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1.1 Introduction

In-situ bioelectronics, a rapidly evolving field focusing on the development of electronic devices that can operate within the body for on-site sensing, stimulation, and powering, holds great promise for revolutionizing the field of medicine in a variety of ways (Figure 1.1a) [1, 2]. For example, the development of biosensors that can achieve on-site quantification of biomarkers closely related to the development and progression of colorectal cancer (CRC) would enable early CRC detection preventing CRC from progressing, thereby increasing the five-year relative survival rate up to 90% [3]. The development of neural probes that can form intimate interfaces with neurons without provoking a severe foreign body response would enable chronic neuron stimulation and recording, facilitating fundamental understanding of neural activities and offering long-term treatment for neuropsychiatric disorders, traumatic injuries, and inflammatory conditions [4-6]. The advancement of energy harvesting devices that can provide sustainable power supply to cardiac pacemakers could prolong their lifespan and help maintain or restore a normal heart rhythm with electrical impulses [7–9]. However, the key challenge faced by *in-situ* bioelectronics stems from the fundamentally contradictory properties between electronic components and biological systems, which induce foreign body responses due to mechanical, chemical, and biological mismatches. Specifically, electronic components are typically made of metals, silicon, glass, ceramics, and plastics that are hard, dry, and abiotic; in contrast, biological systems are composed of living tissues that are soft, wet, and dynamic.

Hydrogels, as polymer networks infiltrated with water, exhibit intriguing multiphysics phenomena associated with mechanical, electrical, chemical, and thermal

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Figure 1.1 Overview of the *in-situ* hydrogel electronics. (a) *In-situ* bioelectronics revolutionized the field of medicine in a variety of ways. (b) Hydrogels form long-term, high-efficacy, multi-modal bridging interfaces between electronic components and biological systems. (c) Three key components of *in-situ* hydrogel bioelectronics include stretchable hydrogel conductors, electrochemical hydrogel biosensors, and flexible hydrogel biobatteries.

couplings [10, 11]. Hydrogels are typically constituted of two phases: one phase of interconnected polymer networks giving the solid-like network elasticity, and the other phase of infiltrated water molecules endowing the fluid-like transport property [12–15]. Due to the unique combination of polymer networks and water molecules, hydrogels show their superior softness, wetness, responsiveness, biocompatibility, and bioactivity, therefore having been regarded as an ideal material candidate to form long-term, high-efficacy, multi-modal bridging interfaces between electronic components and biological systems (Figure 1.1b) [16, 17].

Recently, a nascent field named hydrogel bioelectronics has rapidly evolved, exploiting hydrogels as key components for electronic devices that seamlessly interact with biological systems. The generic idea for the design of hydrogel bioelectronic device is to embed functional electronic components such as conductors, microchips, transducers, resistors, and capacitors inside or attached to the surface of a highly stretchable and tough hydrogel matrix, providing a soft, wet, and biocompatible environment interfacing with biological tissues [18]. As the hydrogel bioelectronic device is stretched, flexible electronic components can deform together with the device while rigid electronic components maintain their undeformed shapes, which involves large deformation of hydrogels around rigid electronic components [19, 20]. Therefore, to maintain reliability and functionality of the device, the hydrogel matrix and the interface between hydrogel and functional components need to be tough and robust. Pioneered by Gong et al., hydrogels with fracture toughness higher than 10^4 J/m² have been widely available [21–23]; additionally, initiated in the past 10 years, hydrogel adhesions to diverse substrates can successfully achieve interfacial fracture toughness above 10^3 J/m², on the same order as their biological counterparts [24–29]. The recent development of tough hydrogels and tough hydrogel adhesions as well as their synthesis and fabrication techniques have enabled various soft material technologies in the form of diverse hydrogel bioelectronics [18, 30, 31].

More recently, an emerging class of *in-situ* bioelectronics that combines hydrogel technologies with electronic components to create devices that can interact with harsh environments within the body, which we define as *in-situ hydrogel bioelectronics*, have been recently proposed with great promise in potentially addressing the limitations faced by existing *in-situ* bioelectronics [16, 31, 32]. Despite the promise, the complicated chemical, biological, and mechanical factors in the physiological environment pose significant challenges to the reliability and functionality of hydrogel electronic devices when operating within the body. What are the key properties of hydrogels and how to rationally design these properties for developing electronic devices that can operate in the physiological environment for reliable on-site sensing, stimulation, and powering? These are unanswered questions, even considering a growing number of reviews on hydrogel bioelectronics [16, 33–37].

This chapter aims to provide an overview of the design principles, implementation mechanisms, and manufacturing/fabrication techniques, particularly centering on extreme mechanics of hydrogels, for developing three key components of *in-situ* hydrogel bioelectronics (Figure 1.1c): (i) stretchable hydrogel conductors, (ii) electrochemical hydrogel biosensors, and (iii) flexible hydrogel biobatteries. The chapter is organized as follows. Section 1.2 will discuss polymer mechanics for rationally designing extreme mechanical and physical properties of hydrogels crucial for the development of in-situ hydrogel bioelectronics. Section 1.3 will discuss the multiscale orthogonal design of stretchable hydrogel conductors and strategies for implementing the orthogonal design. Section 1.4 will discuss the principles for achieving high-specificity and high-sensitivity of electrochemical hydrogel biosensors, specifically focusing on selective transport design in hydrogels and electrochemical sensing performance at hydrogel-electrode interfaces. Section 1.5 will briefly review the recent efforts in developing flexible hydrogel biobatteries by harvesting mechanical energy, chemical energy, and thermal energy. Section 1.6 will conclude the chapter with a set of future opportunities by integrating interdisciplinary efforts in ingestible sensors, neural interfaces, miniature robots, and data analytics.

1.2 Extreme Properties of Hydrogels by Polymer Network Design

Due to the unique combination of solid-like polymer networks and fluid-like water molecules, hydrogels exhibit superior softness, wetness, responsiveness,



Figure 1.2 Extreme properties of hydrogels by polymer network design. (a) Schematics of unconventional polymer networks such as ideal polymer networks, interpenetrating polymer networks, semi-crystalline polymer networks, micro-/nanofibrous polymer networks. (b) Schematics of mechanical and physical properties critical for the development of *in-situ* hydrogel bioelectronics, including elasticity, fracture, fatigue, and mass transport.

biocompatibility, and bioactivity, therefore being exploited as key material candidates for developing *in-situ* hydrogel bioelectronics. These properties have been intensively studied by understanding the nonlinear elasticity [38], swelling [39–43], poroelasticity [44, 45], viscoelasticity [46–50], fracture [51–53], and fatigue [54–57] of hydrogels, following the pioneering work in the field of polymers and soft materials [58–69]. Despite these unique mechanical and physical properties of common hydrogels, the development of *in-situ* hydrogel bioelectronics also requires hydrogels to possess extreme mechanical and physical properties, such as tunable elastic modulus, extremely high values of fracture toughness and fatigue threshold, and tunable molecular transport. In this section, we will discuss the polymer mechanics to rationally push the limits of mechanical and physical properties of hydrogels including elastic modulus, fracture toughness, fatigue threshold, and mass transport that are crucial for the development of *in-situ* hydrogel bioelectronics (Figure 1.2).

1.2.1 Elastic Modulus

Elastic modulus is one of the most important properties of hydrogels used in biomedical applications, as it governs their ability to withstand deformations while maintaining compliance without damaging the surrounding soft tissues or causing adverse foreign body responses. There are several methods available to measure the elastic modulus of hydrogels, including tensile/compression, rheology, and indentation tests, as illustrated in Figure 1.3a. The shear elastic modulus of a hydrogel *G* can be measured by identifying the initial slope of the stress–strain curve in tensile or compression tests *E*, namely $G = \alpha E$, where α is a dimensionless perfector

1.2 Extreme Properties of Hydrogels by Polymer Network Design 7



Figure 1.3 Common experimental methods to characterize mechanical and physical properties of hydrogels. (a) Tensile, rheology, and indentation tests to measure the elastic modulus of hydrogels. (b) Pure shear, T-peel, and tearing tests to measure the fracture toughness of hydrogels. (c) Pure shear fatigue tests to measure the fatigue threshold of hydrogels Γ_0 by plotting the crack extension da/dN versus the applied energy release rate G. (d) 1D transport assay, fluorescence recovery after photobleaching (FRAP), and particle tracking methods to measure the diffusivity of particles (e.g. ions, monomers, proteins, viruses) in hydrogels.

depending on samples' dimensions and material incompressibility [70]. Rheology test is the other method to measure the shear elastic modulus of a hydrogel, which is often preferred for soft hydrogels. Rheology tests can provide information on both storage and loss modulus across a range of frequencies, allowing for a quantitative decomposition of elastic and viscous contributions during deformation [71]. Indentation tests can also be used to characterize the shear elastic modulus, but the results may be affected by water migration and stress state. Therefore, a theoretical analysis is necessary to accurately interpret the indentation results and decouple nonlinear elasticity, viscoelasticity, and poroelasticity in hydrogels [72].

Assuming the polymer network of a hydrogel is fully amorphous with negligible molecular entanglements and crystalline domains, the shear elastic modulus of a hydrogel *G* can be theoretically predicted following the classical affine and phantom network theories of elasticity [12, 66],

$$\frac{G}{kT} = C\phi^{1/3}n\tag{1.1}$$

where ϕ is the volume fraction of dry polymers, *n* is the number of elastically active polymer chains per unit volume of dry polymers, *k* is the Boltzmann constant, *T* is the absolute temperature, and *C* is a constant that has a value of 1 when the polymer network of the hydrogel follows affine deformation and $1 - \frac{2}{f}$ when the polymer network of the hydrogel follows phantom deformation with *f* being the functionality of the polymer network (i.e. the number of chains connected to a junction point). By substituting the typical values of C = 1, $\phi = 0.1$, $n = 10^{24} - 10^{26}$ m⁻³, and $kT = 4.11 \times 10^{-21}$ J, the shear elastic modulus of a hydrogel is estimated on the order of 1–100 kPa.

Recent experiments have shown that the measured shear modulus of a hydrogel is consistently below the theoretical predictions by Eq. (1.1) due to the presence of molecular defects in a real hydrogel. Given the capability of counting the numbers

of various orders of molecular defects in hydrogels, Olsen and Johnson and coworkers [73] developed a real elastic network theory to quantify the impact of molecular defects on shear elastic modulus,

$$\frac{G}{kT} = \frac{f-2}{f} \sum_{i} \varepsilon_{i} n_{i}$$
(1.2)

where n_i is the number of the type *i* polymer chains per unit volume of the hydrogel associated with a specific type of molecular defects, ε_i is the elastic effectiveness of the type *i* polymer chains. ε_i accounts for the elastic contribution of each polymer chain, having a value of 1 if the polymer chain is an ideal chain with no impact from defects and a value smaller than 1 if the polymer chain is a defective chain affected by surrounding molecular defects. The real elastic network theory suggests the critical role of polymer network architecture on hydrogel's elastic modulus. The design principle for achieving tunable elastic modulus of hydrogels is to engineer the type and density of molecular defects, ubiquitous in synthetic hydrogels and biological tissues [74] (Figure 1.4a).

1.2.2 Fracture Toughness

Fracture toughness, the energy required to propagate a unit area of crack surface under monotonic load, defines the ability of a material to resist crack extension under stress. Fracture toughness of a hydrogel is crucial for maintaining reliability of in-situ hydrogel bioelectronics [10, 11]. As illustrated in Figure 1.3b, the fracture toughness of a hydrogel can be measured through pure shear, T-peel, and tearing tests, originally proposed by Rivlin and Thomas for measuring the fracture toughness of rubbers [75]. In a pure shear test, the fracture toughness of a hydrogel Γ is determined by the critical energy release rate applied on a notched sample G_c , $\Gamma = G_{c} = W(\lambda_{c})H$, where W is the strain energy density stored in the unnotched sample with λ_c being the critical stretch for crack extension in the notched sample, and H is the sample height [53]. T-peel tests can also be used to measure the fracture toughness of hydrogels [76], where a hydrogel layer with a precut is sandwiched between two inextensible backing layers. The fracture toughness of the hydrogel is then calculated via $\Gamma = 2F_{SS}/b$, where F_{SS} is the steady-state plateau force, and b is the specimen's thickness. Tearing tests, also known as trouser tests, are another commonly adopted method for measuring hydrogel toughness [77]. Unlike pure shear and T-peel tests with the crack deformed in Mode I, the crack in a tearing test is deformed in Mode III (out-of-plane shear loading). The fracture toughness of a hydrogel in a tearing test is determined by $\Gamma = 2F_{SS}/b$, where F_{SS} is the steady-state plateau force, and b is the specimen's thickness.

Common hydrogels are intrinsically brittle [78]. The intrinsic fracture energy of a hydrogel Γ_0 can be calculated following the classical Lake–Thomas theory [12, 61, 79],

$$\Gamma_0 = \phi^{2/3} \cdot n\sqrt{Nb} \cdot NU = \phi^{2/3} nbN^{3/2}U$$
(1.3)

where ϕ is the volume fraction of dry polymers, $n\sqrt{Nb}$ is the number of elastically active polymer chains per unit area of crack surface, with *n* being the number of



Design principle: Introducing delocalized damage around crack tip by fracturing high-energy phases

Figure 1.4 Design principles for extreme properties of hydrogels. (a) Achieving tunable elastic modulus by engineering type and density of molecular defects in hydrogels. (b) Achieving high fracture toughness of hydrogels by introducing both bulk hysteretic dissipation and near-crack dissipation in stretchy polymer networks. (c) Achieving high fatigue threshold of hydrogels by introducing high-energy phases to impinge fatigue crack extension. (d) Achieving tunable diffusivity in hydrogels by leveraging the synergy of reversible bonds and network elasticity.

elastically active polymer chains per unit volume of dry polymers, *N* being the number of Kuhn monomers in each polymer chain, *b* being the length of each Kuhn monomer, *NU* is the energy required to fracture a polymer chain, with *U* being the energy required to fracture a single Kuhn monomer. It is commonly assumed that the dry polymers of a hydrogel satisfy the volume conservation following $Nnb^3 = 1$ with b^3 being the volume of a Kuhn monomer. By imposing the volume conservation, the

intrinsic fracture energy of a hydrogel can be calculated via,

$$\Gamma_0 = \phi^{2/3} b^{-2} N^{1/2} U \tag{1.4}$$

By substituting the typical values of $\phi = 0.1$, $b = 10^{-9}$ m, N = 100 - 10, 000, and U = 100kT = 4.11×10^{-19} J, the intrinsic fracture energy of a hydrogel Γ_0 is estimated as low as 1-10 J/m², orders of magnitude lower than biological tissues around 1000 J/m².

Lin and Zhao recently developed a defect-network fracture model to predict the intrinsic fracture energy of polymer networks containing various types of topological defects including cyclic loops and dangling chains [80]. The defect-network fracture model is inspired by the real elastic network model [73] or, more generally, the phantom network model [66, 81] discussed in Section 1.2.1. Analogous to the previous elastic models, the key idea of this fracture model is to introduce effectively longer fractured chains to account for the energy of the fractured chains on the crack path. However, physically different from the previous elastic models, the effectively longer chains in the fracture model do not change the density of the layer of fractured chains. The intrinsic fracture energy of the polymer network with defects normalized by that of the corresponding defect-free ideal network can be expressed as:

$$\overline{\Gamma} = \sum_{X} (\gamma_X - 1) n_X^{\text{affected}} - \sum_{X} n_X^{\text{inactive}} + 1$$
(1.5)

where $\gamma_X \ge 1$ is the fracture effectiveness to account for the contribution to intrinsic fracture toughness by a single polymer chain affected by defect *X*, n_X^{affected} is the number density of affected chains due to the presence of defect *X*, n_X^{inactive} is the number density of inactive chains due to the presence of defect *X*. The defect-network fracture model indicates a competing effect due to the presence of topological defects: toughening by increasing effective chain length and weakening by introducing inactive chains. While the defect-network fracture model predicts that the presence of defects can potentially enhance intrinsic fracture energy of hydrogels by a few times, such enhancement is not sufficient to ensure the reliable use of hydrogels in engineering applications.

In the past 20 years, the fracture toughness of hydrogels has been significantly enhanced above 1000 J/m², making tough hydrogels key load-bearing components for devices and machines [82, 83]. The generic toughening mechanism of hydrogels is to incorporate two physical processes: one is the scission of a layer of polymer chains on the crack tip, and the other is the bulk hysteretic dissipation around the crack tip such as Mullins effect and viscoelasticity [22, 23, 51, 78, 84–91]. The first process defines the intrinsic fracture energy Γ_0 as discussed in Eqs. (1.3)–(1.5), and the second process defines the bulk hysteretic dissipation's contribution Γ_D^{bulk} . Conceptually, the total fracture toughness of a tough hydrogel can be expressed as:

$$\Gamma = \Gamma_0 + \Gamma_D^{\text{bulk}} \tag{1.6}$$

which is often named the bulk dissipation model. The value of $\Gamma_{\rm D}^{\rm bulk}$ can be estimated by $\Gamma_{\rm D}^{\rm bulk} = U_{\rm D}L_{\rm D}$ with $U_{\rm D}$ being the energy for breaking sacrificial bonds per unit volume of the material and $L_{\rm D}$ being the length of the process zone around crack tip where breaking sacrificial bonds occurs. $U_{\rm D}$ can be estimated by the bond

energy of one sacrificial bond times the number of sacrificial bonds per unit volume of the material, on the order of 10^6 J/m^3 . Since the value of L_D can reach 1 mm [92], the value of Γ_D^{bulk} can reach 1000 J/m², with the orders of magnitude larger than Γ_0 .

The bulk dissipation model has been widely used to qualitatively explain the toughening mechanisms in diverse soft tough materials [21, 23, 93–95], but recent study shows that the bulk dissipation model significantly underestimates the toughness enhancement of tough hydrogels [93]. The missing term is attributed to a near-crack dissipation that does not rely on bulk hysteresis [96–98]. To account for both bulk hysteretic dissipation and near-crack dissipation in soft tough materials, an extreme toughening model was recently proposed, indicating that the total fracture toughness of a tough hydrogel exhibiting both bulk hysteretic dissipation and near-crack dissipation and near-crack dissipation and near-crack dissipation at the total fracture toughness of a tough hydrogel exhibiting both bulk hysteretic dissipation and near-crack dissipation can be expressed as:

$$\Gamma = \Gamma_0 + \Gamma_D^{\text{bulk}} + \Gamma_D^{\text{tip}} \tag{1.7}$$

where Γ_0 is the intrinsic fracture energy, Γ_D^{bulk} is the bulk hysteretic dissipation's contribution to fracture toughness, and Γ_D^{tip} is the near-crack dissipation's contribution to fracture toughness. A governing equation for the extreme toughening model can be further derived as:

$$\frac{\Gamma}{\Gamma_0} = \frac{\beta}{1 - \alpha h_{\rm m}} \tag{1.8}$$

where $\beta = (\Gamma_0 + \Gamma_D^{\text{tip}})/\Gamma_0 \ge 1$ is a dimensionless number to account for the near-crack dissipation due to molecular entanglements, $0 \le \alpha \le 1$ is a dimensionless number depending on the stretch-dependent hysteresis of the bulk materials ($\alpha = 1$ for highly stretchable materials), and $0 \le h_m < 1$ is the maximum stress-stretch hysteresis of the bulk material. While the cause of the near-crack dissipation is not fully understood, the reported experiments have shown the potential for achieving high values of β up to 10, suggesting the crucial toughening role by the near-crack dissipation. To summarize, the design principle for achieving high fracture toughness in hydrogels is to introduce both bulk hysteretic dissipation (e.g. Mullins effect and viscoelasticity) and near-crack dissipation (e.g. molecular entanglements) in stretchy polymer networks, as depicted in Figure 1.4b.

1.2.3 Fatigue Threshold

Fatigue threshold, the energy required to propagate a unit area of crack surface under cyclic load, defines the ability of a material to resist fatigue crack extension under stress [55, 99, 100]. Fatigue threshold of hydrogels is critical for achieving longevity of *in-situ* hydrogel bioelectronics. As illustrated in Figure 1.3c, the measurement of fatigue threshold of hydrogels requires cyclic loading of unnotched and notched hydrogel specimens. The fatigue tests for hydrogels are typically performed in a chamber with controlled humidity or water bath to ensure the sample reaches an equilibrium state [54, 56]. By monitoring the crack extension Δa at a controlled

energy release rate *G*, one can identify a critical energy release rate at the intersection of abscissa axis as the measured fatigue threshold Γ_0 .

While hydrogels have been made tough with high toughness above 1000 J/m^2 , as discussed in Section 1.2.2, these tough hydrogels still suffer from fatigue fracture when subjected to prolonged cyclic loading [54, 79, 97, 101]. The experimental findings conclude that the resistance to fatigue crack propagation after prolonged cycles of loads is the energy required to fracture a single layer of polymer chains (i.e. the intrinsic fracture energy of the hydrogel), which is unaffected by the additional dissipation mechanisms introduced in tough hydrogels [56]. To address the challenge of fatigue failures in conventional tough hydrogels, we and others have proposed a general design principle for fatigue-resistant hydrogels (Figure 1.4c) – inducing delocalized damage around the crack tip by fracturing high-energy phases, such as nanocrystals [28, 56, 100, 102–104], micro–/nanofibers [105], and macro-fibers [92, 106] in hydrogels. Additionally, hierarchical molecular structure design such as introducing bi-continuous phase networks can suppress fatigue-induced crack advance [57, 107].

The fatigue threshold of fatigue-resistant hydrogels containing high-energy phases can be qualitatively calculated by modifying the Lake–Thomas theory:

$$\Gamma_0 = \phi^{2/3} n l_d N U \tag{1.9}$$

where ϕ is the volume fraction of dry polymers, n is the number of elastically active polymer chains per unit volume of dry polymers, NU is the energy required to fracture a polymer chain with N being the number of Kuhn monomers in each polymer, U being the energy required to fracture a single Kuhn monomer, and l_d is the length scale of the delocalized damage and understood as the crack processing zone length at the threshold. In conventional hydrogels, l_d is equal to the length of a single layer of polymer chains, i.e. $l_d = \sqrt{Nb}$; in fatigue-resistant hydrogels, l_d is larger than the length of a single layer of polymer chains, i.e. $l_d > \sqrt{Nb}$. Since the parameters ϕ , N, and U are intrinsic properties of hydrogels mainly rely on the mechanisms that can significantly enlarge the value of l_d , which can be achieved by introducing intrinsically high-energy phases above mentioned.

1.2.4 Mass Transport

Diffusion in a hydrogel, movement of particles (e.g. ions, monomers, proteins, and viruses) through the hydrogel, is a ubiquitous phenomenon in nature and a fundamental process that governs the working principles in diverse applications. For example, the distinct diffusion of different biomarkers in hydrogels determines the sensing sensitivity and sensing specificity in electrochemical hydrogel biosensors discussed in Section 1.4; the stress-induced diffusion of ions in nano channels of hydrogels governs the stress-voltage coupling in flexible hydrogel batteries discussed in Section 1.5. The diffusivity of particles in hydrogels can be commonly characterized by 1D transport assay, fluorescence recovery after bleaching (FRAP), and particle tracking methods (Figure 1.3d). The mode of diffusion in a hydrogel is determined by the mesh size of the hydrogel. When the hydrogel's mesh size is much larger than the size of the substance, the substance moves as Brownian diffusion, the diffusivity of which is governed by the viscosity of solvent (i.e. water) in the hydrogel [108]:

$$D = \frac{kT}{3\pi\eta d} \tag{1.10}$$

where *k* is the Boltzmann constant, *T* is the absolute temperature, η is the dynamic viscosity of water, and *d* is the diameter of the substance. By substituting the typical values of $kT = 4.21 \times 10^{-21}$ J, $\eta = 8.9 \times 10^{-4}$ Pa · s, and d = 1 nm, the diffusivity of the substance in a hydrogel with large mesh size is estimated as 10^{-10} m/s². When the hydrogel's mesh size is on the same order as the size of the substance, the diffusivity of the substance in a hydrogel is governed by the hydrogel's mesh size ξ [109]:

$$\frac{D}{D_0} \sim \exp\left(-\frac{d}{\xi}\right) \tag{1.11}$$

where D_0 is the diffusivity of the substance in water. For a hydrogel with its mesh size two times the substance diameter (i.e. $d/\xi = 2$), the diffusivity of the substance reduces by 10 times on the order of 10^{-11} m/s^2 .

Existing efforts have been mostly focused on Brownian diffusion of particles in a fluid, the diffusivity of which is governed by the viscosity of fluids. In contrast, when particles diffuse in the polymer networks of a hydrogel, the transport of particles is a discrete process of particles making stochastic hops between neighboring sites, namely hopping diffusion, the diffusivity of which is governed by the energy required to overcome the elasticity of polymer networks [110]. Such network elasticity can be readily tuned by mechanical deformation applied on the hydrogel. This implies the potential of harnessing mechanical deformation as a new design space to program particle transport in hydrogels. Specifically, for a particle with size *d* confined in a cage formed by polymer chains with mesh size ξ slightly smaller than particle size (i.e. $\xi < d$), the particle can still escape from one cage to the other neighboring cage by overcoming the energy barrier due to network elasticity. The probability for such escape *P* is determined by the energy barrier E_e via $P = \int_d^{\infty} \frac{e^{-E_c/kT}}{\xi} dx$ where *k* is the Boltzmann constant and *T* is the absolute temperature, resulting in a reduced hopping diffusivity [110],

$$D_{\text{network}} = D_0 P \tag{1.12}$$

where D_0 is the particle diffusivity in a hydrogel free of network elasticity. The fundamental understanding of how mechanical deformation modulates particle transport in soft materials remains an unexplored research topic but will potentially lead to multiple previously inaccessible technologies that rely on mechano-transport design in hydrogels, including but not limited to force-sensitive cargo for on-demand drug delivery, and strain-programmable tissue adhesive for prolonged tissue repair.

In addition to network elasticity, hopping diffusion in hydrogels can also be induced by reversible interactions between particles and functional groups in hydrogels [111]. During a hopping event, multipoint particle attachment results in caging. As one or more of the attached points can be released, the particle can escape from one cage to the other neighboring cage, resulting in hopping

diffusivity,

$$D_{\text{reversible}} = \sum_{j=2}^{N} k_{\text{off}} P_j D_0.$$
(1.13)

where P_j is the probability of a particle maintaining *j* number of bonding sites, *N* is the number of bonding sites on the particle, D_0 is the particle diffusivity with one bonding site (N = 1), and k_{off} is the dissociate rate of the reversible interaction. Harnessing hopping diffusion by reversible interactions enables selective transport of chemical and biological cargoes based on reversible cargo-barrier interactions [112, 113]. Overall, the principle to tune particle diffusivity in hydrogels is to leverage the synergy of network elasticity and reversible bonds (Figure 1.4d).

1.3 Stretchable Hydrogel Conductors

Hydrogel-based electronic materials that can conduct electricity while being able to stretch and deform without fracture, known as stretchable hydrogel conductors, hold great potential as an alternative to traditional metallic conductors in the development of *in-situ* hydrogel bioelectronics due to their unique combination of tissue-like properties and electrical conductivity [18, 114]. Despite the promising potential, existing stretchable hydrogel conductors face a technical challenge in reconciling superior mechanical toughness and high electrical conductivity, limiting their use in *in-situ* hydrogel bioelectronics that require both mechanical and electrical properties [115–117]. Currently, most stretchable hydrogel conductors consist of a mixture of electrical conductivity due to low connectivity between electrical phases in the hydrogel matrices. Although increasing the amount of conductive filler can improve the electrical conductivity by forming a percolation network, this approach significantly compromises the hydrogel's mechanical properties by reducing stretchability and fracture toughness.

1.3.1 Multiscale Orthogonal Design

The design principle of stretchable hydrogel conductors is depicted in Figure 1.5, where the integration of electrical and mechanical phases is accomplished by a bi-continuous structure with an orthogonal design of each phase. To achieve high electrical conductivity, various electrical fillers such as ions [118, 119], conductive monomers [120, 121], conductive polymers [122–124], conductive nanoparticles [125, 126], and conductive macro fillers [127, 128] have been used. On the other hand, to achieve superior mechanical elasticity, stretchable hydrogels with interpenetrating polymer networks [21, 23, 129], hybrid crosslinkers [130–132], high-functionality crosslinkers [133–135], monodispersed polymer chains [136, 137], and meso-/macro composites [138–140] have been designed. While high electrical conductivity and superior mechanical elasticity have been



Figure 1.5 Design of stretchable hydrogel conductors relying on a bi-continuous integration of electrical and mechanical phases. (a) The high electrical conductivity in the electrical phase can be achieved by introducing electrical fillers including ions, conductive monomers, conductive polymers, conductive nanoparticles, and conductive macro fillers. (b) The superior mechanical elasticity in the mechanical phase can be achieved by designing stretchable hydrogels with interpenetrating polymer networks, hybrid crosslinkers, high-functionality crosslinkers, monodispersed polymer chains, and meso-/macro composites. Source: Reproduced with permission from Ref. [78]. Copyright 2014 The Royal Society of Chemistry.

demonstrated individually, the integration of these two phases is challenging due to their significant mechanical mismatch. The elastic modulus of the electrical phase (~1 GPa) is typically much larger than that of the mechanical phase (hydrogels, ~1–100 kPa), leading to stress concentrations at their interface and surface delamination when highly deformed. Furthermore, the fracture toughness of the electrical phase (~10 J/m²) is typically much lower than that of the mechanical phase (tough hydrogels, 1000 J/m²), resulting in microcrack formation and decreased electrical conductivity. Despite numerous individual efforts to enhance electrical conductivity and mechanical elasticity, implementing the orthogonal design through the bi-continuous integration of the electrical and mechanical phases remains a challenge.

1.3.2 Implementations of the Orthogonal Design

Figure 1.6 illustrates the typical ways in which the orthogonal design of stretchable hydrogel conductors can be implemented. A biocompatible, highly stretchable, and robust hydrogel provides a soft and wet matrix for electronic components, drastically different from dry elastomer/polymer matrices used in conventional stretchable electronics. Electronic components including stretchable, rigid, and brittle conductors are either embedded inside or attached to the surface of the hydrogel. When the stretchable hydrogel conductor is stretched, stretchable electronic components can deform together with the conductor, while rigid electronic components maintain their undeformed shapes, which requires robust interfaces between electronic components and hydrogel



Figure 1.6 General strategies to implement the orthogonal design of stretchable hydrogel conductors. (a) Electronic components including stretchable, rigid, and brittle conductors are embedded inside or attached on the surface of the hydrogel. (b) As the stretchable hydrogel conductor is stretched, stretchable electronic components can deform together with the conductor, but rigid electronic components maintain their undeformed shapes, which requires robust interfaces between electronic components and hydrogel matrix. Additionally, brittle electronic components require serpentine structural designs to reduce the energy for driving the crack formation in electronic components. Source: Reproduced with permission from Ref. [18]. Copyright 2015 John Wiley & Sons, Inc.

matrix. In addition, brittle electronic components require serpentine structural designs to minimize the energy for driving crack formation in the electronic components.

While a stretchable hydrogel conductor with intrinsically stretchable electronic components does not require special interfacial or structural design, it requires material innovation for the design of intrinsically stretchable conductive materials. Conventional conductive materials have difficulty forming long-term stable and conformable interfaces with both biological and synthetic hydrogels due to severe modulus mismatch. For example, carbon-based conductive materials like graphene have moduli of 1 TPa; metallic conductive materials like silicon have moduli of 50 GPa; even conducting polymers such as poly(3.4-ethylenedioxythiophene) polystyrene sulfonate (PEDOT: PSS), polyaniline (PANI), and polypyrrole (PPy), possessing superior flexibility compared to inorganic conductors, typically have moduli on the order of 1 GPa [141]. PEDOT: PSS-based hydrogels [13, 142], owing to their favorable compatibility, processability, and reproducibility, represent one of the most common material candidates for intrinsically stretchable electronic components. Figure 1.7a summarizes the recent efforts in reconciling electrical conductivity and mechanical stretchability in PEDOT: PSS hydrogels through interpenetrating polymer networks [142], dispersing conductive nanofillers [148], and engineering PEDOT: PSS nanofibrils [13, 149]. Liquid metal, a metal alloy including gallium, indium, and tin, is another class of intrinsically stretchable conductive materials, recently embedded in hydrogels [150, 151]. Although liquid metal shows unique properties such as high conductivity, excellent thermal stability, and ease of processing, it is also corrosive, reactive with water or oxygen, and requires safety precautions.



on interpenetrating polymer networks, dispersing conductive nanofillers, and engineering PEDOT: PSS nanofibrils. (b) The implementation of a stretchable hydrogel conductor with rigid conductors requires strong and tough interfaces between the electronic components and the hydrogel matrix such as TSP recently developed by Jin and Bao and coworkers shown in the left image (Source: Reproduced with permission from Ref. [143]. Copyright 2022 Springer PEDOT: PSS-based hydrogels shown in the left image (Source: Ref. [13]/Springer Nature/CC BY 4.0). The design of PEDOT: PSS hydrogels typically relies advanced manufacturing techniques such as micro-/nanofabrication (Source: Ref. [1441), 3D printing (Source: Ref. [145]), morphable 3D meso-structures Figure 1.7 Representative examples to implement the orthogonal design of stretchable hydrogel conductors. (a) The implementation of a stretchable hydrogel conductor with stretchable conductors requires material innovations for the design of intrinsically stretchable conductive materials such as implementation of a stretchable hydrogel conductor with brittle/rigid conductors also requires serpentine structural design, which typically relies on Nature). The design of strong and tough interfaces typically relies on covalent bonds, strong physical crosslinks, and connector polymers. (c) The (Source: Ref. [146]), and syringe-injectable mesh electronics (Source: Ref. [147]).

When incorporating rigid or nonstretchable electronic components into a stretchable hydrogel, the design of strong and tough interfaces between the electronic components and the hydrogel matrix becomes crucial. Figure 1.7b summarizes the recent efforts to develop strong interfacial linkages (e.g. covalent bonds [152, 153], strong physical crosslinks [154–156], and connector polymers [157, 158]) and introduce bulk dissipation mechanisms (such as viscoelasticity and Mullins effect) to create stretchable hydrogel conductors with rigid electronic components. For instance, Jin and Bao and coworkers recently proposed a method for designing a tough interface between a brittle semiconducting thin film and a stretchable substrate that effectively delayed microcrack formation in the film when stretched [143]. They achieved the tough interface using strong interfacial linkages through surface chemistry and dynamic bulk dissipations via a tough self-healing polymer matrix (TSP) that repeatedly dissipates energy via dynamic bond breakage and reformation.

In addition to interfacial design, serpentine structural design is another key feature of many stretchable hydrogel conductors with rigid or brittle electronic components. This undulating design allows the electronic components to stretch and deform together with the hydrogel matrix while minimizing the strain on the electronic components and preventing them from breaking. Figure 1.7c provides an overview of notable efforts in designing stretchable hydrogel conductors with serpentine-structured brittle electronic components. Implementing the serpentine structure design requires advanced manufacturing techniques, including micro-/nanofabrication [144, 159], 3D printing [145, 160, 161], stereographic design [146, 162, 163], and syringe injection [147, 164, 165].

1.4 Electrochemical Hydrogel Biosensors

Electrochemical biosensors have tremendous potential for medical diagnostics as they detect and measure the presence of biological molecules by converting biological events into electrical signals [166, 167]. However, previous efforts have primarily focused on graphene-based sensing materials and metallic electrodes, which have limitations when targeting *in-situ* detection. For instance, the presence of high ionic strength in body fluids produces pronounced screening of electrical fields, significantly reducing the sensitivity of the electrical chemical detection [168]. Additionally, biomarker concentrations in body fluids are typically about 10 ng/ml, which is too low to be detected by most existing electrical biosensors [169]. Recent studies show that introducing hydrogels into existing electrochemical biosensors can provide two unique benefits (Figure 1.8a) [170, 171]. First, hydrogels can act as a protective and selective layer for highly sensitive and specific *in-situ* detection, preventing interferences from biological environments (Figure 1.8b). Second, hydrogels can serve as surface modification to potentially push the limits of electrochemical properties of sensing materials, including Debye length, surface capacitance, and band gap (Figure 1.8c).



Figure 1.8 Design of electrochemical hydrogel biosensors. (a) Illustration of an electrochemical hydrogel biosensor for *in-situ* detection biomolecules. (b) Hydrogels act as a protective and selective layer for sensing materials. (c) Hydrogels serve as surface modification to potentially push the limits of electrochemical properties of sensing materials.

1.4.1 Selective Transport Design of Hydrogels

Efforts to tailor the transport properties of hydrogels mainly rely on two approaches: filtering by size (Figure 1.9a) and filtering by covalent interaction (Figure 1.9b) [172]. However, neither approach is suitable for selectively controlling the transport of biomolecules in body fluids. Filtering by size is impractical due to the small size difference between most biomolecules in body fluid; while filtering by covalent interaction is possible to distinguish the transport between target and nontarget biomolecules, such covalent interaction is only limited at the outmost surfaces of hydrogels and very challenging to achieve an enhanced transport throughout hydrogels due to the resistance from these covalent interactions. Recent studies have shown that introducing reversible interactions between target biomolecules and functional groups in hydrogels has the potential to achieve selective transport of target biomolecules across hydrogels [111, 112, 173, 174] (Figure 1.9c). The dissociation and reforming rates of the reversible interactions determine the mode and speed of biomolecular transport across hydrogels. Additionally, the network



Figure 1.9 Selective transport design in hydrogels. (a) Filtering by size. (b) Filtering by permanent interaction. (c) Filtering by reversible interactions.

elasticity of hydrogels plays an important role in controlling transport. Future opportunities may focus on the synergistic role of network elasticity and reversible interaction in independently controlling the transport of various biomolecules in hydrogels. Such fundamental knowledge may also aid in understanding the selective transport mechanisms employed in biological systems, such as sperm selection by mucus layers.

1.4.2 Electrochemical Design of Hydrogel-2D Material Interfaces

The other challenge faced by *in-situ* electrochemical biosensors is their inability to detect biological events in high ionic strength solutions, ubiquitous in body fluids [166, 175]. The presence of 0.9 wt% mobile ions (Na⁺, K⁺, Cl⁻) in body fluids leads to significant screening of electrical fields, which greatly reduces the sensitivity of electrochemical detection. For instance, the Debye length for 2D sensing materials in body fluids is typically below 1 nm, making it almost impossible to detect proteins (typically around 10 nm) above the Debye length [168]. A potential solution is to engineer the interface between the hydrogel and 2D material to enhance the field-effect sensing performance. Recent studies have shown that surface-functionalized 2D materials can achieve a 10-fold increase in Debye length from 0.82 to 9.6 nm in high ionic strength solutions [170]. One possibility is to graft crosslinked polymer networks of hydrogels onto existing sensing materials and explore the potential of optimizing the network topology and charge density in hydrogels to push the limits of electrochemical properties such as Debye length, surface capacitance, and band gap.

1.5 Flexible Hydrogel Biobattery

A flexible biobattery is a device that converts low-grade energy within the human body into usable energy and is essential for developing self-powered *in-situ* bioelectronic devices [176–179]. Hydrogels are a promising material for use in these devices because they are porous and can modulate ion and electron transport while minimizing potential leakage compared to traditional organic aqueous electrolytes. Furthermore, hydrogels are biocompatible and soft, which reduces damage to surrounding tissues. However, the power outputs of flexible hydrogel biobatteries are still relatively low and do not meet the power requirements of most *in-situ* bioelectronic devices. Figure 1.10a shows the power range and operation time requirements of common biomedical devices [180]. To develop the next generation of self-sustaining *in-situ* bioelectronic devices, high-performing hydrogels must be leveraged to harness various forms of energy within the human body (Figure 1.10b). This potential remains largely unexplored but is highly desirable. This section will discuss recent efforts in exploring the potential of powering *in-situ* bioelectronic devices through mechanical, chemical, and thermal energy harvesters (Figure 1.10c).



Figure 1.10 Flexible hydrogel batteries to self-power *in-situ* bioelectronic devices. (a) Examples of common implantable medical devices, with their required power supply and operation time (Source: Ref. [180]/Oxford University Press). (b) Schematic illustrations of the energy harvester working inside the intestine and (c) three forms of energy harvesting, including mechanical, chemical, and thermal energy, generated within the human body.

1.5.1 Mechanical Energy Harvester

Mechanical energy harvesters are devices that can convert the mechanical energy of the human body or organs into electrical energy for powering *in-situ* bioelectronic devices. One prime example of a mechanical energy harvester is the triboelectric nanogenerators (TENGs) [181–183], which generates an electrostatic potential difference between two materials of diverse polarities due to the triboelectric effect, causing a transfer of charges and the formation of an electric potential difference between them (Figure 1.11a) [177, 190–192]. Figure 1.11b presents a representative example of using ultrasound to induce vibrations and harness triboelectricity for in-body powering [184, 185]. Another way to harvest mechanical energy in the body is by leveraging the fluid-electro-mechanical coupling



Figure 1.11 Schematic illustration of the working mechanism of (a) mechanical energy harvester with (b) examples (Source: Ref. [184] and Ref. [185]), (c) chemical energy harvester with (d) examples (Source: Ref. [186] and Ref. [187]), and (e) thermal energy harvester with (f) examples (Source: Ref. [188] and Ref. [189]. © 2022/Elsevier).

of electrokinetic streams in porous materials to generate electricity [193, 194]. Electrokinetic mechanical energy harvesters typically involve applying external forces, such as pressure, to drive the movement of micro-/nanofluidic water across a porous membrane, thereby causing the motion of ions to produce electricity. Unlike conventional TENGs, electrokinetic mechanical energy harvester can harvest low-frequency body motions while potentially producing high power output [195].

1.5.2 Chemical Energy Harvesters

Chemical energy harvesters are devices that convert chemical energy into electrical energy through chemical reactions in an electrolyte (Figure 1.11c) [196, 197]. These devices usually have two electrodes and an electrolyte, where the electrolyte acts as a mediator for the chemical reactions. The chemical reactions that occur between the positive and negative electrodes cause the flow of electrons within the electrolyte, thus generating electrical energy that can be used in the circuit. Hydrogel, with its watery nature, is an excellent carrier for chemicals, and acts as the electrolyte [198–200]. Drawing inspiration from the electric eel's power generation mechanism, Yang and Mayer and coworkers harnessed the gradients of ions in hydrogels to develop soft, flexible, transparent, and biocompatible hydrogel biobatteries, generating 110 V at open circuit or 27 mW/m² per hydrogel cell, which

is a significant achievement in the realm of electrochemical energy harvesting via hydrogels (Figure 1.11d) [186, 187]. Despite the abundant chemical reactions that occur within the human body (such as those that occur during digestion resulting in pH differences), there has been no significant progress in *in-situ* chemical energy harvesting. The main barriers are related to unstable ion concentrations and uncontrolled ion types in body fluids. The stability and size of the device are still the main challenges.

1.5.3 Thermal Energy Harvesters

Thermal energy harvesters using thermoelectric materials (Figure 1.11e), such as thermoelectric hydrogels, offer the potential to harvest low-grade body heat and power *in-situ* bioelectronic devices. These soft and biocompatible thermoelectric hydrogels are regarded as favorable alternatives to conventional thermoelectric materials [201–205]. Recent studies by Chen and Liu and coworkers have demonstrated a giant positive thermopower of 17 mV/K in an ionic thermoelectric hydrogel by harnessing synergistic thermo-diffusion and thermo-galvanic effects. The thermos-diffusion effect is dominated by the presence of ions (KCl, NaCl, and KNO₃), while the thermo-galvanic effect is governed by a redox couple (Fe(CN)₆^{4–})/(Fe(CN)₆^{3–}), also adopted in other thermoelectric hydrogels (Figure 1.11f) [188, 189]. While thermoelectric hydrogels have significant potential for *in-situ* bioelectronics, the low power output and poor mechanical properties of these materials remain key limitations. Overcoming these limitations through further research and development will be crucial to fully exploit the unique advantages of thermoelectric hydrogels in *in-situ* bioelectronics.

1.6 Concluding Remarks

Over the past few years, we have seen many exciting advances and examples in the field of *in-situ* hydrogel bioelectronics that suggest great potential of high-performing hydrogels for many important applications. We will conclude this chapter with a set of opportunities by integrating interdisciplinary efforts in various areas of *in-situ* hydrogel bioelectronics, including ingestible sensors, neural interfaces, miniature robots, and data analytics. Ingestible sensors are one area where hydrogel-based bioelectronics can be leveraged (Figure 1.12a). These sensors can be designed to be swallowed and pass through the gastrointestinal tract, allowing for noninvasive monitoring of various biomarkers in real time. With the integration of hydrogel-based sensors, these devices can provide more accurate and reliable data, as hydrogels can respond to changes in pH, temperature, and other environmental factors [32]. Neural interface technology is another area where hydrogel-based bioelectronics can be applied (Figure 1.12b) [206-209]. By using hydrogels as a platform for neural interfaces, these devices can be made more biocompatible and less invasive, reducing the risk of rejection or other complications. With the integration of hydrogel-based sensors and actuators, these



Figure 1.12 Future opportunities of *in-situ* hydrogel bioelectronics. Schematic illustrations of (a) ingestible sensors, (b) neural interfaces, (c) miniature robots, and (d) predictive analytics.

interfaces can provide more accurate and precise control over prosthetic devices or assistive technologies, allowing for more natural movements and interactions with the environment. Miniature robots are also an exciting area of research for *in-situ* hydrogel bioelectronics (Figure 1.12c) [210, 211]. With the use of hydrogels, these robots can be made more flexible and compliant, allowing for safer and more effective integration with the body. By incorporating hydrogel-based sensors and actuators, these robots can be controlled and manipulated to perform targeted drug delivery, tissue engineering, and surgical procedures. Finally, the integration of data analytics and machine learning algorithms is crucial for unlocking the full potential of *in-situ* hydrogel bioelectronics (Figure 1.12d) [212, 213]. With the vast amounts of data generated by these devices, there is a need for advanced analytics tools to help interpret and make sense of the data. By leveraging these tools, we can gain new insights into biological systems and develop more effective treatments and therapies.

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