Dark sulfur: Quantifying unpolymerized sulfur in inverse vulcanized polymers

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ABSTRACT: Inverse vulcanization allows polymeric materials to be formed from excess elemental sulfur, promoting polysulfide chains to be stabilised between organic comonomers, rather than reforming as crystalline S8, resulting in high sulfur content materials with interesting properties. The techniques used to determine if free unreacted sulfur remains in the polymers only detect the crystalline, and not the amorphous form. A detailed study is presented on the quantification of free amorphous sulfur within inverse vulcanized polymers, in which free sulfur is shown to increase over a period of aging. Post-aging regeneration by stimulating homolytic S-S cleavage in the polymer, is investigated.

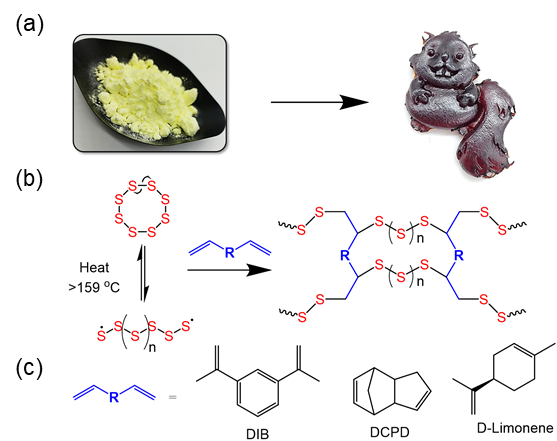
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Sulfur is a naturally occurring element that has been used by humankind for hundreds of years. Records of its use reach back to Roman times, in Pliny the Elder’s discussions of minerology in his Naturalis Historia. 1 Elemental sulfur has been employed for its antibacterial properties, treating ailments such as acne vulgaris, dermatitis, and warts. 2, 3 Indeed, Pliny refers (without evidence) to sulfurs use in the treatment of leprosy, asthma, and scorpion stings. In modern times however, the biggest use of sulfur is in the production of sulfuric acid, notably in the contact process. 4, 5 Sulfur itself is today produced mainly as a by-product of crude oil purification in the manufacture of petrochemicals.6 However, this process yields large quantities of sulfur in excess of that for which uses can currently be found. It is therefore desirable to identify new applications for sulfur to prevent the increasing build-up of waste, and in this regard Inverse Vulcanized Polymers may provide a solution. The seminal publication by Chung *et al*. introduced the process of Inverse Vulcanization7 in which sulfur is co-polymerized with a diene crosslinker to yield a polymeric product of high sulfur content (>50 wt.%). Not only would utilization of excess sulfur be beneficial, but these novel materials possess a variety of properties, with demonstrable benefits in terms of antibacterial capacity,8, 9 optics,10 mercury capture,11, 12 and as encapsulating agents for fertilizer components. 13 Significant interest has also attached to the use of these polymers in Li-S batteries14 due to their theoretical high energy storage capabilities (1762 mAhg-1). 15 The different properties of sulfur-based polymer materials, including self-repair,16 high specific capacity,17 and tunable glass transition temperatures (*T*g, the temperature above which the polymer transitions from a glass-like to an amorphous state),18 make them very attractive for a range of applications.

The variation in properties of these polymers can result from changes in synthesis conditions, sulfur loading, or crosslinker identity.18 Petrochemically derived molecules such as 1,3-diisopropenyl benzene (DIB),7 dicyclopentadiene (DCPD),19 and methyl styrene20 provide a range of crosslinked and linear polymers with highly varied physical characteristics. Such is the potential for variety in viable crosslinkers that renewable oils and plant-based derivatives such as canola oil21 or limonene22 can be used, enhancing the green credentials of this process.

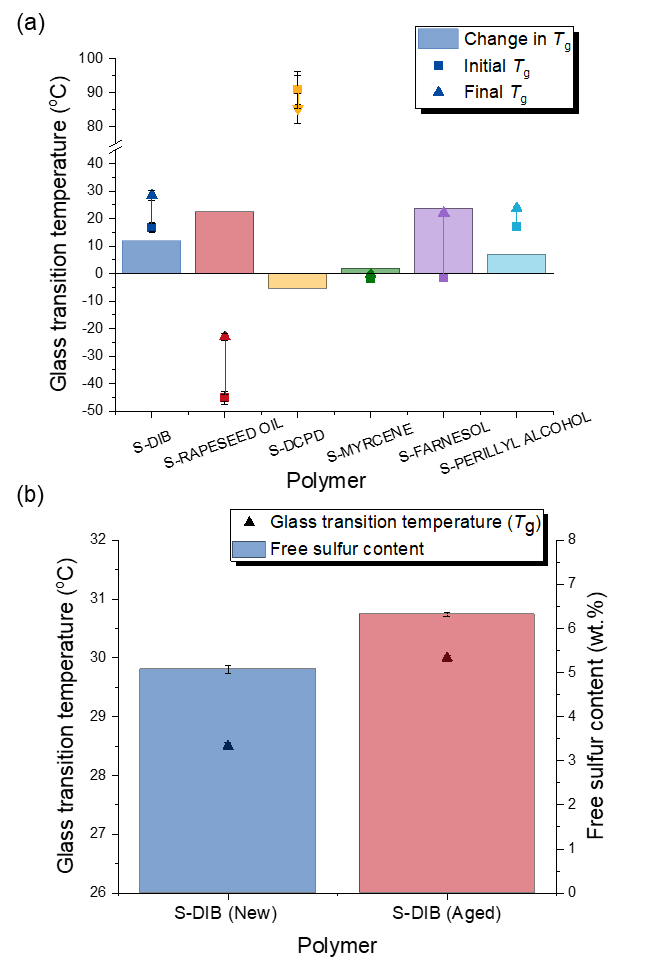
Polymer aging is a major issue,23 and increasing interest in the development of deliberately purposed sulfur-based polymers necessitates improved understanding of the mechanisms of this aging. The *T*g of a polymer is a key characteristic affecting properties such as mechanical strength and brittleness.24 While it has been shown that the *T*g of these polymers is variable owing to reasons such as difference in crosslinker identity and synthesis conditions, it is shown herein that this polymer property is subject to aging over time.

Much of the previously published work has assumed that the measure of a successful polymerisation, that is to say one in which all sulfur is polymerised and no free sulfur remains in the polymer, could be characterised by powder x-ray diffraction (PXRD) or differential scanning calorimetry (DSC), concluding the absence of crystalline S8. 25, 26 Whilst non-crystalline free sulfur has previously been observed in inverse vulcanized polymers by means of Raman spectroscopy, it has never been fully analysed or quantified.27 In order to quantify this free sulfur a high-performance liquid chromatography (HPLC) method has been developed allowing free sulfur to be extracted from the polymer. Here is reported an investigation of the aging, and unpolymerized amorphous S8 content, of a range of inverse vulcanized polymers, as well as the potential to regenerate their properties. This sulfur species is referred to as ‘dark sulfur’ – amorphous sulfur that is not observable by preestablished analytical techniques, and not stabilised within the polymer structure between two organic moieties. While it is not clear what form this amorphous unreacted sulfur takes, it is predicted to exist as a variety of variants, including polysulfides of different lengths and radicals, as previously reported by Wadi *et al.* 28Therefore it is defined as any sulfur, not covalently bonded to an organic moiety, and is non-crystalline and thus undetectable by the previously reported methods of DSC and PXRD.

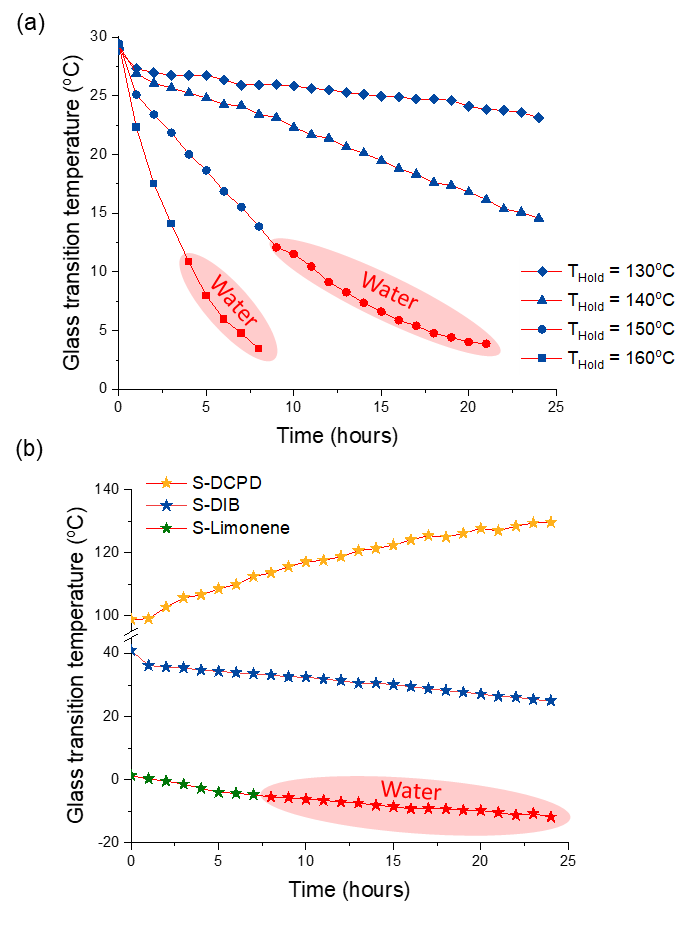
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**Figure 1.** (a) The conversion of elemental sulfur to a poly-S-crosslinker polymer material (b) The reaction mechanism of elemental sulfur reacting with a generic crosslinker molecule to form a sulfur based polymer via inverse vulcanization (c) The chemical structures of 1,3-diisopropenyl benzene (DIB), dicyclopentadiene (DCPD), and D-Limonene, the 3 main crosslinkers considered in this work.

Sulfur polymers were synthesised by the reaction of S8 and organic comonomer in equal mass at 170 °C under agitation at 600 rpm. Upon reaching homogeneity, this polymer solution was oven cured at 140 °C for 24 hours. See SI (Figure S1) for more details. During our investigations into polymer aging, it was observed that the *T*g of the assessed polymers increased over time, except for S-DCPD (Figure 1). No loss of mass was noted, nor an increase in crystalline sulfur (measured by DSC) unless crystalline sulfur was present upon the commencement of aging (Figure S2-6). After aging for six months (with changes in room temperature monitored for 30 days during this period (Figure S7)), all polymers remained insoluble in toluene, with no alterations in the infrared spectra (FTIR) suggesting that no structural change in the organic moieties was taking place, or the formation of such species as thiols (Figure S8-10). This suggests that there may be a change in sulfur rank over time. The sulfur rank (the number of sulfur atoms per chain between organic crosslinker sections) may be variable across the polymer upon synthesis, and consists of a number of weaker S-S linkages end capped by stronger C-S bonds. The longer the sulfur chain length, the weaker the S-S linkage towards the centre of the chain. Over time these weak bonds in the middle of the chain may naturally cleave, liberating sulfur into the polymer matrix. The polymer chain may then reform, thus maintaining the degree of crosslinking.

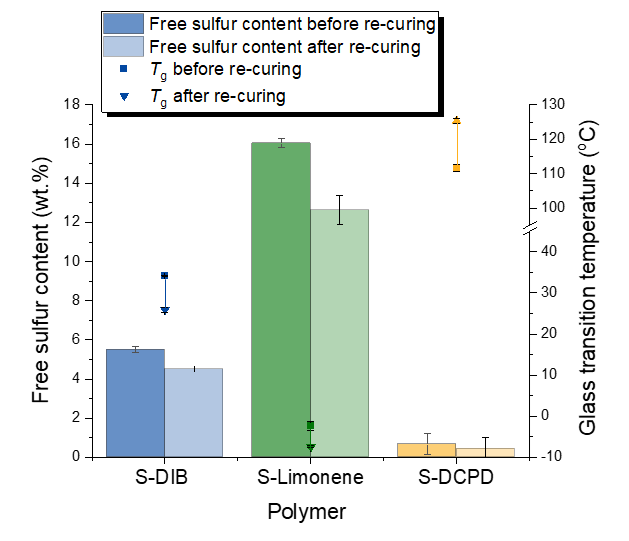
**Figure 2.** (a) The change in *T*g of a range of polymers, selected for having a range of initial values of *T*g. Polymers were synthesised with a sulfur loading of 50 wt.% to 50 wt.% comonomer. Aging time = 6 months (b) A comparison between aged and new S-DIB, showing an increase in both *T*g and dark sulfur content (from HPLC). Aging time = 4 months.

This reformation leads to the observed increase in *T*g, as the linear sections become shorter. To confirm the existence of this amorphous sulfur, polymer samples were analysed by thin layer chromatography (TLC) and HPLC. Solvent extraction of free sulfur was carried out in toluene, with solutions at concentrations of 5 mg mL-1 prepared, and passed through 25 mm 0.45 µm nylon syringe filters. TLC analysis was carried out in a hexane mobile phase. This suggested that, while no free crosslinker was present in the polymer samples, unpolymerized sulfur was present in both new and aged polymers, despite no crystalline sulfur having been observed in the DSC analysis (Figure S11) A method for the analysis of elemental sulfur by HPLC was then developed, adapted from the method detailed by Buchanan *et al*. for the analysis of free sulfur in coal.29 Elemental sulfur in the form of S8 is eluted from the column at a residence time of 7.3 minutes (Figure S12-13). It should be noted that the temperature increase in the HPLC analysis may lead to the formation of further polysulfides, so that if oligomeric species were extracted into the solvent with the dark sulfur the quantity of sulfur observed may be slightly greater than the real value. However, this will only have a minor impact on the obtained value. Comparison of both aged and new S-DIB confirmed an increase in the quantity of free sulfur of 1.2 wt.% in a polymer with a sulfur loading of 50 wt.% after a period of 4 months (Figure 2). Analysis of the new S-DIB was conducted immediately after synthesis.

**Figure 3.** The change in *T*g with re-curing of (a) an aged sample of S-DIB at hold temperatures of 130-160 °C (b) newly synthesised samples of S-DCPD, S-DIB, and S-Limonene at a hold temperature of 140 °C. Water appears in the DSC traces of certain samples and conditions, suggesting the polymer chains are mobile enough to allow the release of trapped moisture.

This coincides with an increase in glass transition temperature of 1.5 °C over this time. This contributes to the suggestion that sulfur may be liberated from the sulfur chains, shortening the sulfur rank and thus increasing the *T*g. This conclusion regarding polymer aging has significance for the development and consideration of inverse vulcanized polymers, and further research is being carried out to fully understand how aging may affect polymer properties and application performance.

It may be possible to reverse this aging due to the dynamic nature of the S-S bonds in the longer polymer chains. The application of heat may allow for homolytic S-S cleavage and subsequent re-insertion of this ‘dark’ sulfur into the polymer chains. This would result in a predicted decrease in the *T*g and amorphous sulfur content. An experiment was therefore designed using DSC, whereby 5 mg of a polymer was re-cured by heating to a designated temperature and held for 1 hour inside the DSC, characterised by temperature sweep, and then similarly cured further. This was repeated over a 25 hour period (the first hour allowing the sample to reach thermal equilibrium). It was observed that upon heating the *T*g decreased linearly, with higher hold temperatures yielding a faster rate of *T*g decrease (Figure 3).

**Figure 4.** The change in *T*g and free sulfur contents of newly synthesised S-DIB, S-Limonene, and S-DCPD after re-curing for 24 hours at 140 °C in sealed containers.

As it was observed that new samples of S-DIB also contain dark sulfur, this experiment was repeated for newly synthesised samples of S-DCPD, S-DIB, and S-Limonene, chosen to represent polymers of a wide range of initial *T*g and sulfur rank. S-DIB and S-Limonene showed a decrease in *T*g suggesting the reinsertion of sulfur into the polymer chains. S-DCPD showed an increase in *T*g. The average estimated sulfur rank of S-DCPD is 1.94, thus suggesting the polymer is made up of a series of strong disulfide linkages. The strength of these disulfides is such that the bonds may not homolytically cleave, therefore preventing the structural rearrangement of the polymer, and elongation of the polymer chains. It is predicted that S-DIB30 and S-Limonene contain longer polysulfide chains with weaker S-S bonds that may cleave under applied heat, allowing for rearrangement (Figure S14). It must also be noted that water began to accumulate in the DSC traces after a period of time, obscuring the *T*g of some samples. It is posited that this water originates from within the polymer, where it has accumulated over the course of the aging period from adventitious moisture (Figure S15-16). The application of heat allows for increased mobility of the elongating polymer chains and the water to escape the polymer. Water appears in the new sample of S-Limonene as it has a *T*g lower than room temperature, therefore existing in a form where the polymer chains are more mobile, allowing for the entrance of moisture rapidly into the sample. Further investigations into this water would be greatly beneficial, as this may link to sulfur blooming, a common problem observed in many sulfur containing materials.

To analyse a change in dark sulfur with re-curing, a series of polymers of S-DCPD, S-DIB, and S-Limonene were held in an oven at 140 °C for 24 hours. After this period of re-curing the *T*g of all polymers increased (Figure S17). As the polymers were open to the air, it was possible for the unpolymerized sulfur to migrate to the surface and escape the polymer, further evidenced by the appearance of sulfur crystals in the extraction vents of the oven. The polymers were then able to cure further and rearrange into a more crosslinked structure. This investigation was repeated with the polymer samples contained in sealed DSC pans. It was subsequently observed that the *T*g of the sealed polymers decreased with a decrease in dark sulfur (figure 4), in agreement with the data presented in Figure 3. Once again S-DCPD shows an increase in *T*g, correlating with further curing of the polymer and the inability to break the sulfur chain linkages. Little difference in re-curing was observed with alterations in gas flow over the sample, thus confirming the re-curing effect is derived from the application of heat (Figure S18). This data suggests that inverse vulcanized polymers may boast regenerative capabilities, with the potential to reverse the aging that may occur during the polymer lifetime.

In summary, it has been determined that the aging of inverse vulcanized polymers is not simply limited to changes observable by the eye. This investigation into polymer aging has confirmed the existence of free amorphous S8 in both new and aged polymers showing that, even in the absence of crystalline sulfur, a polymerization may not be considered as complete. There is a possibility that this sulfur may be incorporated back into the polymer chains through the application of heat, in a sealed system, above the melting point of sulfur. This free sulfur was quantified by HPLC, though the authors accept the limitations of this method, such as the possibility that the solvent extraction method may itself affect the polymers. Independent analyses and further research are currently being conducted to assess the effects of dark sulfur on polymer properties, whether there is any notable benefit e.g. as a reservoir for sulfur release in various applications such as fertilisers and for antimicrobial purposes or detriment e.g. enhanced sulfur blooming of dark sulfur, and determine other methods of analysis. We suggest this area of discovery may benefit from further investigation into other analytical methods for analysis of amorphous sulfur, ideally in-situ.

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Conflict of Interest Disclosure Statement

The PhD student, Joseph Dale, who undertook the majority of this work is sponsored by the company Arkema, which has been specified in the acknowledgements.

Supporting Information

Sample preparation, DSC, storage temperature analysis, FTIR, TLC, HPLC, predicted average sulfur ranks of polymers. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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