¹. Depending on the number and lengths of polymer chains between adjacent crosslinks and the types of bonds between polymer chains and crosslinks, the corresponding hydrogel can have different capabilities of mechanical dissipation and stretchability and therefore different fracture toughness.

The nano-/micro-fibrous hydrogels have also been used to implement the design principle for tough hydrogels. The nano-/micro-fibers can be made intrinsically stretchable (**Figure 13c**), and their re-orientation and re-alignment in hydrogels under deformation further enhance the stretchability of the hydrogels (**Figure 13c**)^{740,743,745-748}. The fracture of the nano-/micro-fibers and pull-out of the nano-/micro-fibers from the hydrogel matrices can dissipate substantial mechanical energy. A combination of the high stretchability and the mechanical dissipation enabled by the nano-/micro-fibrous polymer networks implements the design principle for tough hydrogels.

In addition to the abovementioned UNP architectures, the UPN interactions have also been widely used to implement the design principle for tough hydrogels⁷⁴⁹. The strong physical crosslinks such as crystalline domains and glassy nodules naturally act as high-functionality crosslinks for the corresponding UPN architectures (**Figure 13b**), which lead to tough hydrogels as discussed above.

The weak physical crosslinks^{566,584,623,750-757} and dynamic covalent crosslinks⁷⁴³ have been added into polymer networks with long polymer chains (i.e., sparsely-crosslinked polymer networks via permanent covalent bonds) to design tough hydrogels. The weak physical crosslinks and dynamic covalent crosslinks act as reversible crosslinks in these hydrogels (**Figure 13d**). As the hydrogel deforms, many of these reversible crosslinks dissociate or de-crosslink to dissipate substantial mechanical energy and the sparsely-crosslinked long-chain polymer network still sustains the high stretchability of the polymer network (**Figure 13d**). A synergy of the mechanical dissipation and the high stretchability enabled by the hybrid reversible and covalent crosslinks implements the design principle for tough hydrogels.

The weak physical crosslinks and dynamic covalent crosslinks have also been added into UPN architectures such as the interpenetrating polymer networks (**Figure 13a**)^{490,563,758-765}, polymer networks with high-functionality crosslinks (**Figure 13b**)^{739,741,742,766-769}, and nano-/micro-fibrous polymer networks (**Figure 13c**)^{745,746,767,770} to further toughen the resultant hydrogels, leveraging these reversible bonds' capability of dissipating additional mechanical energy. Furthermore, unlike irreversibly-fractured polymer chains, the dissociated weak physical crosslinks and dynamic covalent crosslinks may re-associate due to their reversible nature, potentially endowing the tough hydrogels with recoverable dissipation over cyclic loads⁴⁹⁰. For further detailed discussion on the design principle and implementation strategies for tough hydrogels, a recent review paper is recommended⁶⁷.



Figure 13. Implementation strategies for tough hydrogels with UPNs. Schematics of the implementation strategies with **a**. interpenetrating or semi-interpenetrating polymer networks, **b**. polymer networks with high-functionality crosslinks, **c**. nano-/micro-fibrous polymer networks, and **d**. polymer networks with reversible crosslinks. The bottom panel of **c** is a microscope image of a fibrous fibrin hydrogel⁷⁴⁸. The bottom panel of **c** is reproduced with permission from Ref⁷⁴⁸. Copyright 2009 American Association for the Advancement of Science.

5.2. Strong: synchronize stiffening and fracture of multiple polymer chains

Tensile strength. Multiple types of strengths such as tensile strength, compressive strength and shear strength have been used to characterize the strength of a material. We will focus on tensile strengths of hydrogels in this paper due to two reasons: 1). The tensile, compressive and shear deformations of a sample are related to one another. For example, the uniaxial compression of a sample is equivalent to the biaxial tension of the sample; the pure shear of a sample is equivalent to the sample being elongated in one direction and shortened perpendicularly. 2). The tensile strength is easier to measure than the shear strength, and the tensile strength is less affected by boundary conditions in the measurement (such as friction) than the compressive strength.

Since soft materials such as elastomers and hydrogels usually do not yield plastically, their tensile strengths are commonly defined as the stresses at which the ultimate tensile failure occurs in the uniaxial tensile test. In addition, since the hydrogel samples usually undergo large deformation before failure, the tensile strength can be defined based on either the nominal stress or the true stress (**Figure 14a**),

$$s_f = \frac{F_f}{A}, \ \sigma_f = \frac{F_f}{a}$$
(22)

where F_f is the tensile force at the failure of the sample, A and a are the cross-section areas of the sample in the reference (undeformed) and current (deformed) states, respectively, and s_f and σ_f are the nominal and true tensile strengths, respectively. The nominal tensile strengths of hydrogels with conventional polymer networks and even tough hydrogels are commonly lower than 1 MPa^{70,490}, much lower than the tensile strengths of engineering materials such as metals and biological tissues such as tendons.

Design principle for strong hydrogels. A generic principle for the design of strong hydrogels is to make a substantial number of polymer chains in the polymer network to stiffen and fracture simultaneously (**Figure 14b**). Following this principle, the nominal and true tensile strengths of the polymer network can be evaluated as,

$$s_f = M_f f_f, \ \sigma_f = m_f f_f \tag{23}$$

where f_f is the force required to fracture a single polymer chain, which is on the order of a few nanonewton³⁸⁵, and M_f and m_f are the numbers of simultaneously fractured polymer chains per unit area of the polymer network at the undeformed and deformed states, respectively. It has been evaluated that s_f and σ_f can reach up to 1 GPa and 10 GPa, respectively, in an ideal scenario where all polymer chains in the polymer network fracture simultaneously⁷⁷¹.

In realistic situations, almost all materials contain defects in the forms such as notches, micro-cracks, cavities, impurities, and missing polymer chains or crosslinks. The presence of defects usually significantly reduces the tensile strengths of the materials⁷⁷²⁻⁷⁷⁴. Without loss of generality, let's assume the largest defect in the tensile sample is a notch with length D in the

undeformed state perpendicular to the tensile direction (Figure 14c). The tensile strength of the sample generally increases with the decrease of the defect size D up to a critical value D_c , below which the tensile strength is defect-insensitive (Figure 14c). A scaling relation for the critical defect size D_c can be expressed as^{773,774}

$$D_c \sim \frac{\Gamma}{W_c} \tag{24}$$

where W_c is the work for tensile failure of a unit volume of the defect-insensitive sample, and Γ is the fracture toughness of the sample.

In order to achieve strong hydrogels, it is highly desirable for the hydrogel samples to have defect-insensitive tensile strengths⁷⁷¹. According to Eq. 24, a tougher material (i.e., higher Γ) can be insensitive to larger defects due to a larger critical defect size D_c . For example, the critical defect size is on the order of a few nanometers for glass and ceramics, a few micrometers for brittle hydrogels, and a few millimeters for tough elastomers and hydrogels^{773,774}. Furthermore, it is a common strategy to set the characteristic size of the sample (e.g., the diameter of the sample in Figure 14c) to be similar to or smaller than the critical defect size D_c , so that the tensile strength of the sample is guaranteed to be insensitive to any possible defects in the sample⁷⁷¹.



Figure 14. Design principle for strong hydrogels: synchronize stiffening and fracture of multiple polymer chains. a. definition and measurement of the tensile strength. A and a are the cross-section areas of the sample in the undeformed and deformed states, respectively, and F is the applied tensile force. b. the simultaneous stiffening and fracture of multiple polymer chains give a high tensile strength⁷⁷¹. F_f is the tensile force at the failure of the sample. c. the nominal tensile strength s_c increases with the decrease of the defect size D up to a critical value D_c , below which the tensile strength is defect-insensitive^{773,774}.

Implementation strategies for strong hydrogels. The UPNs with high-functionality crosslinks such as nano-crystalline domains have been widely used for the design of strong hydrogels⁷⁵. As the hydrogels undergo large deformation, relatively short polymer chains are

gradually pulled out of the nano-crystalline domains, so that the polymer chains bridging adjacent nano-crystalline domains tend to have similar lengths and therefore stiffen and fracture simultaneously – implementing the design principle for strong hydrogels (**Figure 15a**).

The nano-/micro-fibrous polymer networks are another type of UPN architecture that implements the design principle for strong hydrogels (**Figure 15b**). The diameters of the nano-/micro-fibers can be readily controlled below the critical defect size D_c . Bundles of polymer chains in the nano-/micro-fibers can be designed to stiffen and fracture simultaneously to endow the fibers with high tensile strengths up to the ideal strengths (**Figure 15b**)^{76,740,743,747,775}. Consequently, the resultant nano-/micro-fibrous hydrogels can reach extremely high tensile strengths (**Figures 15c** and **d**). Notably, biological hydrogels such as tendons, ligaments and muscles commonly adopt nano-fibers and micro-fibers, often in hierarchical architectures, to achieve high tensile strengths (**Figure 2**).

In addition to the aforementioned UPN architectures, the UPN interactions can facilitate the implementation of the design principles for strong hydrogels. The strong physical crosslinks such as crystalline domains allow the pull-out of polymer chains from them to achieve simultaneous stiffening and fracture of multiple polymer chains (**Figure 15a**)⁷⁵. The weak physical crosslinks such as the hydrogen bonds can facilitate the alignment and self-assembly of polymer chains into bundles (**Figure 15b**), which tend to stiffen and fracture simultaneously to give high tensile strengths of the hydrogels.

On a structural level, high-strength macro-fibers made of polymers^{776,777}, steel⁷⁷⁸, glass^{779,780} and wood⁷⁸¹ have been utilized to strengthen hydrogels, and the tensile strengths of the resultant hydrogels are primarily determined by the strengths of the macro-fibers.



Figure 15. Implementation strategies for strong hydrogels with UPNs. Schematics on the implementation with **a.** polymer networks with high-functionality crosslinks, **b.** nano-/micro-fibrous polymer networks. **c.** confocal (left) and SEM (right) images of a fibrous PVA hydrogel with aligned fibers⁷⁶. **d.** nominal stress-stretch curves of the fibrous PVA hydrogels with aligned and randomly-oriented fibers⁷⁶. **c** and **d** are reproduced with permission from Ref⁷⁶. Copyright 2019 National Academy of Sciences.

5.3. Resilient: delay dissipation

Resilience. Resilience of soft materials such as elastomers and hydrogels is commonly defined as the ratio of energy released in deformation recovery to the energy required to induce the deformation of the materials⁷⁸². Let's consider a cylindrical sample under the uniaxial tensile test over a loading-unloading cycle (**Figure 16a**). The energy released in the unloading and the dissipated energy per unit volume of the sample are denoted as W_R and W_D , respectively. Therefore, the resilience *R* and the hysteresis ratio *H* of the material can expressed as (**Figure 16a**)^{73,782}

$$R = \frac{W_R}{W_R + W_D}, \ H = \frac{W_D}{W_R + W_D} = 1 - R$$
(25)

The resilience *R* and the hysteresis ratio *H* depend on the material properties and the loading conditions such as the applied stretch and the applied stretch rate. The resilience of soft materials has been measured with many experimental methods such as the cyclic tensile test and the dropping-ball test^{73,782}.

Design principle for resilient and tough hydrogels. Once a material is deformed to fracture, the elastic energy stored in the material is mostly dissipated^{73,74,783}, giving low energy recovery and thus low resilience of the fractured material. Therefore, the high resilience of hydrogels can only be designed up to the fracture of the hydrogels. A generic principle for the design of resilient hydrogels is to minimize the mechanical dissipation of the hydrogels within certain ranges of deformation that is commonly experienced by the hydrogels, or in short, to delay dissipation⁷³. Without loss of generality, we define a critical stretch for polymer chains in a hydrogel λ_R , below which the hydrogel can release most of the stored elastic energy during deformation recovery (i.e., $W_D \approx 0$, **Figure 16b**)⁷³. Therefore, according to **Eq. 25**, the hydrogel will give a high resilience under the condition,

$$\lambda \le \lambda_R \le \lambda_{\rm lim} \tag{26}$$

where λ and λ_{lim} are the stretch and the stretch limit of polymer chains in the hydrogel, respectively.

The design principle for resilient hydrogels also reconciles a pair of seemly contradictory properties, fracture toughness and resilience, in the following manner. The hydrogel is highly resilient under moderate deformation with $\lambda \leq \lambda_R$ (Figure 16b); however, when a crack attempts to propagate in the hydrogel, the chain stretch in the process zone around the crack tip can be much higher than λ_R , inducing substantial mechanical dissipation to toughen the hydrogel (Figure 16c). Indeed, biological hydrogels such as heart valves delay the mechanical dissipation up to supraphysiological deformation levels to achieve both high fracture toughness and high resilience (Figure 2)^{784,785}. Synthetic elastomers⁷⁸⁶, hydrogels⁷³ and hydrogel composites⁷⁷ have also been made both tough and resilient by following the design principle of delaying dissipation (Figures 16d-f).



Figure 16. Design principle for resilient and tough hydrogels: delay dissipation. a. definition and measurement of resilience. The relation of nominal stress *s* and stretch λ of a sample is measured under uniaxial tension in a loading-unloading cycle. W_R and W_D are the energy released in the unloading and the dissipated energy per unit volume of the sample, respectively. The resilience can be calculated as $R = W_R / (W_R + W_D)$. **b.** when the stretch is below a critical stretch λ_R , the hydrogel releases most of the stored elastic energy during deformation recovery, giving high resilience; when the stretch is above λ_R , the hydrogel dissipates substantial mechanical energy, giving high fracture toughness⁷³. **c.** the stretch in the process zone around the crack is usually much higher than λ_R , dissipating substantial mechanical energy and giving high fracture toughness⁷³. **d.** the nominal stress vs stretch curves of a PAAm-alginate hydrogel with $\lambda_R = 5^{-73}$. **e.** the measured deformation around a crack in the PAAm-alginate hydrogel with $\lambda_R = 5 \cdot$ **f.** the stretch in the process zone can be much higher than $\lambda_R = 5^{-73}$. **b** and **c** are reproduced with permission from Ref⁷³. Copyright 2014 Elsevier. **d**, **e** and **f** are reproduced with permission from Ref⁷³. Copyright 2014 Elsevier.

Implementation strategies for resilient and tough hydrogels. The ideal polymer networks are one common UPN architecture to implement the design principle for resilient hydrogels^{74,783}. Because the polymer chains in the ideal polymer networks have relatively uniform lengths and no entanglement, the hydrogels with the ideal polymer networks usually can be deformed without significant mechanical dissipation up to stretch limits, giving high resilience (Figure 17a)^{74,783}. It is also expected that the polymer networks with slidable crosslinks may be able to implement the design principle for resilient hydrogels, because the energy dissipated for sliding the crosslinks during their reconfiguration may be negligibly low. Despite being resilient, the ideal polymer networks and polymer networks with slidable crosslinks are not tough, since their fracture toughness is still the intrinsic fracture energy Γ_0 for fracturing a layer of polymer chains as discussed in Section 4.1.

The multimodal polymer networks including the interpenetrating polymer networks, semiinterpenetrating polymer networks, and polymer networks with high-functionality crosslinks usually begin to dissipate mechanical energy at very small deformation, because of the fracture and/or de-crosslink of very short polymer chains in the polymer networks. Such "early" dissipation gives narrow ranges of resilient deformation for the hydrogels in practical applications⁷³. To address the issue of "early" dissipation, Lin et al. have pre-stretched the interpenetrating polymer networks up to λ_R to fracture and/or de-crosslink susceptible short polymer chains and thus deplete possible dissipation mechanisms within the deformation range of λ_R (**Figure 17b**)⁷³. In subsequent tests, if the chain stretch in the hydrogel is below λ_R , the hydrogel is highly resilient, due to the lack of mechanical dissipation within this range (**Figure 17b**). However, as the polymer chains are stretched beyond λ_R , for example, in the process zone around the crack tip, some of the polymer chains will be further fractured and de-crosslinked to dissipate mechanical energy and toughen the hydrogel (**Figure 17b** and **Figures 16 c, e, h**). It is expected that other multimodal polymer networks such as the semi-interpenetrating polymer networks and polymer networks with high-functionality crosslinks⁵⁰ can be pre-stretched in a similar way to implement the design principle for resilient hydrogels (**Figure 17c**). Notably, when pre-stretching the multimodal polymer networks, the fracture and de-crosslink of polymer chains should be irreversible, so that the dissipation mechanism is irrecoverable once depleted⁷⁶.

The nano-/micro-fibrous polymer networks can naturally implement the design principle for resilient hydrogels by constituting the hydrogels with resilient nano-/micro-fibers (**Figures 17d** and e)⁷⁶. In addition, because the energy required to fracture and pull out the nano-/micro-fibers can be much higher than the energy for fracturing amorphous polymer chains, the resilient nano-/micro-fibrous hydrogels can also be tough⁷⁶.

Besides the abovementioned UPN architectures, some UPN interactions can also facilitate the implementation of the design principles for resilient hydrogels. The strong physical crosslinks such as crystalline domains provide the high-functionality crosslinks for some UPN architectures, which can be pre-stretched to give resilient hydrogels (**Figure 17c**)⁷⁵. Notably, the weak physical crosslinks and dynamic covalent crosslinks may be unsuitable to implement the design principle for resilient hydrogels, because of their reversible and dissipative nature⁷³.

On a structural level, resilient elastomeric macro-fibers have been embedded into resilient hydrogel matrices to give resilient yet tough hydrogel composites⁷⁷.



Figure 17. Implementation strategies for resilient and tough hydrogels with UPNs. a. ideal polymer networks are resilient up to fracture due to the lack of dissipation mechanism. b. pre-stretching interpenetrating polymer networks to λ_R can make them both resilient and tough. c. pre-stretching polymer networks with high-functionality crosslinks to λ_R can make them both resilient and tough⁷³. d. nano-/micro-fibrous polymer networks with resilient fibers can be both resilient and tough. e. the nominal stress-stretch curve of a resilient and tough nano-fibrous PVA hydrogel⁷⁶. e is reproduced with permission from Ref⁷⁶. Copyright 2019 National Academy of Sciences.

5.4. Tough adhesion: integrate tough dissipative hydrogels and strong interfacial linkages

Interfacial toughness. Interfacial toughness, or so-called practical work of adhesion, has been commonly used to characterize the capability of the interface of two adhered materials to resist fracture under mechanical loads. One common definition for the interfacial toughness between two adhered materials is the energy required to propagate a crack along the interface or in either material over a unit area measured in the undeformed state of the materials (**Figure 18a**)⁷⁸⁷. Depending on whether the crack propagates along the interface or in either material, the failure mode is called an adhesive failure or cohesive failure, respectively (**Figure 18a**). Quantitatively, the interfacial toughness Γ^{inter} can be expressed as,

$$\Gamma^{\text{inter}} = G_c = -\frac{dU}{dA} \tag{27}$$

where U is the total potential energy of the system, A is the crack area measured in the undeformed state, and G_c is the critical energy release rate that drives interfacial crack propagation. According to Eq. 27, the unit for the interfacial toughness is joule per meter squared (i.e., J m⁻²).

The interfacial toughness of soft materials such as elastomers and hydrogels has been measured with many experimental methods such as the 90-degree peeling test, the T-peeling test and the lap-shear test^{727,787}. For example, in the 90-degree peeling test, a layer of a hydrogel with thickness *T*, width *W* and length *L* ($L \gg W \gg T$) is bonded on a substrate, and a notch is introduced on the interface along the length direction (**Figure 18a**). The detached part of the hydrogel is further peeled off the substrate, while maintaining vertical to the substrate (**Figure 18a**). The measured force reaches a plateau $F_{plateau}$ as the peeling process enters the steady state, and the interfacial toughness is determined by dividing the plateau force $F_{plateau}$ by the width of the hydrogel sheet *W*, i.e. $\Gamma^{inter} = F_{plateau} / W$.

If a hydrogel with a conventional polymer network is strongly bonded on a substrate (e.g., via covalent bonds), the interfacial toughness is on the level of the hydrogel's fracture toughness or intrinsic fracture energy Γ_0 . This is because the fracture toughness of the hydrogel poses an upper limit for the interfacial toughness, since the cohesive failure mode may occur (**Figure 18c**)⁴⁹. Therefore, evaluated with typical parameters of conventional polymer networks, the interfacial toughness of the hydrogel is bounded by a few tens of joule per meter squared. If the hydrogel is adhered on the substrate via a low density of weak physical crosslinks such as hydrogen bonds and

electrostatic interactions, the interfacial toughness can be even lower, since the adhesive failure mode may occur (**Figure 18b**)⁴⁸.

Design principle for tough adhesion of hydrogels. As discussed in the previous part, if a hydrogel adheres to a substrate via a low density of weak physical crosslinks, a crack can easily propagate along the hydrogel-substrate interface, resulting in low interfacial toughness (**Figure 18b**). Therefore, the design of tough adhesion of hydrogels first requires strong interfacial linkages between the hydrogels and the adhered substrates, such as covalent bonds^{49,788,789}, strong physical crosslinks^{48,78,790}, connector polymers^{26,50,791,792}, and mechanical interlocks^{793,794}. In addition, because the interfacial crack can tilt into the bulk hydrogel and develop the cohesive failure mode (**Figure 18c**), the design of tough adhesion of hydrogels further requires high fracture toughness of the hydrogel matrices⁴⁹.

Overall, the design principle for tough adhesion of hydrogels is to integrate tough dissipative hydrogel matrices and strong interfacial linkages⁴⁹. When attempting to detach the tough hydrogel from the substrate, the strong interfacial linkages will hold the interfacial crack tip, allowing the bulk hydrogel to develop a process zone with substantial mechanical dissipation (**Figure 18d**). Quantitatively, the total interfacial toughness can be expressed as^{49,789}

$$\Gamma^{\text{inter}} = \Gamma_0^{\text{inter}} + \Gamma_D^{\text{inter}}$$
(28)

where Γ_0^{inter} , Γ_0^{inter} , and Γ_D^{inter} are the total interfacial toughness, the intrinsic interfacial toughness due to strong interfacial linkages, and the contribution of mechanical dissipation in the process zone to the total interfacial toughness, respectively.

Tough adhesion of biological hydrogels in animal bodies such as cartilages, tendons and ligaments on bones generally relies on the integration of tough hydrogels and strong interfacial linkages. However, only recently has this design principle been proposed⁴⁹ and implemented^{26,47,49,50,788,791,792,795} for tough adhesion of synthetic hydrogels on diverse substrate materials, including metals, ceramics, glass, silicone, elastomers, hydrogels and biological tissues. This is because the role of tough dissipative hydrogel matrices has been underexplored or underestimated in adhesion of hydrogels^{49,796,797}. Notably, strong interfacial linkages and/or bulk dissipation of the adherents have also been widely employed for tough bonding of engineering materials such as metals⁷⁹⁸ and rubbers^{799,800} on substrates.



Figure 18. Design principle for tough adhesion of hydrogels: integrate tough dissipative hydrogels and strong interfacial linkages. a. definition of interfacial toughness, and the 90-

degree peeling test to measure the interfacial toughness. *F* is the peeling force, $F_{plateau}$ is the plateau peeling force, and *W* is the width of the sample. The interfacial toughness can be calculated as $\Gamma^{inter} = F_{plateau} / W$ based on the values of $F_{plateau}$ and *W* measured in the 90-degree peeling test. **b**. weak interface can give the adhesive failure mode. **c**. brittle hydrogel matrix can give the cohesive failure mode. **d**. integration of tough dissipative hydrogels and strong interfacial linkages gives tough adhesion of hydrogels⁴⁹. The contributions of strong interfacial linkages and mechanical dissipation in the process zone to the total interfacial toughness are $\Gamma_0^{inter} = \Gamma_0^{inter} + \Gamma_D^{inter}$, respectively. The total interfacial toughness of the tough adhesion is $\Gamma^{inter} = \Gamma_0^{inter} + \Gamma_D^{inter}$. **d** is reproduced with permission from Ref⁴⁹. Copyright 2016 Springer Nature.

Implementation strategies for tough adhesion of hydrogels. Since the implementation of tough hydrogels has been discussed in Section 5.1, we will focus on how to implement the strong interfacial linkages to bond tough dissipative hydrogels on various substrates in this section. In order to achieve tough adhesion, the intrinsic interfacial toughness Γ_0^{inter} of the interfacial linkages should at least reach the level of the intrinsic fracture energy Γ_0 of tough hydrogels, i.e. over a few tens of joule per meter squared⁴⁹. Given this requirement on the intrinsic interfacial toughness, the strong interfacial linkages have been commonly implemented with covalent bonds, strong physical crosslinks, connector polymers, and mechanical interlocks.

Covalent bonds have been widely adopted to strongly anchor polymer chains in tough hydrogels' UPNs (as discussed in Section 5.1) on various substrates. The commonly-used covalent bonds for tough adhesion of hydrogels include carbon-carbon, carbon-nitrogen, carbon-sulfide, carbon-oxygen, and silicon-oxygen bonds (Figure 5c)⁸⁰¹. In order to form these covalent bonds, the hydrogels and substrates are usually designed to possess functional groups such as crosslinkable unsaturated bond (to form carbon-carbon bond)⁸⁰², amine group (to form carbon-nitrogen bond)⁴⁹, thiol group (to form carbon-sulfide bond)⁸⁰³, hydroxyl and carboxyl group (to form carbon-oxygen bond), and silanol group (to form silicon-oxygen bond)⁸⁰⁴ (Figure 19a). According to the Lake-Thomas model, the intrinsic interfacial toughness Γ_0^{inter} of polymer chains covalently anchored on a substrate can be expressed as

$$\Gamma_0^{\text{inter}} = M^{\text{inter}} N U_f \tag{29}$$

where M^{inter} is the number of covalently-anchored polymer chains on a unit area of the substrate in the undeformed reference state, N is the number of Kuhn monomers per polymer chain, and U_f is the lower value of the energy required to fracture either the Kuhn monomer or the covalent bond on the substrate. According to **Eq. 29**, anchoring longer polymer chains with a higher density of covalent bonds on a substrate will give a higher value of the intrinsic interfacial toughness^{49,789}.

Strong physical crosslinks including crystalline domains, glassy nodules, and high-density physical bonds such as hydrogen bonds can also strongly adhere tough hydrogels on substrates (**Figure 19b**)^{78,757,803-811}. Since the crystalline domains and glassy nodules usually act as high-functionality crosslinks, each of them may anchor multiple polymer chains on the substrate, further enhancing the intrinsic interfacial toughness Γ_0^{inter} .

Connector polymers^{799,812} have been employed to strongly bond elastomers and hydrogels on substrates (**Figure 19c**). In this case, the substrates usually take the form of polymer networks (i.e., elastomers and hydrogels) as well. To provide strong interfacial linkages, the connector polymers can form covalent crosslinks^{26,50}, interlocked loops^{791,792,803,807,813}, and/or strong physical crosslinks⁷⁹² with the polymer networks of both the hydrogels and the substrates. Specifically, the strong physical crosslinks can be crystalline domains, glassy nodules, and/or high-density weak physical crosslinks⁷⁹². The connector polymers can be polymerized from monomers in the two polymer networks^{26,791,804} or can be directly added on the interface of the two polymer networks⁷⁹².

Mechanical interlocks between tough hydrogels and substrates usually occur at length scales from micrometers to millimeters (**Figure 19d**). One commonly-used method is to impinge precursor solutions of tough hydrogels into porous substrates and then form tough hydrogels that are mechanically interlocked with the substrates⁷⁹³. Similarly, the surfaces of the substrates can be roughened or patterned to enhance the strength of mechanical interlocks with tough hydrogels^{794,814,815}. As a special yet interesting case, hydrogels have been fabricated into dried microneedles, which can pierce into a soft substrate such as biological tissues and then swell to form mechanical interlocks⁸¹⁶.

Inspired by the adhesive proteins found in mussels, catechol chemistry has been widely adopted to achieve various types of interfacial linkages between hydrogels and substrates (**Figure 19e**)^{579,790}. Catechol can form both covalent and physical crosslinks with various functional groups (**Figure 19e**). Upon oxidation to quinone, catechol can form a covalent bond with nucleophiles (e.g, amine and thiol) via the Michael addition as well as a strong coordination complex with metal oxides⁸¹⁷. The hydroxyl groups of catechol can form electrostatic interaction with metal oxides as well as hydrogen bonds with hydrophilic substrates. The benzene ring of catechol can further form cation- π interaction with positively charged functional groups, π - π stacking with benzene functional groups, and hydrophobic interaction with hydrophobic functional groups on substrates^{790,817}. While catechol chemistry has been widely utilized for adhesion of hydrogels to various substrates, the interfacial toughness of the adhesion achieved only by catechol-based interfacial linkages is not high⁸¹⁸, highlighting the importance of tough dissipative hydrogel matrices on top of the interfacial linkages to achieve tough adhesion⁸¹⁹.



Figure 19. Implementation strategies for tough adhesion of hydrogels with UPNs. The tough hydrogels with UPNs are bonded on substrates via various types of strong interfacial linkages: **a.** covalent bonds, **b.** strong physical crosslinks, **c.** connector polymers, and **d.** mechanical interlocks. **e.** catechol interactions can implement various types of strong interfacial linkages. **e** is reproduced with permission from Ref^{817,820}. Copyright 2019 Wiley and 2017 Wiley.

5.5. Fatigue-resistant: pin fatigue cracks with intrinsically high-energy phases

Fatigue threshold. The word "fatigue" has been used to describe many symptoms observed in materials under prolonged loads, including materials with or without precut cracks under prolonged static or cyclic loads^{821,822}. In this section, we will focus on the fatigue fracture of hydrogels with precut cracks under cyclic loads (**Figure 20a**), because this is one of the most common failure modes of hydrogels in mechanically dynamic environments such as artificial cartilages⁸²³ and soft robots²⁶. Fatigue threshold has been commonly used to characterize a material's resistance to fatigue crack propagation under cyclic loads. The fatigue threshold is defined as the minimal fracture energy at which fatigue crack propagation occurs under infinite cycles of loads^{55,56}. Quantitatively, the fatigue threshold Γ_{FT} can be expressed as,

$$\Gamma_{FT} = G_c \left(\frac{dc}{dN} \to 0 \right) \tag{30}$$

where G is the energy release rate to drive crack propagation under each cycle of load, G_c is the minimal energy release rate at which crack propagation occurs under infinite cycles of loads (i.e. $dc/dN \rightarrow 0$), c is the length of the crack, N is the cycle number of the applied load, and dc/dN gives the crack extension per cycle.

The fatigue threshold of soft materials such as elastomers and hydrogels have been measured with various experimental methods such as the pure-shear fatigue-fracture test and the single-notch fatigue-fracture test⁷⁵. For example, in the pure-shear fatigue-fracture test, two identical pieces of a hydrogel are fabricated with the same thickness *T*, width *W* and height *H*, where W >> H >> T (Figure 20a). Both pieces of samples are clamped along their long edges (i.e., along the width direction) with rigid plates. The first sample is repeatedly pulled to a stretch of $\lambda_{applied}$ times of its undeformed height to measure the nominal stress *s* vs stretch λ relation, and the corresponding energy release rate can be calculated as $G = H \int_{1}^{\lambda_{applied}} sd\lambda$, which is a function of the cyclic number *N* (Figure 20a). Thereafter, a notch with a length of ~0.5W is introduced into the second sample, which is then repeatedly pulled to the same stretch $\lambda_{applied}$ to measure the crack length *c* as a function of the cyclic number *N*. The pure-shear fatigue-fracture tests are repeated for different values of the applied stretch $\lambda_{applied}$ (i.e., different energy release rate *G*), and a curve of dc/dN vs *G* can be obtained (Figure 20a). The fatigue threshold Γ_{FT} is then determined by intersecting the curve of dc/dN vs *G* with the *G* axis (i.e., when $dc/dN \rightarrow 0$). Notably, the

fatigue-fracture tests of hydrogels are commonly carried out in aqueous environments to avoid dehydration of the hydrogels under prolonged loads^{75,76}. For further discussion on the theory and experiments for the fatigue of hydrogels, a comprehensive review paper on this topic is recommended⁸²².

Design principle for fatigue-resistant hydrogels. As discussed in Section 5.1, a hydrogel can be designed tough by building mechanical dissipation into stretchy polymer networks⁶⁷. The mechanical dissipation in the process zone around the crack tip can dramatically enhance the fracture toughness of the hydrogel. However, the mechanisms for irreversible dissipation such as fracturing polymer chains in the process zone are usually depleted under cyclic loads. The mechanisms for reversible dissipation such as reversible crosslinks, once depleted, usually cannot recover in time to resist fatigue crack propagation in future cycles of loads (Figure 20b)^{55,56,824}. Consequently, the fatigue threshold of hydrogels and elastomers is their intrinsic fracture energy^{55,56,824},

$$\Gamma_{FT} = \Gamma_0 \tag{31}$$

Therefore, it is clear that the design of fatigue-resistant hydrogels usually cannot rely on mechanical dissipation in the bulk hydrogel matrices.

The design principle for fatigue-resistant hydrogels is to make the fatigue crack encounter and fracture objects with energies per unit area much higher than that for fracturing a single layer of polymer chains, or in short, to pin fatigue crack by intrinsically high-energy phases (**Figure** 20c)⁷⁵. The intrinsically high-energy phases that have been exploited for the design of fatigueresistant hydrogels include nano-crystalline domains (**Figure 21a**)⁷⁵, nano-/micro-fibers (**Figure** 21b)⁷⁶, microphase separations (**Figure 21c**)^{822,825}, and macroscale composites (**Figure 21d**)⁷⁷. In addition, because the design of fatigue-resistant hydrogels does not rely on mechanical dissipation in the bulk hydrogels, fatigue-resistant hydrogels usually demonstrate low hysteresis ratio *H* and high resilience *R* (**Eq. 23**)^{75,76,826}. Notably, biological hydrogels such as muscles, tendons and ligaments commonly possess intrinsically high-energy phases such as nano-/micro-fibers, usually arranged in hierarchical architectures, to achieve high fatigue threshold (**Figure 2**).



Figure 20. Design principle for fatigue-resistant hydrogels: pin cracks by intrinsically highenergy phases. **a**. definition of fatigue threshold, and the pure-shear method to measure fatigue threshold. *G* is the energy release rate, *c* is the crack length, and *N* is the cycle number. The fatigue threshold Γ_{FT} is determined by intersecting the curve of dc/dN vs *G* with the *G* axis. **b**. dissipation mechanisms such as reversible crosslinks in tough hydrogels are depleted over cyclic loads, not contributing to the fatigue threshold. **c**. fatigue crack is pinned by intrinsically highenergy phases in fatigue-resistant hydrogels⁷⁵. **b** and **c** are reproduced with permission from Ref⁷⁵. Copyright 2019 American Association for the Advancement of Science.

Implementation strategies for fatigue-resistant hydrogels. The design principle for fatigue-resistant hydrogels has been implemented with the UPNs that possess intrinsically high-energy phases^{75-77,822,825}. In order to effectively pin fatigue cracks, the density of the intrinsically high-energy phases in the UPNs should be sufficiently high⁷⁵.

High-functionality crosslinks such as nano-crystalline domains can effectively play the role of intrinsically high-energy phases in the UPNs (**Figure 21a**). The energy required to pull out a polymer chain from a nano-crystalline domain can be multiple times higher than that to fracture the same polymer chain, and the energy required to mechanically damage the nano-crystalline domains can be multiple times higher than that to fracture the corresponding amorphous polymer chains (**Figure 21e**)⁷⁸. Consequently, it has been shown that enhancing the crystallinity of a PVA hydrogel from 0.2 wt. % to 18.9 wt. % by dry-annealing the hydrogel can increase its fatigue threshold from 10 Jm⁻² to 1,000 Jm⁻², reaching the level of fatigue-resistant biological hydrogels such as cartilages⁷⁵. Since the size of the crystalline domains in the PVA hydrogel has been

measured to be a few nanometers, it is the nano-crystalline domains that play the role of intrinsically high-energy phases (**Figure 21a**). It is expected other UPNs with sufficiently high densities of high-functionality crosslinks such as crystalline domains and glassy nodules can also implement the design principle for fatigue-resistant hydrogels. It should be further noted that hydrogels with high densities of rigid crystalline domains and glassy nodules can be much stiffer than common hydrogels⁷⁵, and such high stiffness may be undesirable for many applications of hydrogels.

Nano-/micro-fibers can also act as intrinsically high-energy phases in the UPNs to implement the design principle for fatigue-resistant hydrogels (**Figure 21b**). The energy required to fracture a nano-/micro-fiber can be much higher than that to fracture the corresponding amorphous polymer chains, because of the synergistic elongation and stiffening of the bundled polymer chains in the fiber⁷⁶. Based on this implementation strategy, it has been shown that introducing nano-fibers into a PVA hydrogel by freeze-thawing the hydrogel can enhance its fatigue threshold from 10 Jm⁻² to 310 Jm⁻². In particular, if the nano-fibers are aligned perpendicular to the fatigue crack by pre-stretching the hydrogel, the measured fatigue threshold further increases to 1,250 Jm⁻² (**Figures 21f** and **g**)⁷⁶. In addition, because the nano-fibers can be made compliant, stretchable and strong by using a low density of nano-crystalline domains to bundle polymer chains (**Figure 15b**), the resultant nano-fibrous hydrogel integrates high compliance, stretchability and strength together with high fatigue threshold – mimicking the combinational mechanical properties of biological muscles⁷⁶.

Phase separations in hydrogels can also enhance the fatigue threshold of the hydrogels^{822,825}, possibly because the energy required to fracture the separated phases is higher than that to fracture the corresponding amorphous polymer chains. The UPN interactions including reversible covalent bonds and weak physical crosslinks play critical roles in inducing the phase separations in the hydrogels^{822,825}.

On a structural level, macroscale resilient elastomer fibers have been embedded in a resilient hydrogel to form a macroscale composite⁷⁷. Since it requires much higher energy to fracture the elastomer fibers than a layer of amorphous polymer chains, a fatigue threshold over 1,000 Jm⁻² has been achieved for the macroscale composite (**Figure 21d**)⁷⁷.



Figure 21. Implementation strategies for fatigue-resistant hydrogels with UPNs. Fatigue cracks can be pinned by intrinsically high-energy phases including **a**. nano-crystalline domains⁷⁵, **b**. nano-/micro-fibers⁷⁶, **c**. micro-phase separations⁸²⁵, and **d**. macro-fibers⁷⁷. **e**. Molecular dynamic simulation for pulling a polymer chain out of a PVA nano-crystalline domain and for fracturing the same polymer chain⁷⁸. *d* is the displacement of one end of the polymer chain, and *U* is the energy required to achieve the displacement. **f**. confocal microscope image of a crack pinned by nano-fibers in a nano-fibrous PVA hydrogel, and **g**. measurement of the fatigue threshold of the nano-fibrous PVA hydrogel⁷⁶. *G* is the energy release rate, *c* is the crack length, and *N* is the cycle number. **a** is reproduced with permission from Ref⁷⁵. Copyright 2019 American Association for the Advancement of Sciences, **b** is reproduced with permission from Ref⁸²⁵. Copyright 2020 National Academy of Sciences, **d** is reproduced with permission from Ref⁷⁷. Copyright 2019 Elsevier, **e** is reproduced with permission from Ref⁷⁸. Copyright 2020 Springer Nature, **f** and **g** are reproduced with permission from Ref⁷⁶. Copyright 2019 National Academy of Sciences.

5.6. Fatigue-resistant adhesion: strongly bond intrinsically high-energy phases on interfaces

Interfacial fatigue threshold. The interfaces of adhered materials can suffer from fatigue failure under prolonged loads, including interfaces with or without precut cracks under prolonged static or cyclic loads. In this section, we will focus on the fatigue fracture of hydrogels adhered on
substrates with precut cracks on their interfaces under cyclic loads (**Figure 22a**). Depending on whether the fatigue crack propagates along the interface or tilts into the hydrogel under cyclic loads, the failure mode is called adhesive failure or cohesive failure, respectively (**Figure 22a**)⁷⁸. Interfacial fatigue threshold has been commonly used to characterize the capability of adhered materials to resist interfacial fatigue crack propagation following either failure mode under cyclic loads. The interfacial fatigue threshold is defined as the minimal fracture energy at which interfacial crack propagation occurs under infinite cycles of loads^{78,827-829}. Similar to the fatigue threshold, the interfacial fatigue threshold Γ_{FT}^{inter} can be expressed as,

$$\Gamma_{FT}^{\text{inter}} = G_c \left(\frac{dc}{dN} \to 0 \right)$$
(32)

where G is the energy release rate to drive interfacial crack propagation under each cycle of load, G_c is the minimal energy release rate at which interfacial crack propagation occurs under infinite cycles of loads (i.e. $dc/dN \rightarrow 0$), c is the length of the crack, N is the cycle number of the applied load, and dc/dN gives the crack extension per cycle.

The interfacial fatigue threshold of soft materials such as elastomers and hydrogels has been measured with many experimental methods such as the cyclic 90-degree peeling test, the cyclic T-peeling test and the cyclic lap-shear test^{78,827-829}. For example, in the cyclic 90-degree peeling test^{78,827}, a layer of a hydrogel with thickness *T*, width *W* and length *L* (L >> W >> T) is bonded on a substrate, and a notch is introduced on the interface along the length direction (**Figure 22a**). A force *F* is repeatedly applied on the detached part of the hydrogel, while maintaining the detached part vertical to the substrate (**Figure 22a**). The applied force *F* gives the energy release rate G = F/W, where *W* is the width of the hydrogel sheet. The interfacial crack length *c* is then measured as a function of the cyclic number *N*. The cyclic 90-degree peeling tests are repeated for different values of the applied force *F* (i.e., different energy release rate *G*), and a curve of dc/dNvs *G* can be obtained (**Figure 22a**). The interfacial fatigue threshold Γ_{FT}^{inter} is determined by intersecting the curve of dc/dN vs *G* with the *G* axis (i.e., when dc/dN=0). Notably, the interfacial fatigue fracture tests of hydrogels are commonly carried out in aqueous environments to avoid dehydration of the hydrogels under prolonged loads⁷⁸.

Design principle for fatigue-resistant adhesion of hydrogels. As discussed in Section 5.4, tough adhesion of hydrogels on substrates relies on the integration of tough dissipative hydrogel matrices and strong interfacial linkages (Figure 18)⁴⁹. The strong interfacial linkages can hold the interfacial crack tip, while the mechanical dissipation in the process zone around the crack tip can dramatically enhance the total interfacial toughness of the adhesion. However, similar to the situation in fatigue fracture of hydrogels^{55,56,824}, because the mechanical dissipation in bulk hydrogel matrices is usually depleted or not timely accessible after cyclic loads, such dissipation usually cannot contribute to resisting interfacial fatigue crack propagation (Figures 22b and c)^{78,829}. Consequently, the interfacial fatigue threshold of hydrogels and elastomers is their intrinsic interfacial toughness^{78,829},

$$\Gamma_{FT}^{\text{inter}} = \Gamma_0^{\text{inter}} \tag{33}$$

Because interfacial cracks can tilt into bulk hydrogels and develop the cohesive failure mode (**Figure 22c**), the design of fatigue-resistant adhesion of hydrogels first requires fatigue-resistant hydrogel matrices that possess sufficiently high densities of intrinsically high-energy phases⁷⁵. Notably, hydrogel matrices that are only tough but not fatigue-resistant are unsuitable for the design of fatigue-resistant adhesion, owing to the depletion of dissipation over cyclic loads (**Eq. 33**). To further avoid the adhesive failure mode under cyclic loads (**Figure 22b**), fatigue cracks on the interfaces also need to be pinned by intrinsically high-energy phases strongly bonded on the interfaces as well (**Figure 22d**).

Therefore, the design principle for fatigue-resistant adhesion of hydrogels, in short, is to strongly bond intrinsically high-energy phases on interfaces⁷⁸. While the intrinsically high-energy phases that have been exploited for the design of fatigue-resistant adhesion include nano-crystalline domains⁷⁸ and long polymer chains⁸²⁸, other candidates such as nano-/micro-fibers can be explored in the future. Not surprisingly, biological hydrogels including tendons, ligaments and cartilages all rely on strongly bonding nano-crystalline domains and nano-/micro-fibers on their interfaces with bones to achieve fatigue-resistant adhesion⁷⁵⁷.



Figure 22. Design principle for fatigue-resistant adhesion of hydrogels: strongly bond intrinsically high-energy phases on interfaces. a. definition of interfacial fatigue threshold, and the 90-degree cyclic peeling test to measure the interfacial fatigue threshold. *F* is the applied peeling force, *W* is the width of the sample, *G* is the energy release rate, *c* is the crack length, and *N* is the cycle number. The interfacial fatigue threshold Γ_{FT}^{inter} is determined by intersecting the curve of dc / dN vs *G* with the *G* axis. b. fatigue-crack propagation along the interface gives the adhesive failure mode. c. fatigue-crack propagation in the hydrogel gives the cohesive failure mode. d. fatigue-crack pinned by intrinsically high-energy phases on the interface and in the bulk hydrogel⁷⁸. d is reproduced with permission from Ref⁷⁸. Copyright 2020 Springer Nature.

Implementation strategies for fatigue-resistant adhesion of hydrogels. The design of fatigue-resistant hydrogels relies on achieving sufficiently high densities of intrinsically high-energy phases in the UPNs, as discussed in **Section 5.6**. In this section, we will focus on how to strongly bond the intrinsically high-energy phases on substrates to implement the design principle for fatigue-resistant adhesion.

High-functionality crosslinks such as nano-crystalline domains usually can play the role of intrinsically high-energy phases in the UPNs. For example, nano-crystalline domains in PVA hydrogels have been strongly bonded on diverse substrates including glass, ceramics, metals and elastomers via high-density hydrogen bonds (**Figure 23a**)⁷⁸. Molecular dynamic simulations show that the energy required to pull a polymer chain out of the interface is much higher than the energy required to fracture the same polymer chain or to pull the same polymer chain out of the nano-crystalline domain, implying high intrinsic interfacial toughness of the adhesion $\Gamma_0^{\text{inter 78}}$. As a result, the interfacial fatigue threshold of the PVA-substrate system measured in phosphate-buffered saline reaches up to 800 Jm⁻², similar to those of tendon/ligament/cartilage-bone interfaces (**Figure 23d** and **e**). In addition, the failure mode of the PVA-substrate systems observed in the interfacial fatigue fracture tests follows the cohesive failure, indicating the critical role of intrinsically high-energy phases in the bulk hydrogels (i.e., fatigue-resistant hydrogels) for the design of fatigue-resistant adhesion⁷⁸.

It is expected that covalent bonds may be able to strongly bond the intrinsically high-energy phases such as nano-crystalline domains and nano-/micro-fibers on substrates as well (**Figure 23b**). In addition, curing precursor solutions of fatigue-resistant hydrogels on porous, roughened or patterned substrates can lead to mechanical interlocks that may strongly bond intrinsically high-energy phases on the hydrogel-substrate interfaces (**Figure 23c**)⁸³⁰.



Figure 23. Implementation strategies for fatigue-resistant adhesion of hydrogels with UPNs. The intrinsically high-energy phases can be strongly bonded on the substrates via **a.** high-density physical bonds such as hydrogen bonds⁷⁸, **b.** covalent bonds, and **c.** mechanical interlocks. **d.** measurements of the fatigue thresholds of tough adhesion and fatigue-resistant adhesion of hydrogels on substrates⁷⁸. **e.** photos of interfacial crack propagation in a cyclic peeling test for tough adhesion (top) and fatigue-resistant adhesion (bottom) of hydrogels on substrates⁷⁸. **a, d** and **e** are reproduced with permission from Ref⁷⁸. Copyright 2020 Springer Nature.

5.7. Implementation strategies for extreme mechanical properties with unconventional polymer networks

While the design principles and implementation strategies for hydrogels with extreme mechanical properties have been discussed in **Sections 5.1-5.6**, we will provide an overview of the design process and implementation strategies in this section.

Since the design principles discussed in Sections 5.1-5.6 are general and abstract, it is usually more intuitive to begin the design process with specific UPN architectures and/or UPN interactions than with a design principle. The commonly-used UPN architectures that can give extreme mechanical properties include interpenetrating polymer networks, semi-interpenetrating polymer networks, polymer networks with high-functionality crosslinks, and nano-/micro-fibrous polymer networks. The commonly-used UPN interactions that can give extreme mechanical properties include various types of strong physical crosslinks, weak physical crosslinks, and dynamic covalent crosslinks. Let's imagine that the selected UPN is subjected to the relevant modes of mechanical loads such as tension, compression, shear, fracture, fatigue, and/or peeling. If the selected UPN under mechanical loads seems to be able to implement the design principle for the desired property, one can proceed to select polymers and crosslinks such as those discussed in Section 2 and Section 4 for the design and fabrication of the hydrogel. Furthermore, it may be difficult to initiate the design process by considering both UPN architectures and UPN interactions simultaneously; in this case, we can first test whether a UPN architecture will likely implement the design principle, and then further add UPN interactions into the UPN architecture to facilitate the implementation. For example, in order to design a fatigue-resistant hydrogel, we can begin with a polymer network with high-functionality crosslinks, because a sufficiently high density of high-functionality crosslinks can act as intrinsically high-energy phases to pin fatigue cracks. Furthermore, strong physical crosslinks such as crystalline domains and glassy nodules can be added into the polymer network as the intrinsically high-energy phases to facilitate the implementation of the design principle. Indeed, dry-annealed PVA with high densities of nanocrystalline domains has been selected to implement the design principle for fatigue-resistant hydrogels⁷⁵.

Alternative design and implementation strategies are through the mimicry of the UPNs of biological hydrogels that possess the desired extreme mechanical properties (**Figure 2**). Because biological hydrogels have exploited various types of UPNs to implement the design principles discussed in **Sections 5.1-5.6**, we can simply begin the design process by replicating biological hydrogels' UPNs (**Figure 2**). However, biological hydrogels' UPNs, commonly featuring hierarchical and gradient structures (**Figure 2**), can be more complex than the UPNs discussed in **Section 4**; therefore, we should only mimic the essential characteristics of the biological UPNs that enable the desired mechanical properties. As an example, tendons, ligaments and cartilages all feature fatigue-resistant adhesion on bones, owing to nano-/micro-fibers and nano-crystalline domains strongly anchored on the interfaces (**Figure 2**). By strongly anchoring nano-crystalline

domains in synthetic PVA hydrogels on substrates, bioinspired fatigue-resistant adhesion of PVA hydrogels on diverse solid substrates have been recently achieved (**Figures 23d** and e)⁷⁸.

Hydrogel	Young's modulu	Water conten	Nominal tensile	Fracture toughness	Fatigue thresho	Resilie nce	Refere nce
	s (kPa)	t (vol. %)	strength (kPa)	(J m ⁻²)	ld (J m ⁻²)	(%)	
Freeze-thawed PVA	100	85	1,200	100-500	310	30	76,481,831
Directional freeze-thawed PVA	30-100	88-90	300-1,200	160-420	-	40-50	832
PAMPS- PAAm	100- 1,000	90	1,500	1,000- 3,000	200- 400	30-50	70,733,833, 834
PAAm- alginate	10-100	90	170	8700	53	30-40	490
Agar/PAAm	100	80	1,000	-	-	40-50	835
Prestretched PAAm- alginate	2	90	-	1,900	-	95	73
Polyampholyte	100- 2,200	50-70	100-2,000	1,000- 4,000	67-71	10-20	825,836
PNaSS/PMPT C	10-1,000	50	3,300	7,700	-	10-20	566
Tetra-PEG	18	95	10	10-30	-	~100	74,333,837
Dry annealed PVA	300- 9,000	58-75	1,000-9,000	1,000- 9,000	300- 1,000	40-50	75
Mechanically trained PVA	200	84	5,000	1,200	1,200	~100	76
Elastomer- hydrogel composites	200	60-70	180-250	4,500	1,290	90	77
Dual- crosslinked p(AAm- <i>co</i> - AAc)	500- 3,500	60-70	3,000-6,500	-	-	20-40	838
Nanoclay reinforced PNIPAM	0.8-26	90	27-300	-	-	-	839
Wood hydrogel	200,000	65	500-35,000	-	-	-	781

Table 3. Typical examples of hydrogels that possess extreme mechanical properties

Fiber	660-	100-	200-1,000	6,000-	-	70-80	777
reinforced	6,370	700		32,000			
PAAm-							
Alginate							
PA-GF	-	38	-	100,000-	-	-	840
				400,000			
Wool	50-7,000	86	200-700	-	-	-	778
reinforced							
PAAm-							
alginate							
PNAGA	50-200	70	100-1,100	200-1,100	-	60	750

- Indicates not reported

	n • 1	1	61 1	1 11 •	41 4	4	• •
l able 4	I vnical	examples	of hvdra	ngel adhesions	that noccess e	vfreme mech	anical properties
	Jpicar	champies v	or nyur (55 ^{cl} autorious	mar possess c		annear properties

Hydrogel adhesion	Water content (vol. %)	Young's modulus (kPa)	Interfacial toughness (J m ⁻²)	Interfacial fatigue threshold (J	Refere nce
	(())	((0)	m ⁻²)	
PAAm-	90	10-100	1,000-1,700	68	78,831
alginate to					
non-porous					
surfaces			-10		17
Tissue double-	90	7.5-15	710	-	4/
sided tapes to					
tissues					50.820
PAAm-	90	12	580	24.4	50,829
alginate to					
tissues					700
PAAm-	90	10-100	1,000	-	/88
alginate to					
elastomers					
PAMPS-	90	-	200-900	-	793
PAAm to					
porous					
surfaces					
Dry annealed	38-68	-	8,000	800	78
PVA to glass					
PDA-clay-	80	100-200	-	-	841
PAAm to					
glass					
Dopa-modified	90	2	-	-	842
PEG to porcine					
skin					

PAAm to	90	-	1,400	300	828
polyester cloth					

- Indicates not reported.

6. Design of hydrogels with extreme physical properties

In addition to the extreme mechanical properties discussed in **Section 5**, the design of hydrogels that possess extreme physical properties has attracted escalating research interests in recent years. Examples of hydrogels' extreme physical properties under development and exploration include high electrical conductivity⁶⁴², patterned magnetization⁸⁴³, high refractive index and transparency^{844,845}, tunable acoustic impedance⁴⁴, and self-healing⁸⁴⁶. Unlike the extreme mechanical properties discussed in **Section 5**, many of the extreme physical properties do not have embodiments in biological hydrogels. Nevertheless, these extreme physical properties can be of similar importance as the extreme mechanical properties to hydrogels' various applications, especially to the nascent applications of hydrogel machines²⁰. In this section, we will briefly discuss the design principles and implementation strategies for hydrogels to possess these extreme physical properties, while bearing in mind that many works in this field are still in the initial stage of development.

6.1. Electrically conductive: percolate electrically conductive phases

Electrical conductivity is critical for hydrogels' nascent applications such as bioelectrodes for stimulation and recording of neural activities in bioelectronics⁶⁴² and electrodes for supercapacitors and batteries in energy storage^{20,847}. However, the electrical conductivity of common hydrogels is less than a few Siemens per meter, on the same level as that of saline water⁶⁴². Compared to metals, carbon and conducting polymers, common hydrogels are usually deemed to be electrically nonconductive.

The design principle for electrically conductive hydrogels is to embed electrically conductive phases such as liquid metals, metallic nanowires, carbon nanotube, graphene and conducting polymers in hydrogel matrices and make the conductive phases form percolated networks, or in short, to percolate electrically conductive phases (Figure 24a)^{642,848,849}. In particular, conductive hydrogels based on conducting polymers have attracted great interests recently, owing to their unique polymeric nature as well as favorable electrical and mechanical biocompatibility^{31,32,116,495,850-853}. and For example, properties, stability, poly(3,4ethylenedioxythiophene):poly(styrene sulfonate) (PEDOT:PSS) has been made into pure conducting polymer hydrogels that achieve high electrical conductivity over a few thousand Siemens per meter and superior biocompatibility (Figure 24c)^{31,32,116}. In addition to electrically conductive hydrogels, ionically conductive hydrogels have also been intensively developed as stretchable and transparent ionic conductors for various applications (Figure 24b)²⁷. The conductive phases in ionically conductive hydrogels are usually high concentrations of salt ions. For a further detailed discussion on various types of conductive hydrogels, a recent review paper is recommended⁶⁴².



Figure 24. Design principle for hydrogels with high electrical conductivity: percolate electrically conductive phases. **a.** hydrogels with percolated electrically conductive fillers. **b.** hydrogels with ionically conductive salt solvents. **c.** hydrogels based on conducting polymers. The bottom panel of **a** is reproduced with permission from Ref⁸⁴⁹. Copyright 2013 American Association for the Advancement of Science. The bottom panel of **b** is reproduced with permission from Ref²⁷. Copyright 2014 American Association for the Advancement of Science. The bottom panels of **c** are reproduced with permission from Ref¹¹⁶ (left) and Ref³¹ (right). Copyright 2020 Springer Nature and 2019 Springer Nature.

6.2. Magnetized: embed magnetic particles and pattern ferromagnetic domains

Soft materials such as elastomers and hydrogels with ferromagnetic domains or magnetization have been intensively developed and explored for biomedical applications such as drug delivery and minimally invasive surgery^{28,38,843,854-857}, owing to their mechanical compliance, potential biocompatibility, and capability of fast deformation under applied magnetic fields. Common hydrogels are usually diamagnetic and do not contain ferromagnetic domains, possessing similar magnetic properties as water. Therefore, subjected to applied magnetic fields, common hydrogels cannot be actuated to deform, exert forces, or release substances.

The design principle for hydrogels to possess patterned magnetization is to embed magnetic components such as hard-magnetic, soft-magnetic or super-paramagnetic particles in the hydrogels matrices where ferromagnetic domains may be further patterned, or in short, *to embed magnetic particles and pattern ferromagnetic domains* (Figure 25) ^{38,843,854-857}. In particular, hard-magnetic particles such as neodymium iron boron (NdFeB) particles after magnetic saturation can

retain their magnetization under actuation magnetic fields, because of the high coercivity of the hard-magnetic particles (**Figure 25a**). Therefore, patterned ferromagnetic domains can be programmed into elastomers and hydrogels embedded with hard-magnetic particles. Subjected to actuation magnetic fields, the elastomers and hydrogels with the patterned ferromagnetic domains can quickly transform among various shapes^{38,854-857}. Recently, 3D printing has been further employed as an effective method to program complex 3D shapes as well as domain patterns in ferromagnetic elastomers and hydrogels (e.g., **Figures 25b** and **c**)^{854,857}. It should be noted that magnetic particles can be corrosive in the aqueous environments of hydrogel matrices. To enhance their chemical stability in hydrogel matrices, the magnetic particles have been coated with protective layers such as silica layers (**Figures 25b** and **c**)³⁸.



Figure 25. Design principle for hydrogels and elastomers with patterned magnetization: embed magnetic particles and pattern ferromagnetic domains. a. typical relations of applied magnetic field H and magnetization M for paramagnetic, soft-magnetic, and hard-magnetic materials. M_r and H_c are the residual magnetization and coercivity of the hard-magnetic material, respectively. b. hard-magnetic particles can be embedded into an elastomer/hydrogel matrix, in which ferromagnetic domains can be patterned by 3D printing. c. Photos of the resultant magnetic soft material before and after magnetic actuation. a is reproduced with permission from Ref⁴⁰. Copyright 2020 Elsevier. b and c are reproduced with permission from Ref³⁹. Copyright 2018 Springer Nature.

6.3. High reflective index and transparency: uniformly embed high-refractive-index non-scattering nano-phases.

Various optical applications of hydrogels such as ophthalmic lenses^{21,380,858} and optical fibers^{42,859} require high refractive indices and high transparency of the hydrogels (**Figure 26a**). The refractive indices of common hydrogels are around 1.333, similar to that of water. One general strategy to enhance the refractive indices of hydrogels is to uniformly embed nano-phases such as nano-particles^{844,845} and nano-crystalline domains with high refractive indices in the hydrogel matrices. However, the refractive-index mismatch between the nano-phases and hydrogel matrices may lead

to substantial undesirable light scattering, reducing the transparency of the hydrogels (**Figure 26b**). It has been found that decreasing the size of the nano-phases below one-tenth of the light wavelength (e.g., zinc sulfide nanoparticles with 3 nm diameter) can effectively diminish light scattering to achieve hydrogels with a high refractive index (i.e., 1.49) and high transparency (**Figures 26b-d**)⁸⁴⁴. Overall, the design principle for hydrogels with high refractive indices and transparency is to uniformly embed high-refractive-index non-scattering nano-phases in hydrogel matrices.



Figure 26. Design principle for hydrogels with high reflective indices and transparency: uniformly embed high-refractive-index non-scattering nano-phases. a. high contrast between reflective indices of the hydrogel fiber η_{HF} and tissue fluid η_{TF} can give minimal light leakage. b. uniformly embedding nano-phases such as nano-particles with high refractive indices in the hydrogel matrices can enhance the refractive index of the hydrogel. The size of the nano-phases d_{NC} should be much smaller than the light wavelength λ for minimal scattering and high transparency. c. hydrogels with high reflective indices and transparency can be used as optical fibers in living tissues. d. photo of a hydrogel optical fiber⁸⁵⁹. d is reproduced with permission from Ref⁸⁵⁹. Copyright 2015 Wiley.

6.4. Tunable acoustic impedance: tune densities and bulk moduli of effectively homogeneous hydrogels

Hydrogels have been widely used as the media for sound-wave transmissions such as the coupling agents for imaging and therapeutic ultrasounds. It is highly desirable to design hydrogels that possess tunable acoustic impedance to match the impedance of different materials or varying environments^{26,44}. The acoustic impedance *z* of a homogenous material can be expressed as

$$Z = \sqrt{\rho_{eff} K_{eff}} \tag{34}$$

where ρ_{eff} and K_{eff} are the effective density and bulk modulus of the material, respectively. Because the density and bulk modulus of common hydrogels are almost the same as those of water, the acoustic impedance of common hydrogels also approximates that of water. To achieve tunable acoustic impedance, fluidic channels have been patterned into tough hydrogel matrices recently (**Figure 27a**)⁴⁴. By infusing air, water or liquid metal (i.e., eutectic gallium-indium) into the fluidic channels, the effective density, bulk modulus and thus acoustic impedance of the hydrogel can be dramatically varied to approximate the acoustic impedance of air, water and many solids on demand (**Figure 27b**)⁴⁴. In order to approximate a homogeneous material, the fluidic channels should be uniformly distributed in the hydrogel and the characteristic sizes of the fluidic channels (i.e., channel diameter and distance between adjacent channels) should be much smaller than the acoustic wavelengths. Overall, a generic design principle for hydrogels with tunable acoustic impedance is to tune densities and bulk moduli of effectively homogeneous hydrogels.



Figure 27. Design principle for hydrogels with tunable acoustic impedance: tune densities and bulk moduli of effectively homogeneous hydrogels. a. by infusing air, water or liquid metal (i.e., eutectic gallium-indium) into the fluidic channels inside a hydrogel matrix, the effective density, bulk modulus and thus acoustic impedance of the hydrogel can be dramatically varied. b. the hydrogel can approximate the acoustic impedance of air, water and many solids on demand⁴⁴. **a** and **b** are reproduced with permission from Ref⁴⁴. Copyright 2019 Wiley.

6.5. Self-healing: form new crosslinks and/or polymers at damaged regions

A salient feature of many biological hydrogels is their capability of healing after injury. The capability of self-healing can potentially bestow synthetic hydrogels with merits such as damage mitigation and long-term robustness. However, the healing processes in biological hydrogels mostly rely on the functions of biological cells, which usually do not exist in synthetic hydrogels. In absence of living components, a generic strategy to achieve self-healing in engineering materials is to form new materials and/or interactions in the vicinity of damaged regions in the materials⁸⁶⁰.

In particular, for soft materials such as elastomers and hydrogels, the new materials formed in the vicinity of damaged regions are usually new crosslinks and/or polymer chains. Therefore, the design principle for self-healing hydrogels is to form new crosslinks and/or polymers at damaged regions (**Figure 28a**). The commonly-used crosslinks for self-healing of hydrogels include weak physical crosslinks such as hydrogen bonds^{750,765,767,861}, ionic bonds^{305,496,862}, metal coordinations^{305,767,852}, hydrophobic interactions⁸⁶³ and guest-host interactions⁷⁵²; and dynamic covalent bonds such as olefin metathesis^{46,864}. Once two newly-formed surfaces in a damaged hydrogel are brought into contact with each other under certain conditions such as specific temperature and pH, new crosslinks can form on the interface, endowing the hydrogel with the self-healing capability (**Figures 28a** and **c**). Besides weak physical and dynamic covalent crosslinks (i.e., reversible crosslinks), the self-healing of hydrogels can also be achieved through interdiffusion of polymer chains to form entangled chains that span the crack surfaces (**Figures 28a** and **c**)^{865,866}. In fact, self-healing processes in hydrogels mostly involve both mechanisms of chain entanglement and reversible crosslinking⁸⁶⁷.

Notably, when the surfaces of two intact self-healing hydrogels are brought in contact under certain conditions such as specific temperature and pH, an adhered interface can also be formed between the two hydrogels. Therefore, strictly speaking, most of the existing self-healing hydrogels are self-adhesive hydrogels, because the damage of the hydrogels is not required to induce the process of self-healing. Recently, Matsuda et al. have reported a self-growing or self-reinforcing hydrogel, in which the scission of polymer chains can induce mechanoradicals that trigger the polymerization of monomers in the solvent of the hydrogel (**Figure 28b**)⁸⁶⁸. Consequently, the hydrogel self-grows or self-reinforces after moderate damage, analogous to mechanical training of a muscle (**Figure 28d**). This strategy may be adopted for the future design of truly self-healing (instead of self-adhesive) hydrogels where the healing is triggered by the damage. For a further detailed discussion on various types of self-healing hydrogels, a recent review paper is recommended⁸⁴⁶.



Figure 28. Design principle for self-healing hydrogels: form crosslinks and/or polymers at damaged regions. a. reversible crosslinks and polymer chain entanglements form on the interfaces between two pieces of hydrogels for self-healing or self-adhesion. **b.** damage of a hydrogel induces new polymerization and crosslinking, giving self-reinforcement or self-growth of the hydrogel⁸⁶⁸. **c.** photos of a self-healing hydrogel based on oppositely charged polyelectrolytes^{496,836}. **d.** photos of a self-reinforcing or self-growing hydrogel⁸⁶⁸. **b** and **d** are reproduced with permission from Ref⁸⁶⁸. Copyright 2019 American Association for the Advancement of Science. **c** is reproduced with permission from Ref⁸³⁶. Copyright 2013 Springer Nature.

6.6. Implementation strategies for extreme physical properties with unconventional polymer networks

While the implementation strategies for hydrogels with extreme mechanical properties exploits various types of UPNs as discussed in **Section 5**, it seems hydrogels with extreme physical properties mainly reply on one implementation strategy: functional nano-/micro-/macro-fillers. The functional fillers range from percolated conductive phases for high electrical conductivity, to

magnetic particles for magnetization, to high-refractive-index non-scattering nano-phases for high refractive index and transparency, to fillers with tunable densities and bulk moduli for tunable acoustic impedance, and to reversible crosslinks and damage-triggered polymerization for self-healing capability.

7. Design of hydrogels with multiple combined properties

In addition to the extreme mechanical and physical properties discussed in **Sections 5** and **6**, respectively, chemical and biological properties of hydrogels also play critical roles in various applications of hydrogels. In fact, many nascent applications of hydrogels such as hydrogel living devices commonly require that a set of combined mechanical, physical, chemical and biological properties simultaneously coexist in hydrogels^{20,24,46}. In order to achieve multiple combined properties, we will propose a general strategy for the orthogonal design of hydrogels guided by the corresponding design principles, which will then be implemented with UPNs in a synergistic manner.

7.1. Orthogonal design principles

In **Sections 5** and **6**, we have discussed the design principles for hydrogels to achieve a variety of extreme mechanical and physical properties, which are summarized as the following.

- Tough: build dissipation into stretchy polymer networks.
- Strong: synchronize chain stiffening and fracture.
- Resilient: delay dissipation.
- Tough adhesion: integrate tough dissipative hydrogels and strong interfacial linkages.
- Fatigue-resistant: pin fatigue cracks with intrinsically high-energy phases.
- Fatigue-resistant adhesion: strongly bond intrinsically high-energy phases on interfaces.
- Electrically conductive: percolate electrically conductive phases.
- Magnetization: embed magnetic particles and pattern ferromagnetic domains.
- High reflective index and transparency: uniformly embed high-refractive-index non-scattering nano-phases.
- Tunable acoustic impedance: tune densities and bulk moduli of effectively homogeneous hydrogels.
- Self-healing: form crosslinks and/or polymers at damaged regions.

Since the abovementioned design principles are general and material-independent, they have been widely deployed for the design of biological hydrogels, synthetic hydrogels, and other engineering materials. In addition, based on the discussions in **Sections 5** and **6**, these design principles do not contradict or exclude one another in general. For example, the seeming

contradiction between high toughness and high resilience of hydrogels has been reconciled by the design principles of building dissipation into stretchy polymer networks and delaying dissipation, respectively. Therefore, the design of multiple combined mechanical and physical properties of hydrogels can potentially follow the corresponding design principles in an orthogonal and independent manner as illustrated in **Figure 29a**. For example, a tough, electrically conductive, and self-healing hydrogel can be potentially designed by following the orthogonal design principles of building dissipation into stretchy polymer networks, percolating electrically conductive phases, and forming crosslinks and/or polymers at damaged regions, respectively³² (**Figure 29b**). The hydrogel can further form tough adhesion on substrates by following the design principle of integrating tough hydrogels and strong interfacial linkages⁸⁵¹.

In addition, although chemical and biological properties of hydrogels are beyond the scope of the current review, it is expected that the design of hydrogels' chemical and biological properties will likely follow a set of design principles that are orthogonal with one another and with the design principles for mechanical and physical properties as well. Consequently, a set of orthogonal design principles will potentially guide the rational design of future hydrogels that possess multiple combined mechanical, physical, chemical and biological properties (**Figure 29**).

7.2. Synergistic implementation strategies

The orthogonal design principles for hydrogels to achieve multiple combined properties will be implemented with UPNs in a synergistic manner, meaning that one type of UPN can implement multiple design principles. As discussed in **Section 4**, the commonly-used UPNs include:

- Ideal polymer networks
- Polymer networks with slidable crosslinks
- Interpenetrating and semi-interpenetrating polymer networks
- Polymer networks with high-functionality crosslinks
- Nano-/micro-fibrous polymer networks and the UPN interactions:
- Strong physical crosslinks
- Weak physical crosslinks
- Dynamic covalent crosslinks.

Each UPN architecture or interaction usually can implement (or facilitate the implementation of) multiple design principles. For example, the nano-/micro-fibrous polymer networks can integrate high stretchability and mechanical dissipation (Section 5.1), delay dissipation (Section 5.3), synchronize stiffening and fracture of polymer chains (Section 5.2), and act as intrinsically high-energy phases (Section 5.5) to implement the design principles for tough, resilient, strong, and fatigue-resistant hydrogels, respectively. By strongly bonding the nano-/micro-fibrous of substrates (Section 5.4 and Section 5.6), the corresponding nano-/micro-fibrous

polymer networks can achieve tough and fatigue-resistant adhesion as well. Furthermore, the nano-/micro-fibers can also be made functional such as electrically conductive or with high reflective index (Section 6.1) to implement the design of the corresponding extreme physical properties. Not surprisingly, biological hydrogels indeed frequently employ nano-/micro-fibrous polymer networks, supplemented by other UPN architectures and interactions, to achieve multiple combined extreme mechanical and physical properties necessary for their robustness and wellbeing over the lifetime (Figure 2).

Last but not least, it should be emphasized that the design principles and implementation strategies for hydrogels discussed in this paper are based on generic polymer networks; therefore, they should be applicable to other soft materials comprised of polymer networks including elastomers and organogels as well. For example, the design principle and implementation strategy for tough hydrogels have been used to design tough elastomers⁷³⁷. We expect the current review will provide a solid and systematic foundation for the rational design of various types of polymeric soft materials including hydrogels, elastomers and organogels to achieve multiple combined extreme properties for diverse applications. Furthermore, we hope the current review will provoke interdisciplinary discussions on a fundamental question: why does nature select soft materials, especially hydrogels embodied in unconventional polymer networks (**Figures 1** and **2**), to constitute the major components of animal bodies?



Figure 29. Orthogonal design principles and synergistic implementation strategies for the design of hydrogels with multiple combined extreme properties. a. schematics of the orthogonal design principles and the synergistic implementation strategies. b. example of the design of a tough, self-healing and electrically conductive hydrogel.

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