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Thermally-healable network solids of sulfur-crosslinked poly(4-allyloxystyrene)[†]

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Network polymers of sulfur and poly(4-allyloxystyrene), **PAOS_x** (*x* = percent by mass sulfur, where *x* is varied from 10–99), were prepared by reaction between poly(4-allyloxystyrene) with thermal homolytic ring-opened S₈ in a thiol-ene-type reaction. The extent to which sulfur content and crosslinking influence thermal/mechanical properties was assessed. Network materials having sulfur content below 50% were found to be thermosets, whereas those having >90% sulfur content are thermally healable and remeltable. DSC analysis revealed that low sulfur-content materials exhibited neither a *T_g* nor a *T_m* from –50 to 140 °C, whereas higher sulfur content materials featured *T_g* or *T_m* values that scale with the amount of sulfur. DSC data also revealed that sulfur-rich domains of **PAOS₉₀** are comprised of sulfur-crosslinked organic polymers and amorphous sulfur, whereas, sulfur-rich domains in **PAOS₉₉** are comprised largely of α-sulfur (orthorhombic sulfur). These conclusions are further corroborated by CS₂-extraction and analysis of extractable/non-extractable fractions. Calculations based on TGA, FT-IR, H₂S trapping experiments, CS₂-extractable mass, and elemental combustion microanalysis data were used to assess the relative percentages of free and crosslinked sulfur and average number of S atoms per crosslink. Dynamic mechanical analyses indicate high storage moduli for **PAOS₉₀** and **PAOS₉₉** (on the order of 3 and 6 GPa at –37 °C, respectively), with a mechanical *T_g* between –17 °C and 5 °C. A **PAOS₉₉** sample retains its full initial mechanical strength after at least 12 pulverization-thermal healing cycles, making it a candidate for facile repair and recyclability.

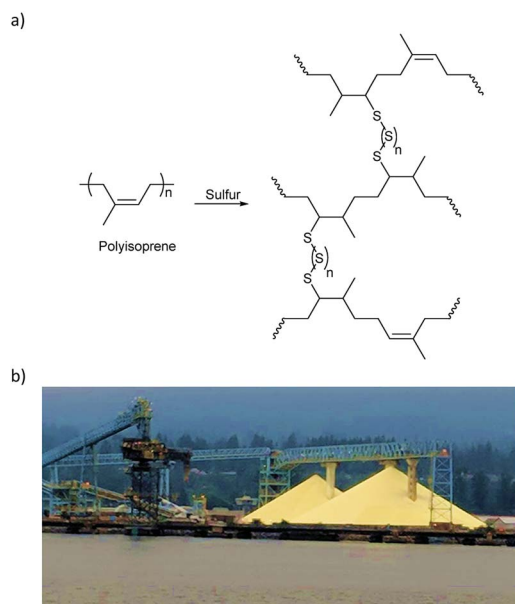
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Introduction

Vulcanization is a process developed by Charles Goodyear¹ in which the mechanical properties of natural rubber (polyisoprene) are dramatically enhanced by heating it in the presence of up to 5 wt% elemental sulfur (as S₈ at STP). The increase in material strength of a rubber following vulcanization results from covalent crosslinking between polymer chains.² Such crosslinking is derived from homolytic thermal ring opening of S₈ to afford sulfur-centered radicals³ that subsequently add to C=C bonds, leading to polysulfide crosslinks between polymer chains (Scheme 1a). Without the dramatic increase in mechanical strength endowed upon rubber products by polysulfide crosslinks formed during vulcanization, the modern automobile tire would not be accessible, and early automobiles



Scheme 1 (a) Sulfur vulcanization of polyisoprene. (b) By-product sulfur at North Vancouver Sulphur Works, originating from tar sand-processing facilities in Alberta, Canada.

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[†] Electronic supplementary information (ESI) available: Proton NMR of **PVP** and **PAO**; IR spectra of **PAOS_{10–99}**; DSC curves of **PVP**, S₈, **PAO**, and **PAOS_{10–99}**. See DOI: 10.1039/c8ra06847j



would have been severely limited in the types and duration of tasks they could perform.

Despite the dramatic structural improvement mediated by sulfur as a minor component by mass of vulcanized rubber, elemental sulfur is currently underutilized as a bulk component of materials. The underutilization of sulfur is especially surprising considering the titanic quantities of S_8 (Scheme 1b) that are produced by petrochemical industries. Petrochemical by-product sulfur is produced by the Claus process,^{4,5} wherein H_2S is oxidized to S_8 . Sulfur removal from fuels is necessary to prevent catalyst poisoning in subsequent steps of the petroleum-refining process and to attenuate the environmental impact of acid rain that results from combustion of sulfur-rich fossil fuels.⁶ Currently, the primary use of the S_8 generated from fossil fuels is in the production of sulfuric acid, with lesser quantities being used in fertilizers/pesticides, vulcanization of rubber, and as an additive to asphalt.^{7–9} Even after consumption of S_8 by these processes, however, 7 million tons of S_8 accumulate each year as unused waste.¹⁰ Vast quantities of S_8 have been stockpiled at industrial centers and storage sites, so S_8 is an inexpensive, abundant, and readily accessible feedstock for valorization.^{6,11–13} Furthermore, the formation of S–S bonds is thermally reversible, a property that could be exploited to generate recyclable thermoplastics and thermally self-healing materials.¹⁴

Although elemental sulfur itself has very poor mechanical properties, significant advances have been achieved with the advent of inverse vulcanization.¹⁵ Inverse vulcanization is a term coined by Pyun for a process in which S_8 is the bulk material and organic small molecules form the covalent crosslinks between polysulfide chains.^{6,15–27} Pyun's breakthrough studies on such high sulfur-content materials have inspired a flurry of related studies to better employ byproduct sulfur.¹² Materials prepared by inverse vulcanization can retain some of the advantageous properties of S_8 . For example, S_8 exhibits high electrical resistivity ($2 \times 10^{23} \mu\Omega \text{ cm}$) and low thermal conductivity ($0.205 \text{ W m}^{-1} \text{ K}^{-1}$). In addition, S_8 actively resists several material degradation pathways, as it is resistant to corrosion by strong acids. It also repels insects and rodents,²⁸ as well as having antibacterial and anti-microbial properties,^{29–34} and has long been used as a treatment for timbers to prevent decay.^{35,36} Moreover, if an organic small molecule used in inverse vulcanization has more than one C=C bond, then that molecule can crosslink multiple polysulfide chains to afford a networked covalent composite (NCC) or polymer material.

Many of the inverse vulcanized materials prepared by Pyun exhibit significantly greater durability compared to sulfur itself, even for materials containing >90% sulfur by weight. Because even small amounts of an alkene-functionalized organic small molecule can afford materials with dramatically enhanced mechanical properties relative to pure S_8 , we envisioned that using an alkene-functionalized organic polymer would afford an NCC with enhanced networking. We hypothesized that using this strategy to increase networking in NCCs would increase the mechanical strength of the NCCs. Herein we report the synthesis of allyloxy-functionalized polystyrene, its crosslinking with S_8 *via* inverse vulcanization, and the thermomechanical

properties of the resulting NCCs, **PAOS_x** (x = percent by mass sulfur in the material), wherein x is varied from 10 to 99.

Experimental section

General considerations

All NMR spectra were recorded on a Bruker Avance spectrometer operating at 300 MHz for protons. Thermogravimetric analysis (TGA) was recorded on a TA SDT Q600 instrument over the range 20 to 800 °C, with a heating rate of 5 °C min^{−1} under a flow of N_2 (100 mL min^{−1}). Differential Scanning Calorimetry (DSC) was acquired using a Mettler Toledo DSC 3 STAR^e System over the range of −50 to 140 °C, with a heating rate of 10 °C min^{−1} under a flow of N_2 (200 mL min^{−1}). Each DSC measurement was carried out over five heat-cool cycles to confirm consistent results following the first heat-cool cycle. The data reported were taken from the second cycle of the experiment. Dynamic Mechanical Analysis (DMA) was performed using a Mettler Toledo DMA 1 STAR^e System in dual cantilever mode. DMA samples were cast from silicone resin molds (Smooth-On Oomoo® 30 tin-cure). The sample dimensions were 1.5 × 8 × 50 mm. Clamping force was 1 cN m and the temperature was varied from −60 to 88 °C with a heating rate of 2 °C min^{−1}. The measurement mode was set to displacement control with a displacement amplitude of 5 μm and a frequency of 1 Hz. Fourier transform infrared spectra were obtained using a Shimadzu IRAffinity-1S instrument operating over the range of 400–4000 cm^{−1} at ambient temperature using an ATR attachment. Poly(4-vinylphenol) (M_w = 11 000 g mol^{−1}, Sigma Aldrich), allyl bromide (Oakwood Chemical), elemental sulfur (99.5+%, Duda Energy, LLC) were used without further purification.

Preparation of poly(4-allyloxystyrene) (PAO)

The procedure that follows is based on a reported method.³⁷ To a 500 mL round bottom flask equipped with a Teflon-coated magnetic stir bar was added 22.489 g (185.87 mmol) of allyl bromide, 20.000 g (166.44 mmol) of poly(4-vinylphenol) (**PVP**) and 200 mL of acetonitrile. The contents of the flask were stirred for 5 min to facilitate complete dissolution of **PVP**. The flask was then equipped with a water-cooled reflux condenser and placed under an atmosphere of N_2 . The flask was heated in an oil bath at 80 °C while stirring. While the heated reaction mixture was stirring, 49.999 g (361.77 mmol) of K_2CO_3 was added portionwise to the reaction mixture. The reaction was then refluxed with stirring for 72 h, after which time the mixture was allowed to cool to room temperature. The flask was opened to the air and deionized water (250 mL) was added to the reaction flask to dissolve the remaining K_2CO_3 .

The reaction mixture was then extracted with dichloromethane (450 mL). The organic layer was collected and washed with 1 M HCl(aq.) (1 × 250 mL), and then with deionized water (1 × 250 mL). The organic layer was collected, and the solvent was evaporated under reduced pressure. The product was collected as a beige solid. ¹H NMR (300 MHz, DMSO-d₆, δ: 1.41 (br, 2H), 1.72 (br, 1H), 4.45 (br, 2H), 5.21 (br, 1H), 5.35 (br, 1H), 5.99 (br, 1H), 6.48–6.69 (br, 4H). Elemental analysis calculated



for C₁₁H₁₂O: C, 82.45; H, 7.56%. Found C, 81.88; H, 7.77%. These data are consistent with those previously reported.³⁷ DSC data for this compound are provided in the ESI.†

General synthesis of PAOS_x

Elemental sulfur was weighed directly into an aluminum reaction vessel. The vessel was heated to 180 °C, over which time the sulfur melted. Once the sulfur turned a viscous dark red-orange color (indicative of thermal ring-opening), the appropriate amount of **PAO** was slowly added to the molten sulfur. The reaction media was manually stirred with a spatula for 60 min. After 60 min the reaction was stopped, the mixture was allowed to cool, and the solid was removed from the reaction vessel. Reagent masses and results of elemental combustion micro-analysis are provided below.

Synthesis of PAOS₉₉

CAUTION: Heating elemental sulfur with organics can result in the formation of H₂S gas. H₂S is toxic, foul-smelling, and corrosive. The general synthesis above was used to synthesize **PAOS₉₉** (99 wt% sulfur) where 9.90 g of elemental sulfur and 0.0994 g of **PAO** were used in the reaction. Elemental analysis calculated: C, 0.82; S, 99.00; H, 0.07%. Found: C, 1.02; S, 98.21; H, 0.0%.

Synthesis of PAOS₉₀

The general synthesis above was used to synthesize **PAOS₉₀** (90 wt% sulfur) where 9.00 g of elemental sulfur and 0.998 g of **PAO** were used in the reaction. Elemental analysis calculated: C, 8.29; S, 90.00; H, 0.69%. Found: C, 8.07; S, 89.37; H, 0.64%.

Synthesis of PAOS₈₀

The general synthesis above was used to synthesize **PAOS₈₀** (80 wt% sulfur) where 2.40 g of elemental sulfur and 0.600 g of **PAO** were used in the reaction. Elemental analysis calculated: C, 16.59; S, 80.00; H, 1.40%. Found: C, 16.32; S, 79.70; H, 1.48%.

Synthesis of PAOS₇₀

The general synthesis above was used to synthesize **PAOS₇₀** (70 wt% sulfur) where 2.10 g of elemental sulfur and 0.900 g of **PAO** were used in the reaction. Elemental analysis calculated: C, 24.89; S, 70.00; H, 2.08%. Found: C, 25.75; S, 63.18; H, 2.24%.

Synthesis of PAOS₆₀

The general synthesis above was used to synthesize **PAOS₆₀** (60 wt% sulfur) where 1.80 g of elemental sulfur and 1.20 g of **PAO** were used in the reaction. Elemental analysis calculated: C, 33.19; S, 60.00; H, 2.79%. Found: C, 34.54; S, 61.91; H, 3.08%.

Synthesis of PAOS₅₀

The general synthesis above was used to synthesize **PAOS₅₀** (50 wt% sulfur) where 1.50 g of elemental sulfur and 1.50 g of **PAO** were used in the reaction. Elemental analysis calculated: C, 41.49; S, 50.00; H, 3.49%. Found: C, 41.50; S, 49.90; H, 3.75%.

Synthesis of PAOS₄₀

The general synthesis above was used to synthesize **PAOS₄₀** (40 wt% sulfur) where 1.20 g of elemental sulfur and 1.80 g of **PAO** were used in the reaction. Elemental analysis calculated: C, 49.78; S, 40.00; H, 4.18%. Found: C, 51.42; S, 37.65; H, 4.81%.

Synthesis of PAOS₃₀

The general synthesis above was used to synthesize **PAOS₃₀** (30 wt% sulfur) where 0.900 g of elemental sulfur and 2.10 g of **PAO** were used in the reaction. Elemental analysis calculated: C, 58.08; S, 30.00; H, 4.89%. Found: C, 60.25; S, 27.08; H, 5.70%.

Synthesis of PAOS₂₀

The general synthesis above was used to synthesize **PAOS₂₀** (20 wt% sulfur) where 0.600 g of elemental sulfur and 2.40 g of **PAO** were used in the reaction. Elemental analysis calculated: C, 66.38; S, 20.00; H, 5.58%. Found: C, 66.55; S, 18.22; H, 6.11%.

Synthesis of PAOS₁₀

The general synthesis above was used to synthesize **PAOS₁₀** (10 wt% sulfur) where 0.300 g of elemental sulfur and 2.70 g of **PAO** were used in the reaction. Elemental analysis calculated: C, 74.67; S, 10.00; H, 6.28%. Found: C, 73.62; S, 9.40; H, 7.10%.

Results and discussion

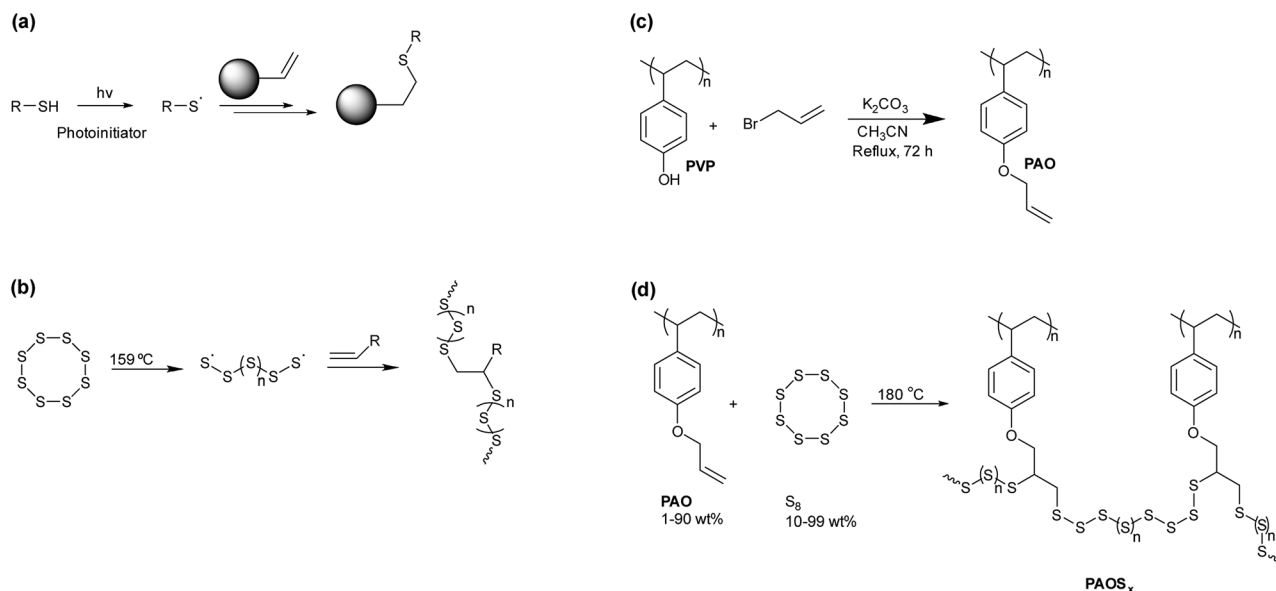
Synthesis

Polystyrene (PS) was selected as the scaffold for the alkene-bearing polymer substrate to access NCCs because a wide variety of functionalized PS derivatives are commercially available with relatively high molecular weights and well-defined polydispersities. The cross-linking mechanism in inverse vulcanization is analogous to the well-known thiol-ene click reaction (Scheme 2a).³⁸ Whereas the thiol-ene reaction requires a thiol group and an organic radical initiator to generate a sulfur-centered radical, in inverse vulcanization elemental sulfur (S₈) undergoes thermal homolytic ring-opening above 159 °C to generate a S-centered radical that can react with an olefin (Scheme 2b).

We reasoned that a PS derivative functionalized with hydroxyl groups could be readily modified with allyl bromide, thus poly(4-vinylphenol) (**PVP**) was employed as the initial starting material. Exhaustive alkylation was achieved upon reaction of **PVP** with allyl bromide in the presence of K₂CO₃, affording the desired PS derivative with alkene side chains, poly(4-allyloxystyrene) (**PAO**), in quantitative yield (Scheme 2c).³⁷

Analysis of **PAO** by ¹H NMR and FT-IR spectroscopy (Fig. 1a) revealed that the signals for the OH proton and the O-H stretching vibration (3300 cm⁻¹) had disappeared. Two new peaks appeared in the FT-IR spectrum at 922 and 995 cm⁻¹ that were consistent with monosubstituted alkene (*i.e.*, R-CH=CH₂) C-H bending modes (Fig. 1b). The ratio of integrals for the allyloxy proton signals to the integral of all PS-backbone proton





Scheme 2 (a) The general synthetic scheme of a thiol–ene reaction. (b) The homolytic ring-opening of S_8 to form a sulfur-centered radical, which then can react with olefins. (c) Synthetic route to prepare PAO. (d) Synthetic route to $PAOS_x$, where x represents the wt% of S_8 .

signals revealed that allylation of hydroxyl groups in **PVP** was quantitative.^{39,40}

Thermal reaction of **PAO** with sulfur (Scheme 2d) was achieved by heating the polymer to 180 °C in the presence of varying amounts of sulfur to afford the desired NCCs comprising **PAO** crosslinked with varying relative wt% of sulfur

($PAOS_x$; $x = 10, 20, 30, 40, 50, 60, 70, 80, 90$, and 99 wt% S). Each $PAOS_x$ was a brown-red solid with a glass-like appearance. The general physical appearance was consistent across the series, but the materials became more brittle with decreasing wt% sulfur. Mass losses in the range of 2–7% were observed during the reactions of **PAO** with S_8 (Table 1), which were attributed to

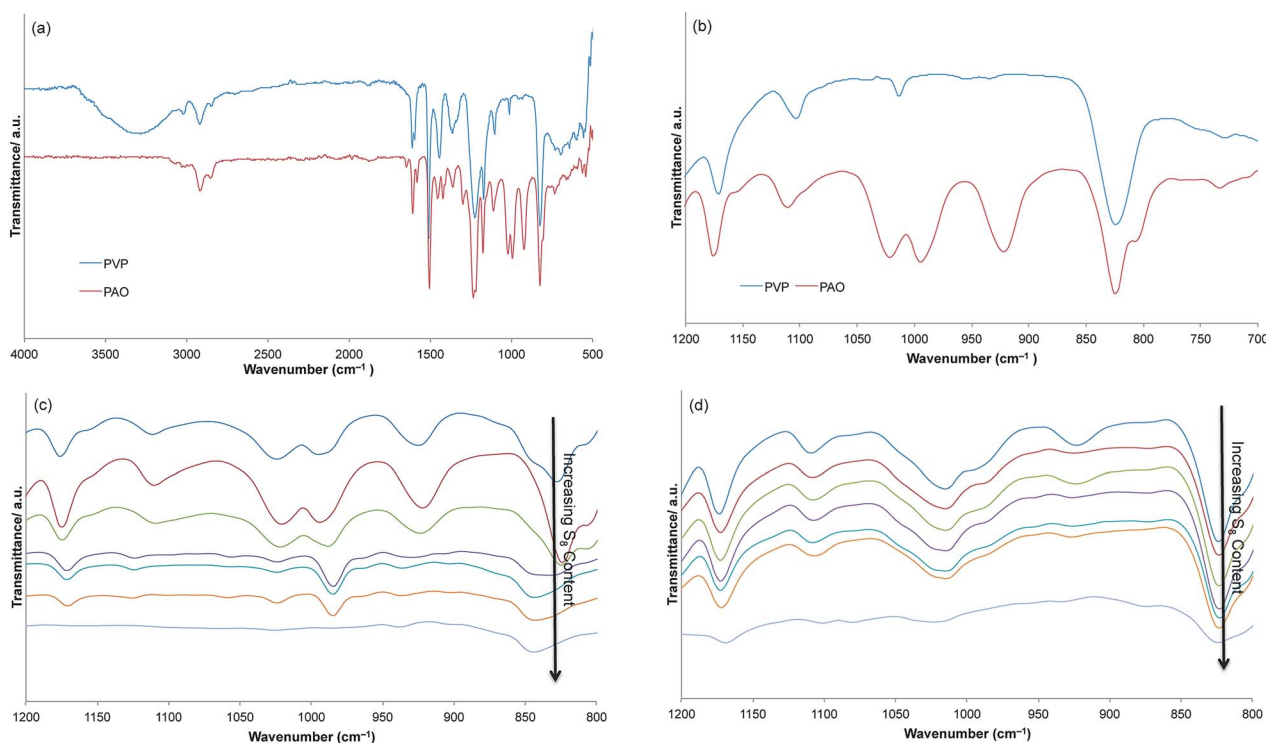


Fig. 1 FT-IR spectra for poly(4-vinylphenol) and **PAO** over the full scan range (a), poly(4-vinylphenol) and **PAO** highlighting the range for the monosubstituted alkene C–H bending mode (b), CS_2 -soluble fractions from $PAOS_x$ ($x = 40–99$, (c)), and CS_2 -insoluble fractions from $PAOS_x$ ($x = 40–99$, (d)).



Table 1 Summary of mass loss in synthesis, TGA, and DSC data

Material	T_d^a (°C)	T_g (°C)	T_m (°C)	$T_{\alpha-\beta}$ (°C)	% Mass loss ^b (±2%)
PVP	310	— ^c	— ^c	— ^c	NA
PAO	395	43.3	— ^c	— ^c	NA
S ₈	210	— ^c	118.7	— ^c	NA
PAOS ₉₉	203	— ^c	117.0	— ^c	— ^d
PAOS ₉₀	191	−37.0	115.8	— ^c	3
PAOS ₈₀	199	— ^c	116.5	105.3	4
PAOS ₇₀	192	— ^c	114.3	— ^c	4
PAOS ₆₀	195	— ^c	114.6	— ^c	3
PAOS ₅₀	238	— ^c	115.2	103.4	2
PAOS ₄₀	241	— ^c	116.6	105.8	2
PAOS ₃₀	244	— ^c	— ^c	— ^c	2
PAOS ₂₀	263	— ^c	— ^c	— ^c	2
PAOS ₁₀	290	— ^c	— ^c	— ^c	7

^a T_d was determined by calculating the temperature at which 5% mass was lost. ^b Mass loss was determined by weighing the mass of the product after the reaction. ^c No visible transition in the range studied. ^d Was not recorded.

the release of H₂S gas, as confirmed by elemental microanalysis. CAUTION: H₂S gas is foul-smelling, toxic, and corrosive and should be trapped as it is produced. Literature examples of H₂S-trapping strategies include zinc oxide-titanium dioxide sorbents,⁴¹ zinc oxide nanoparticles,⁴² reaction with Ag⁺ salts for precipitation as Ag₂S,⁴³ and other systems.⁴⁴ Analysis of PAOS_x by FT-IR spectroscopy revealed that not all C=C bonds in PAO had reacted with S₈, as peaks were still observed at 922 and 995 cm^{−1} in all samples (spectra are provided in the ESI†). Unfortunately, ¹H NMR spectrometric analysis could not be performed due to the poor solubility of PAOS_x in all solvents examined. Combustion analysis was performed for the elements C, H, N, and S for all PAOS_x, which demonstrated that S was efficiently incorporated into PAOS_x.

Given that unreacted C=C linkages remained in each PAOS_x NCC, we hypothesized that there might also be uncrosslinked S₈. Each NCC was therefore stirred with an excess of carbon disulfide (CS₂) to extract and enable quantification of free (non-cross-linked) sulfur. As anticipated, NCCs with higher sulfur contents ($x > 50$ wt%) comprised substantial amounts of CS₂-extractable mass. In contrast, PAOS_x materials containing less than 50 wt% sulfur afforded little to no CS₂-extractable mass, indicating that essentially all the sulfur is covalently incorporated in these cases. The CS₂-soluble and insoluble fractions resulting from extraction of PAOS_x ($x = 99, 90, 80, 70, 60, 50$, and 40) were isolated and then analyzed by FT-IR spectroscopy (Fig. 1d). FT-IR analysis suggested that nearly no organics are present in CS₂-soluble fractions for PAOS_x ($x > 50$). Elemental analysis on the CS₂-extractable fraction of PAOS₉₀ confirmed their identity as being >99% by mass sulfur. The CS₂-insoluble fraction from PAOS₉₀ contained 47% by mass of non-extractable sulfur and the FT-IR spectrum confirms nearly complete consumption of the alkene moiety in the polymer backbone. On the basis of the extent of alkene consumption and the amount of non-extractable (crosslinked) sulfur present, the average crosslink is comprised of five sulfur atoms.

Thermal stability

Thermogravimetric analysis (TGA) of PAOS_x revealed decomposition temperatures (T_d , here defined as the temperature at which 5% mass loss was observed upon heating under an atmosphere of N₂) ranging from 191–290 °C (Fig. 2a). For comparison, the T_d values for S₈ and PAO are 210 and 395 °C, respectively. For lower sulfur content PAOS_x ($x = 10$ –50%), T_d values generally decreased as the wt% of S₈ increased. However, samples with $x > 50$ % exhibited T_d values lower than that of S₈ and did not appear to vary significantly in response to changes in S₈ content. This depression of T_d is attributable to decomposition of an amount of non-crosslinked sulfur corresponding to the amount of CS₂-extractable sulfur (*vide supra*). Higher sulfur content PAOS_x ($x > 70$) exhibit very low char yields due to the clean decomposition of sulfur-rich domains, whereas materials containing <70 wt% sulfur – and comprised by

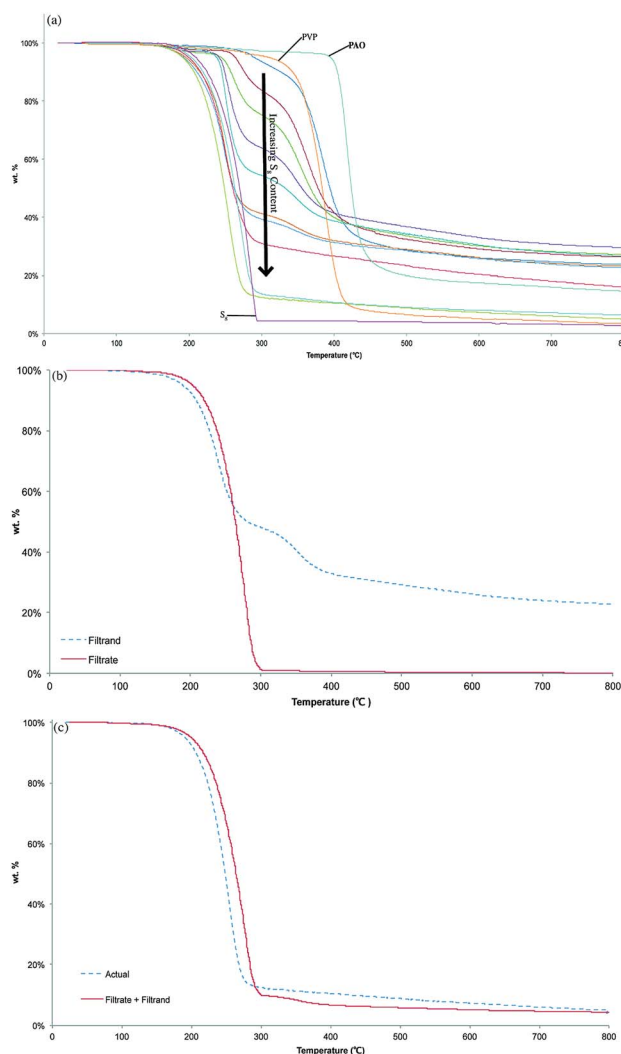


Fig. 2 (a) TGA curves of PVP, PAO, elemental sulfur, and PAOS_x. (b) The TGA curves for the CS₂-extractable (solid line) and CS₂-insoluble fraction (dashed line) of PAOS₉₀. (c) The calculated (solid line) TGA curve for independent thermal decomposition of the two fractions overlaid with the experimentally observed TGA curve (dashed line) from the PAOS₉₀ bulk material.



correspondingly little non-cross-linked sulfur – have char yields ranging from 20–28% (Fig. 2a).

The first degradation step observed for all **PAOS_x** materials was attributed to the loss of the extractable sulfur portion as H₂S gas, which can form *via* hydrodesulfurization reaction between S and the organic components. To confirm this hypothesis, a bulk sample of **PAOS₅₀** was subjected to heating at 250 °C, just above the observed *T_d* for this material. The vessel was equipped with a bubbler containing 1 M AgNO₃(aq.) to trap evolved H₂S gas *via* reaction with AgNO₃ to form insoluble Ag₂S. The mass of Ag₂S isolated from this experiment accounts for >83% of the mass loss observed in the TGA trace for **PAOS₅₀**. The shortfall of isolated Ag₂S is likely a result of gas escape prior to reaction or incomplete collection of the finely-dispersed solid. **PAOS_x** samples with *x* < 70 exhibit a second degradation step at roughly 290 °C attributable to degradation of the organic polymer backbone. The assertion that the covalently-linked and extractable sulfur domains undergo thermal decomposition independently even when commingled in **PAOS₉₀** is further supported by the observation that the curve resulting from summing mass-normalized TGA traces of CS₂-soluble and -insoluble fractions is nearly coincident with that of the TGA trace of the original sample of **PAOS₉₀** (Fig. 2C).

Morphology

Differential scanning calorimetry (DSC) data for fully cured samples of **PAOS_x** are summarized in Table 1. **PAO** is shown to have a *T_g* at 43.3 °C, while pure orthorhombic α -sulfur does not exhibit a glass transition in the range studied. Additionally, sulfur has a melt peak at 118.7 °C. The higher sulfur content **PAOS_x** (*x* = 60–99) exhibit endothermic features at ~106 °C. The feature at ~106 °C in DSC analysis is attributable to an orthorhombic to monoclinic phase change (*T_{α-β}*, for the α -S(s) → β -S(s) process).^{45–47} Older literature has variably attributed this transition to occurring at 95 °C,^{48–51} possibly confused by the significant variability in properties of sulfur based on its thermal history and complex phase behavior.^{36,49,52–55} The peak at ~115 °C is attributed to the melt peak of sulfur-rich domains in the materials. The normalized integral of the sulfur melt peak decreases predictably as the wt% of sulfur is decreased (Fig. S13–S24 in the ESI†), consistent with progressively lower amounts of non-cross-linked S in these materials. More interestingly, there is no apparent *T_g* in the measured temperature range for any of the **PAOS_x** materials except for **PAOS₉₀**. **PAOS₉₀** exhibits a weakly-observable *T_g* at –37.0 °C, a transition which has been suggested in amorphous sulfur as well.^{56–58} The lack of a comparable *T_g* in even more sulfur-rich **PAOS₉₉** suggests that a lower ratio of cross-linking of **PAO** may facilitate continuity of a more purely orthorhombic sulfur state. An alternate explanation is that **PAO** may act as a plasticizer for amorphous sulfur and stabilize the polymeric and orthorhombic domains in the NCC, consistent with the observed decrease in *T_g* for **PAOS₉₀** relative to amorphous sulfur (–30 °C). DSC data show that the lower wt% sulfur materials (**PAOS_{10–30}**) have neither a *T_g* nor a *T_m* in the temperature range studied. The predominating

absence of these thermal transitions or those attributable to either orthorhombic sulfur or polymeric sulfur further supports the formulation of **PAOS_x** as extensively cross-linked materials with oligosulfide chains separating organic polymer chains.

Mechanical properties

One of the primary objectives of the current study was to investigate the extent to which low loading of polymers can endow bulk sulfur with enhanced mechanical strength. On this basis, **PAOS₉₀** and **PAOS₉₉** were selected for study by dynamic mechanical analysis (DMA).

The responses of **PAOS₉₀** and **PAOS₉₉** to flexural stress were studied by DMA in dual cantilever mode with a 1 Hz frequency of oscillation from –60 to 88 °C. The thermal history of sulfur has a dramatic impact on its subsequent thermomechanical properties. To assess whether this would hold true of high sulfur-content NCCs, DMA was undertaken 1.5 h (Fig. 3a–c) and 96 h (Fig. 3d–f) of curing at room temperature after initial melt-casting of samples. The TGA and DSC data discussed in the previous sections is for samples that were cured for at least 96 h. One telling metric in assessing the curing process is the *T_g*. The *T_g* can be elucidated in many different ways from DMA data.⁵⁹ In the present case, the *T_g* was determined from the onset of storage modulus (*T_{g,SM}*) or as a local maximum in either loss modulus (*T_{g,LM}*) or tan δ curves (*T_{g,tan}*). For samples analysed 1.5 h after preparation, the onset of storage moduli (Fig. 3a) for **PAOS₉₀** and **PAOS₉₉** occur at the same temperature (–17 °C), however the magnitude of the storage modulus for **PAOS₉₀** is only 50% that of **PAOS₉₉**. The *T_g* calculated from loss moduli are –4.7 and –6.3 °C for **PAOS₉₀** and **PAOS₉₉**, respectively. The calculated *T_g* from the tan δ peaks are 2.3 °C for **PAOS₉₀** and 5.0 °C for **PAOS₉₉** (Fig. 3c). The magnitude of the tan δ peak provides evidence for effective damping by the material. The damping factor indicates how well a material can dissipate energy through segmental motions or relaxations. Fig. 3c shows that over the entire temperature range studied, **PAOS₉₉** has a higher tan δ value than **PAOS₉₀**. This suggests that **PAOS₉₉** exhibits far more viscous than elastic properties relative to **PAOS₉₀**. **PAOS₉₉** is thus able to dissipate energy more efficiently than its **PAOS₉₀** counterpart. Notably, the tan δ curve for **PAOS₉₉** increases from 60 to 88 °C, a result of the large amounts of free sulfur in the material. These insights further support the inference from DSC data that there is a greater predominance of orthorhombic sulfur in **PAOS₉₉** than in **PAOS₉₀**.

The 96 h-cured samples of **PAOS₉₀** and **PAOS₉₉** were subjected to testing by DMA under identical experimental conditions as were the 1.5 h-cured samples. For comparison, a 96 h-cured sample of pure sulfur is too brittle to be clamped in the DMA instrument even at minimum clamping force. The storage and loss moduli and tan δ curves are shown in Fig. 3d–f. All parameters exhibit modest shifts between 1.5 h and 96 h post-casting. In every case the calculated *T_g* increased over the 96 h time period. Table 2 summarizes the calculated *T_g* values for **PAOS₉₀** and **PAOS₉₉** after 1.5 and 96 h post-casting. In both **PAOS_x** materials, the storage modulus is seen to increase over the 96 h period, whereas, the tan δ values for both **PAOS_x** materials are shown to decrease over the entire temperature range. The increase in



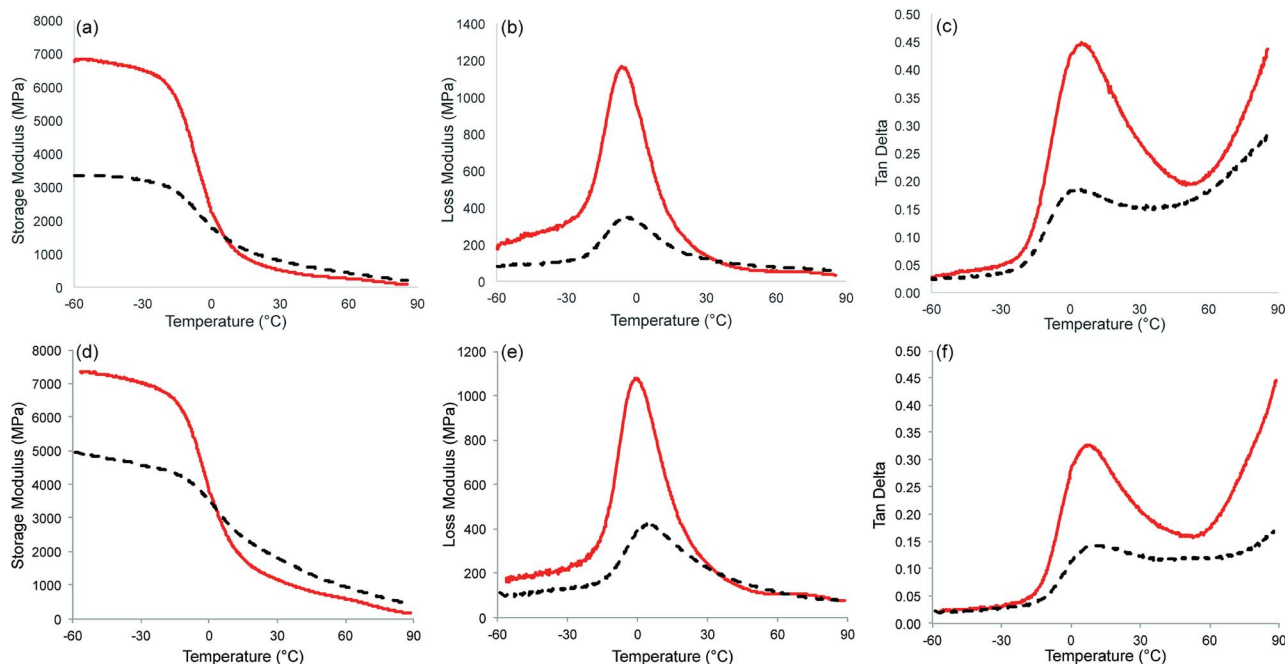


Fig. 3 Temperature dependence of storage and loss modulus and $\tan \delta$ of PAOS₉₀ (dashed line) and PAOS₉₉ (solid line) in dual cantilever mode after 1.5 h of cooling (graphs a–c) and 96 h of cooling (graphs d–f).

Table 2 T_g values for PAOS₉₀ and PAOS₉₉ determined from DMA data

Material	$T_{g,SM}$ (°C)		$T_{g,LM}$ (°C)		$T_{g,tan}$ (°C)	
	1.5 h	96 h	1.5 h	96 h	1.5 h	96 h
PAOS ₉₀	−17	−10	−4.7	5.7	2.3	9.7
PAOS ₉₉	−17	−13	−6.3	−0.5	5.0	7.3

storage modulus is attributed to the slow reversion of free polymeric sulfur back to the thermodynamically-favored, more crystalline orthorhombic S₈.⁶⁰ Moreover, the decrease in magnitude of $\tan \delta$ is an indication that the materials' elastic behaviour concomitantly increases.

Thermal healing and materials comparison

One driving force for the current work was that the thermal reversibility of S–S bond formation offers intriguing possibilities for thermal healing and recycling of high sulfur-content materials. To test the thermal healing/recyclability of PAOS₉₉, a sample of the material was subject to DMA analysis and then pulverized into small pieces prior to reannealing to heal the material, and finally allowing the sample to cool to room temperature over 15 min. After repeating this pulverization-thermal healing process ten times, the mechanical properties of the twelve-times-healed sample were identical to that of the first sample within error of the measurement (Fig. 4).

Another driving force behind this study was to determine whether crosslinking by small amounts of organic material could improve the mechanical integrity of bulk sulfur to levels that make it viable for practical applications: without additives

to stabilize it, polymeric sulfur is unstable, reverting to the S₈ form at STP. Comparison of the mechanical features of polymer-cross-linked high-sulfur content PAOS_{90–99} to other small molecule-cross-linked, high-sulfur content materials is also of interest. Pyun's group has prepared 1,3,5-triisopropenylbenzene/sulfur crosslinked materials.⁶¹ The storage modulus of this small molecularly-cross-linked material comprising 30 wt% 1,3,5-triisopropenylbenzene and 70 wt% sulfur was 1 GPa at 30 °C under a shear stress. Polymer-cross-linked PAOS₉₀ and PAOS₉₉ exhibit a notably higher storage modulus of approximately 2.6 GPa at 30 °C under a flexural stress despite the significantly higher sulfur content in these NCCs.

Compared to other reversibly-crosslinked/thermally healable polymers, PAOS₉₉ has a higher flexural storage modulus than for epoxy shape-memory polymers⁶² and comparable to that of crosslinked and thermally healable (*via* Diels–Alder reaction) poly(dicyclopentadiene)s.⁶³ The improved mechanical strength

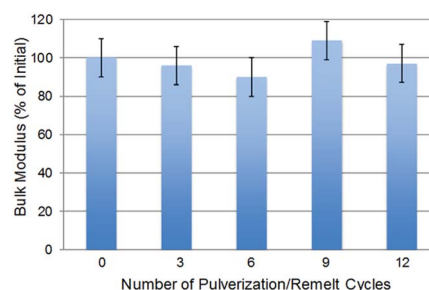


Fig. 4 The bulk modulus (measured at −45 °C) for PAOS₉₉ after pulverization/remelting cycles remains consistent within error for at least twelve cycles.



of polymer-crosslinked **PAOS_x** over other sulfur-rich materials and the similarity in room-temperature storage modulus to polystyrene (1.8 GPa at 30 °C)⁶⁴ provide sound proof-of-principle to justify further efforts to prepare value-added materials from affordable/plentiful by-product sulfur by the strategy employed in this study.

Conclusions

A commercially-available polystyrene derivative was modified to facilitate cross-linking with elemental sulfur. The crosslinked materials, **PAOS_x** were prepared with sulfur content of 10–99% by mass. The thermal and mechanical properties of **PAOS_x** were elucidated. Target high sulfur-content **PAOS₉₉** and **PAOS₉₀** were comprised of a mixture of sulfur trapped in the amorphous state and short-chain sulfur crosslinks between organic polymer chains. The storage moduli of both **PAOS₉₀** and **PAOS₉₉** increased over a 96 h setting time as free polymeric sulfur slowly relaxed to the orthorhombic state. The fully-set mechanical strength of even **PAOS₉₉**, which is 99% by mass sulfur, increased significantly when compared to a pure sulfur sample prepared under the same conditions. **PAOS₉₉** maintains its mechanical integrity after physical breakdown and thermally recasting for at least a dozen cycles, demonstrating the facile recyclability of these materials. **PAOS₉₀** and **PAOS₉₉** materials thus show promise for valorizing elemental sulfur as a component of durable, recyclable material applications for which petrochemicals are currently used. Follow-up studies on improving the mechanical properties of high sulfur-content network solids are currently underway.

Conflicts of interest

There are no conflicts to declare.

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