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## Toughening Hydrogels Through Force-triggered Chemical Reactions that Lengthen Polymer Strands

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**Abstract:** The utility and lifetime of materials made from polymer networks, including hydrogels, depend on their capacity to stretch and resist tearing. In gels and elastomers, those mechanical properties are often limited by the covalent chemical structure of the polymer strands between cross-links, which is typically fixed during the material synthesis. Here, we report polymer networks in which the constituent strands lengthen through force-coupled reactions that are triggered as the strands reach their nominal breaking point. Reactive strand extensions of up to 40% lead to hydrogels that stretch 40-50% further than, and exhibit tear energies twice that of, networks made from analogous control strands. The enhancements are synergistic with those provided by double network architectures, and complement other existing toughening strategies.

#### Main Text:

Polymeric materials such as elastomers gradually break down under repeated use. This process of physical fatigue is due to a series of microscopic tearing events that occur during periods of mechanical strain (1, 2), and it limits the useful lifetime of products ranging from automobile tires to prosthetic implants, and from stretchable displays to soft robots and other devices. Polymer network fatigue therefore contributes to a wide range of undesirable endpoints, including the rate at which commodity rubber products end up in landfills (ca.  $2.5 \times 10^8$  tires/yr in the U.S. alone) (3), the number of replacement surgeries due to failing implants such as artificial knees (4, 5), and the limits to which flexible displays can be stretched before breaking (6). As a result, recent years have seen extensive effort given to strategies that improve the durability of polymer networks (7, 8), including the use of multi-network architectures (7, 9-11), reversible cross-linking (10, 12), noncovalent domain unfolding (13-16), and the addition of fillers (17). All of these approaches have in common that the primary, covalent chemical structure of the network remains fixed up until the point that the polymer strand breaks.

Polymer networks that actively remodel their primary, covalent structure during use therefore have the potential to provide property enhancements that complement existing mechanisms. Recent advances in covalent polymer mechanochemistry now enable the synthesis of polymer strands that extend the covalent bonding along their backbone when stretched (18), as a result of force-coupled chemical reactions that are triggered at or just before the strands reach their nominal breaking point. Because these reactions are typically associated with high forces (~nN), the amount of energy absorbed at the single molecular level can be quite large relative to the energy absorbed by a covalent strand in the absence of the reaction-enabled extension (19-21). Existing models of network fracture draw a direct relationship between the energy required to propagate a crack through a rubbery material (the tear energy, T) and the energy required to stretch an individual polymer strand to the point at which it breaks (the chain fracture energy, U) (22, 23). We therefore hypothesized that by turning a polymer network into a literal network of responsive chemical reactions, reactive strand extension might improve the properties of bulk materials.

A Toughening DN by reactive strand extension

B Chemical designs of polyelectrolytes in first network



**Fig. 1.** The reactive strand extension (RSE) concept in double network hydrogels (strand scission events are not shown). (A) Schematic of double networks RSE and an analogous control polymer. (B) Chemical structures of the first network co-polymers employed in this work.

Here we demonstrate that reactive strand extension (RSE) can be used to improve the stretchability and tear resistance of hydrogels. The RSE concept is shown in Figure 1, and the strand extension is provided by the sodium salts of bicyclo[6.2.0]decane (BCD) mechanophores, which react via a force-coupled [2+2] cycloreversion to release stored length (24). BCD copolymers are formed via the radical addition co-polymerization of the corresponding bicyclic cyclobutene carboxylate monomer 1a. The mechanochemical activity of the sodium salts of BCD was quantified using single-molecule force spectroscopy (SMFS) implemented through an atomic force microscope (AFM). Stretching of the single molecules to the forces required for BCD cycloreversion requires strong attachments of the polymer to the AFM tip, and we found that an acetoacetate group could be introduced to promote rapid and strong bonding to aminefunctionalized AFM tips (25, 26). Co-polymerization of **1a** with acrylamido-2-methylpropane sulfonate (NaAMPS) 2 and acetoacetate methacrylate 3 provided linear co-polymers P1 with 1a:2:3 = 0.22:0.62:0.16. At forces of up to ~1500 pN, the SMFS curves are typical of a flexible polymer strand, as extension initially requires only low forces (order of 1 - 10 pN) to overcome the conformational entropy associated with bond torsions along the polymer main chain. As the extension increases, the available conformational degrees of freedom are exhausted and the trapped polymer strand reaches end-to-end distances that approach its fully extended contour length (order of  $10^2$  pN). Subsequent stretching is dominated by the deformation of bond angles

and bond lengths, and only small fractional extensions are achieved with increasing force, as the polymer nears its physical stretching limits before breaking (order of  $10^3$  pN). With the BCD mechanophore, however, the force-separation curve shows a reproducible transition across 1500-2000 pN, in which the release of stored length of up to ~40% of the initial polymer contour length is observed. This stored length is consistent with that expected from the BCD cycloreversion, and the transition force is similar to that reported in other cyclobutane mechanophores (20, 27, 28). The onset of extension also provides a measure of the scission point of control strands made from **1b** (no fused ring), and CoGEF calculations (see Fig. S2) confirm that the scission behave of the fused and unfused cyclobutanes are effectively indistinguishable (29). Thus, BCD incorporation provides the desired molecular behavior: as the polymer strands reach their nominal stretching limit, the presence of selective mechanophores converts strand scission events into force-coupled covalent rearrangements that allow the strands to survive forces and extensions that would otherwise be catastrophic.



**Fig. 2.** Mechanochemical reactions driving RSE. (A) Mechanically coupled [2+2] cycloreversion leads to chain extension in [6.2.0]bicyclodecane mechanophores and to chain scission in analogous acyclic cyclobutane mechanophores. (B) Representative single molecule force-extension curves of linear RSE co-polymers in water and DMSO.

To see whether this single molecule behavior has an impact on bulk material properties, the mechanochemically active co-polymers were incorporated into double-network (DN) hydrogels, whose robust mechanical properties facilitate the characterization of mechanical response. A

DN hydrogel consists of two interpenetrating networks with contrasting structure: the first network comprises strands in a highly stretched state and is rigid and brittle. The second network comprises strands in a coiled state, and as a result it is soft and stretchable (30, 31). This doublenetwork structure allows multiple strands within the first brittle network to break independently without causing catastrophic breakdown of the entire brittle network, as the local strain amplification created by the breaking of the brittle strand is suppressed by transferring the load to the soft network strands nearby rather than propagating through the first network (31-34). Hydrophilic polymer networks were made via the radical addition co-polymerization of the sodium salts of BCD 1a, NaAMPS 2 (aqueous feed concentrations of 0.3 M and 0.7 M for 1a and 2, respectively) and N,N'-methylenebisacrylamide (MBAA) cross-linker (8 mol% or 4 mol%) of total monomer feed ratio). In the DNs, this BCD-AMPS co-polymer was used as the first network, and sparsely cross-linked polyacrylamide PAAm (AAm, 4 M; MBAA, 0.01 mol%) was used as the soft and stretchable second network. The DN structure is formed by swelling the formed first network in a precursor monomer solution of the second network, followed by polymerization of the second network (see Fig. S1). Two first networks, N1 and N2, of the same BCD-AMPS content (BCD : AMPS = 3 : 17 mol/mol; ~ 40% contour length increase expected from CoGEF calculations, see Fig. S2) but different cross-linking density (8 mol% and 4 mol% of MBAA for N1 and N2, respectively), were prepared (see Table S1 and Fig. S1).

Control networks **N1-con** and **N2-con** were synthesized using acyclic cyclobutene carboxylate **1b**. The structural similarity of **1a** and **1b** was expected to lead to similar reactivity in the polymerization and, thus, similar network structure in **N1** and **N1-con**, or in **N2** and **N2con**. This expectation was verified in multiple ways. First, the reactivity ratios of **1a** and **1b** with NaAMPS were measured and found to be  $r_{Ia} = 0.19 \pm 0.01$ ;  $r_{NaAMPS} = 1.85 \pm 0.09$  and  $r_{Ib} = 0.16 \pm 0.01$ ;  $r_{NaAMPS} = 1.87 \pm 0.09$ , respectively (see Fig. S3). Second, the equilibrium swelling ratios (V/V<sub>0</sub>) of **N1** (12.4 ± 0.1) and **N2** (32.7 ± 0.7) in water are indistinguishable from those of **N1con** (12.1 ± 0.9) and **N2-con** (32.7 ± 1.8), respectively. Third, **N1**, **N1-con**, **N2**, and **N2-con** were disassembled through the hydrolysis of the bis-acrylamide cross-linker **4**, and the gel permeation chromatography (GPC) elugrams of the fragments were found to be very similar (see Fig. S4). Finally, these single networks were too brittle for reliable mechanical characterization, but forming DNs by cross-linking PAAm inside the swollen first networks resulted in moduli of **DN1** (288 ± 4 kPa) and **DN1-con** (290 ± 8 kPa) that fall within experimental uncertainty. The

	E (kPa)	$\boldsymbol{\varepsilon}_{b}$	$\sigma_b$ (MPa)	$W_{\rm b}$ (MJ m <sup>-3</sup> )	$\varepsilon_p$	<i>T</i> (J m <sup>-2</sup> )
DN1	$288\pm4$	10.0	0.92	6.4	2.3	$2840 \pm 103$
DN1-con	$290 \pm 8$	4.9	0.61	2.3	1.7	$1419\pm52$
DN2	$243 \pm 5$	7.0	0.64	2.6	1.3	$554\pm40$
DN2-con	$249 \pm 13$	4.7	0.54	1.6	0.8	276 ± 10
PAAm	$57\pm 6$	5.0	0.18	0.4	/	~10

moduli of **DN2** ( $243 \pm 5$  kPa) and **DN2-con** ( $249 \pm 13$  kPa) are similarly matched. Thus, any differences in ultimate stretch and tear resistance between the active and control networks are attributed to the conversion of scission to molecular extension that is allowed in the BCD copolymers by virtue of the designed release of length stored in the bicyclic ring structure.

The incorporation of the BCD mechanophore leads to substantial changes in the mechanical limits of its double networks. The resistance to tearing was evaluated by introducing a notch into DN films and measuring the strain necessary for the notch to propagate under pure shear loading. Figure 3A, B shows the stress-strain behavior of notched **DN1** and **DN1-con** under uniaxial extension. The critical stretch for crack propagation in **DN1** is  $234 \pm 4\%$ , which is 40% greater than the  $167 \pm 7\%$  observed in **DN1-con**. The gain in stretchability is found as well in **DN2**, for which the critical stretch for crack propagation is  $128 \pm 8\%$ , vs.  $83 \pm 2\%$  in **DN2-con** (53% greater strain). The resistance to tearing can be further quantified in terms of the tear energy *T*, or the energy required per area of crack growth, which was determined via the method of Rivlin and Thomas (*35*). Figure 3D shows that both **DN1** and **DN2** have roughly double the *T* of their respective controls:  $2,840 \pm 103$  J/m<sup>2</sup> vs.  $1,419 \pm 52$  J/m<sup>2</sup> for **DN1** vs. **DN1-con**, and  $554 \pm 40$  J/m<sup>2</sup> vs. 276  $\pm 10$  J/m<sup>2</sup> for **DN2** vs. **DN2-con**.



**Fig. 3.** Response of DNs to uniaxial tension. (A) Single-edge notched **DN1** at stretch ratios  $\lambda$  (mm/mm) of 1.3, 2.5, and 3.2. (B) Single-edge notched **DN1-con** at stretch ratios (mm/mm) of 1.2, 2.4, and 2.6. (C) Representative stress-strain curves (pure shear, 2 mm thick films) of unnotched DNs and a single PAAm network prepared identically to the second network of the DNs. (D) Propagation strain and tear energy of DNs. (E) Hysteresis in cyclic loading/unloading of **DN1**, **DN1-con**, and a single PAAm network prepared identically to the second network of the DNs.

The improved mechanical properties due to reactive strand extension are also observed in unnotched DNs (see Fig. S5). Figure 3C shows the stress-strain behavior of a pristine **DN1** film

under uniaxial extension. The film survives to an ultimate stretch of  $890 \pm 120\%$  before breaking, whereas the acyclic control **DN1-con** fails at a lower stretch of  $479 \pm 99\%$ . The high strains obtained in the unnotched films reveal that RSE changes the internal fracture patterns of the DN. Most notably, **DN1** shows a clear yielding, suggesting that the first network ruptures into a global discontinuous phase that enables the load to be transferred to the second network in the yielded region. On the other hand, **DN1-con** shows a gradual increase in stress without a clear plateau stress region, suggesting that the internal rupture of the first network is relatively random and does not achieve a global discontinuous structure up to sample failure. We attribute this to the fact that fracture of **N1-con** within **DN1-con** begins at lower strain than **N1**. When a small defect forms, that stress largely remains in N1-con, because the second network is still too soft for stress transfer (the local strain at the defect is not great enough). Thus, the fracture propagates further across **N1-con** within the DN and only begins to transfer to the second network after the defect becomes larger. In DN1, fracture of N1 initiates at higher strain, at which point a greater fraction of the stress is transferred to the second network, and so the fracture event is blunted and the next fracture occurs elsewhere, leading to the desired fractal/domain patterns for greater toughness (31).

The presence of RSE should alter the energy dissipation behavior from that observed in conventional DN hydrogels, because it can occur through covalent rearrangement without chain scission. Here, we examined the change in modulus and energy dissipated in sequential loading/unloading cycles response under uniaxial tension, where the maximum strain increases from one loading cycle to the next. When a DN gel is stretched to a given ratio for the first time and then unloaded, there is significant hysteresis (*33*); this typical DN behavior is observed as well in our systems (Figure 3E; low strain behavior expanded in Figure 4A,B). Typically, hysteresis in DN gels is attributed almost entirely to chain scission in the first network (*31*), and it is therefore correlated with a decrease in modulus due to the loss of stress-bearing network



**Fig. 4.** Hysteresis in cyclic loading/unloading of **DN1** (A), and **DN1-con** (B) in small strain range; (C) Fractional loss of Young's modulus of DN1 and DN1-con as a function of strain; (D) Fractional hysteresis as a functional of fractional loss in modulus loss for **DN1** and **DN1-con**.

strands (*36*). RSE suppresses this loss in modulus; greater strains are required to trigger a measurable change in modulus in **DN1** than **DN1-con**, and the loss in modulus in **DN1** consistently lags than observed in **DN1-con** as a function of strain (Figure 4C). RSE should also alter the correlation between energy dissipation and change is modulus, by allowing dissipation without chain scission. As seen in Figure 4D, **DN1** exhibits greater hysteresis per loss in modulus than **DN1-con**. The difference is greatest at the early stages of loading (lower strains), where the ratio of extended but unbroken strands to broken strands is expected to be largest.

Looking forward, the concept of reactive chain extension demonstrated here has numerous implications for network design. First, reactive chain extension occurs only at large strains and only at the specific molecular locations that are about to break without it. Unlike the incorporation of weak, reversible cross-linking or noncovalent interstrand interactions, the enhancement does not come with a concomitant increase in the viscous modulus of the network.

In other words, the energy dissipation is triggered only when needed to resist fracture and need not be overcome in routine functions of the material when fracture is not imminent. Second, the property enhancements are achieved within networks that otherwise have identical structure and modulus to their non-mechanophore controls, meaning that reactive chain extension might be introduced into at least some materials whose properties are already optimized for a given function. We note that the magnitude of the effect seen here is large enough that, if it were translated to fatigue-prone materials, it would have a substantial impact on material life cycle. The correlation between fatigue life of rubbery materials and T has been appreciated for some time (35, 37), with scaling dependences of  $T^2$  noted (38). Even greater property enhancements should be obtainable in the future through the optimization of mechanophore design, polymerization strategy, and multi-network architecture. Notably, relative to control DN gels, RSE provides this fatigue resistance while also suppressing the loss in modulus that accompanies energy dissipation. Third, prior examples of mechanochemically induced changes in macroscopic physical properties involve either activation in solution (21, 39), irreversible plastic deformation (40, 41), and/or the presence of reagents beyond the initial network (11, 39, 41); we are not aware of direct correlations of macroscopic properties and mechanochemical response within an intact strand. Mechanophore designs offer an increasing range of responses (42), and the chemical changes that accompany reactive chain extension might be designed to produce additional, desirable responses. For example, the ring opening of mechanophores similar to those employed here can be reversed photochemically (43), offering the potential for new types of molecular-scale repair that could further extend material lifetime. Finally, whereas reactive chain extension is demonstrated here in the context of DN hydrogels, single chain response should also contribute to the properties of single networks, elastomers, and polymer composites. The investigation of this rich structure-activity space, including the interplay of molecular response and network architecture, presents a rich set of opportunities and challenges that span synthetic, physical, theoretical, and analytical aspects of polymer chemistry.



**Fig. 1.** The reactive strand extension (RSE) concept in double network hydrogels (strand scission events are not shown). (A) Schematic of double networks RSE and an analogous control polymer. (B) Chemical structures of the first network co-polymers employed in this work.



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**Table 1.** Young's modulus (*E*), nominal strain, nominal stress, and input energy density at break  $(\varepsilon_b, \sigma_b, \text{ and } W_b, \text{ respectively})$  of unnotched films in unconstrained uniaxial tension, and crack propagation strain  $(\varepsilon_p)$  and tearing energy (*T*) of films in a single-edge notched geometry; Tearing energy of PAAm gel is listed as reported (44).

	E (kPa)	$\boldsymbol{\varepsilon}_{b}$	$\sigma_b$ (MPa)	$W_{\rm b}$ (MJ m <sup>-3</sup> )	$arepsilon_p$	<i>T</i> (J m <sup>-2</sup> )
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**Fig. 4.** Hysteresis in cyclic loading/unloading of **DN1** (A), and **DN1-con** (B) in small strain range; (C) Fractional loss of Young's modulus of DN1 and DN1-con as a function of strain; (D) Fractional hysteresis as a functional of fractional loss in modulus loss for **DN1** and **DN1-con**.

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