Modelling mercury sorption of a polysulfide coating made from sulfur and limonene

Max J. H. Worthington, Ismi Yusrina Muhti, Maximilian Mann, Zhongfan Jia, Anthony D. Miller, and Justin M. Chalker

a) Institute for Nanoscale Science and Technology, College of Science and Engineering, Flinders University, Bedford Park, South Australia 5042, Australia

b) College of Science and Engineering, Flinders University, Bedford Park, South Australia 5042, Australia

To whom correspondence should be addressed:
tony.miller@flinders.edu.au
justin.chalker@flinders.edu.au

Keywords: mercury, polysulfide, remediation, sorbent, sulfur

Abstract: A polymer made from sulfur and limonene was used to coat silica gel and then evaluated as a mercury sorbent. A kinetic model of mercury uptake was established for a range of pH values and concentrations of sodium chloride. Mercury uptake was generally rapid from pH = 3 to pH = 11. At neutral pH, the sorbent (500 mg with a 10:1 ratio of silica to polymer) could remove 90% of mercury within one minute from a 100 mL solution 5 ppm in HgCl₂ and 99% over 5 minutes. It was found that sodium chloride, at concentrations comparable to seawater, dramatically reduced mercury uptake rates and capacity. It was also found that the spent sorbent was stable in acidic and neutral media, but degraded at pH 11 which led to mercury leaching. These results help define the conditions under which the sorbent could be used, which is an important advance for using this material in remediation processes.

Statement of novelty: Previous studies of the featured mercury sorbent did not detail the scope and limitations at varying pH and salt concentrations, which are critical to know in remediation projects. This is the first study of the effects of pH and sodium chloride on the rate of mercury uptake by a polymer made from sulfur and limonene. Additionally, the first kinetic model of mercury uptake was established for this material. Finally, leaching experiments under different conditions were completed for the first time, establishing that the sorbent is stable at low and neutral pH, but degrades at pH 11 and releases bound mercury from the bulk sorbent.

Graphical abstract

Renewable feedstock

(catalyst) 140 °C

Dry load on silica

[S]ₘ

Solvent processible mercury sorbent

Free-flowing mercury sorbent

Rapid mercury removal from water
1.1 INTRODUCTION

Mercury is a toxic heavy metal encountered in a variety of industrial sectors such as coal combustion, oil and gas refining, and artisanal and small-scale gold mining.\textsuperscript{1, 2} Mercury pollution in aquatic systems is especially problematic, as it can contaminate the food supply through bioaccumulation in seafood, or lead to contaminated drinking water.\textsuperscript{3, 4} Mercury remediation technologies for purifying water are therefore important, but this is challenging because of the high rate of mercury uptake required for filtration applications. High throughput is also important where large volumes of water need to be treated. To meet this need, a number of sorbent technologies have been evaluated including activated carbon, biochar, zeolites, polymers, metal covalent frameworks, and many other nanostructured or functionalised materials.\textsuperscript{5} Among the many classes of mercury sorbents, those functionalised with sulfur are of particular interest due to the strong affinity of sulfur for mercury.\textsuperscript{6} It is perhaps not surprising that there have been many studies on sulfur-functionalised sorbents for mercury remediation.\textsuperscript{7} Among these materials, those sorbents made directly from sulfur are of particular interest, due to the low cost and industrial surplus of elemental sulfur.\textsuperscript{7, 9} Additionally, there has been a resurgence in methods to make polymers from sulfur,\textsuperscript{10-14} which has created many opportunities for using these low-cost and scalable polymers in mercury and heavy metal remediation.\textsuperscript{7, 15-26}

In this study, we investigated one of these sulfur polymers made by direct copolymerisation of elemental sulfur with the renewable terpene limonene. Our lab first reported the synthesis and use of this material in mercury remediation in 2015,\textsuperscript{27, 28} describing its advantageous features. For instance, both sulfur and limonene are low-cost, highly abundant feedstocks produced by the petroleum and citrus industries, respectively. The copolymer made from these building blocks—referred to here as poly(S-r-limonene)—is a low molecular weight oligomer that is fully soluble in organic solvents.\textsuperscript{28} This solubility is useful, for instance, in using the polymer to coat surfaces.\textsuperscript{28, 29} Additionally, at high concentrations of inorganic mercury, the polymer changes colour from red to yellow—a chromogenic response with potential use in mercury sensing or for filtration media that changes colour when it needs to be replaced.\textsuperscript{28} Building upon these foundational studies, the Hasell lab expanded access to this polymer and its utility in mercury remediation. For instance, the addition of an accelerator or catalyst in the synthesis allowed the material to be made at a lower temperature, which reduced by-product formation and increased the safety profile of the synthesis.\textsuperscript{29} Furthermore, Hasell also demonstrated that poly(S-r-limonene), when coated on a silica gel support, is effective at removing mercury from water.\textsuperscript{29} However, these studies were focused more on the poly(S-r-limonene) synthesis rather than mercury sorption, so there is a need to understand the detailed kinetics of mercury sorption and the scope of conditions under which it is effective. Herein, we report our findings on the first detailed kinetic analysis and modelling of this sorbent in both Hg\textsuperscript{2+} uptake and desorption. We also studied the influence of pH and sodium chloride concentrations on mercury sorption, which provided an increased understanding of the scope and limitations of this sorbent. These findings helped establish important guidelines for deploying this unique mercury sorbent in remediation.

1.2 MATERIALS AND METHODS

Additional experimental details, characterization data, and modelling details are provided in the Supporting Information.

1.2.1 Synthesis of poly(S-r-limonene)

D-Limonene (20.00 g, 146.8 mmol) and zinc diethyldithiocarbamate (2.00 g, 5.53 mmol) were added to a 100 mL round bottom flask and heated to 140 °C with stirring. Sulfur (20.00 g, 625.0 mmol) was added over 5 minutes to the reaction mixture. After the addition of sulfur, a heat gun was used to melt any sulfur adhering to the walls of the flask, returning the reagent to the reaction mixture. Over the course of 30 minutes, the reaction appeared to form one phase and changed from orange to dark brown. The reaction was continued for a period of 7 days. To isolate the product, the material was poured from the flask while hot, and then allowed to cool and solidify in a silicone mould. Yields were typically >95% based on mass balance of the starting materials and product.
1.2.2 Coating silica gel with poly(S-r-limonene)

The poly(S-r-limonene) material (13.89 g) was dissolved in 300 mL of dichloromethane and the solution was poured onto 138.9 g silica gel in a 1000 mL round bottom flask. The solvent was then removed by rotary evaporation (40 °C, 800 mbar). In cases where the silica gel adhered together, additional dichloromethane was added and the dry loading repeated to ensure even coverage on the silica. Finally, the silica was dried under high vacuum to remove any remaining solvent. The coated silica was isolated as an orange, free-flowing powder (150 g).

1.2.3 Preliminary assessment of mercury sorption

An aqueous solution of 5 ppm HgCl$_2$ (100 mL) was added to a 250 mL round bottom flask containing 500 mg of silica gel coated with poly(S-r-limonene). The sorbent was prepared as described above and contained a 10:1 mass ratio of silica to the poly(S-r-limonene) coating. The mixture was stirred and 1.00 mL samples of the solution were taken by pipette at 1, 2, 5, 10, 15, and 30 minutes. Any trace solids were removed by centrifugation, and the samples were diluted 10-fold with 5% HCl to stabilise mercury species for subsequent analysis. The experiment was repeated in triplicate. The experiment was also repeated with uncoated silica gel as a control. Mercury concentrations were then determined by cold vapour atomic absorption spectroscopic analysis (CVAAS).

1.2.4 General protocol for kinetic analysis of mercury sorption

Silica gel coated with poly(S-r-limonene) (500 mg, 10:1 silica:poly(S-r-limonene)) was added to 250 mL beaker and mixed with magnetic stirring. Next, an aqueous solution of 5 ppm HgCl$_2$ (100 mL) was added. This solution was prepared at various pH values and sodium chloride concentrations, as described below (1.2.5 and 1.2.6). The solution was sampled over time by drawing 1.00 mL of the solution into a 3 mL syringe equipped with a syringe filter (nylon, 0.45 µm). In this way, the solution was separated from the sorbent during sampling. Samples (1.00 mL) were taken every 10 seconds for the first minute and then at 90, 120, 180, and 300 seconds of total sorption time. The experiment was completed in triplicate. A control experiment was done in which 450 mg of uncoated silica was used as the sorbent. All samples were diluted 10-fold with 5% HCl and then mercury concentrations were determined by CVAAS.

1.2.5 Mercury sorption at varying pH values

Aqueous solutions of 5 ppm HgCl$_2$ were prepared at pH values of 3, 5, 9, and 11. The solution at pH = 3 contained 1 mM HCl, the solution at pH = 5 contained 10 µm HCl, the solution at pH = 9 contained 10 µm NaOH, and the solution at pH = 11 contained 1 mM NaOH. A 5 ppm solution of HgCl$_2$ was also used without adjusting the pH; this sample is referred to as the neutral sample (pH = 6.99). The mercury solutions were added to the silica gel coated with poly(S-r-limonene) and sampled and analyzed by CVAAS according to the general protocol described in 1.2.4.

1.2.6 Mercury sorption in the presence of NaCl

Aqueous solutions of 5 ppm HgCl$_2$ and sodium chloride were prepared. The final concentration of sodium chloride was 6.85 mM or 599 mM. These concentrations of sodium chloride are designed to mimic the salt concentrations in tap water and seawater, respectively. Both solutions were used in mercury sorption experiments following the general protocol for kinetic analysis described in 1.2.4.

1.2.7 Mercury desorption assessment

In a 500 mL beaker, a 20 ppm solution of HgCl$_2$ (200 mL) was added along with silica gel coated with poly(S-r-limonene) (4.00 g). The sorbent was prepared as described above and contains a 10:1 mass ratio of silica to the poly(S-r-limonene) coating. The mixture was stirred for 10 minutes and then the sorbent was isolated by filtration and dried under vacuum before splitting up into 500 mg portions. The sorbent samples, bound to mercury, were then added to 50 mL centrifuge tubes, followed by 50 mL of solutions of varying pH or sodium chloride concentrations (pH of 3, 5, 7, 9, 11, or aqueous solutions of 6.85 mM or 599 mM NaCl, prepared as described in
the sorption experiments). The mixtures were rotated on an end-over-end mixer and sampled at 10 minutes, 3 hours, and then 1, 2, 8, 14, 21, and 28 days. All samples were diluted 10-fold with 5% HCl and then mercury concentrations were determined by CVAAS.

1.2.8 Mercury sorption modelling
Two reaction models were used to analyse the kinetics of the Hg sorption experiments:

1.2.8.1 Single reaction sorption model
Let X denote an aqueous species and let S be an available adsorption site on the surface of the coating. Consider the conceptual reaction model

\[ X[x] + S[s] \rightarrow XS[y] \] (1)

here XS denotes the adsorbed state. The symbols in square brackets [ ] denote the volume or surface concentrations, as appropriate, of the relevant species. In this model any reverse reaction has been assumed extremely slow and so it is neglected. Since this is modelled as an irreversible reaction, it will continue to completion until one of X or S, or both are fully consumed. In what follows X can be thought of as Hg\(^{2+}\) or HgCl\(^2\).

A reaction rate equation for (1) can be written

\[ \frac{dy}{dt} = k_1 x s. \] (2)

If \(x\) is expressed in molar units (M) then \(k_1\) will have units of M\(^{-1}\)s\(^{-1}\).

This rate equation is motivated by the following physical reasoning. The product term \(xs\) is a measure of the likelihood of an entity of X and an entity of S coming into sufficiently close contact that the reaction (1) is possible, while \(k_1\) is a proportionality factor that also expresses the likelihood that a close contact will actually result in the formation of the adsorbed complex XS. It is to be expected that \(k_1\) will depend on a number of factors, in particular, the local chemical environment and temperature. The presence of other species, although they may not have any obvious direct involvement in the reaction, may give rise to various forms of interactions, for example crowding, shielding, attraction or repulsion, which can influence the likelihood of the reaction (1) occurring and so will affect \(k_1\). Likewise, the presence of other nearby entities of X and S may also give rise to interactions and so influence whether a close contact leads to a reaction. So, \(x\) and \(s\) may affect the overall reaction rate not just through the product term \(xs\), but also through \(k_1\). However, it is assumed that \(k_1\) does not change significantly during the course of the reactions considered here and consequently \(k_1\) can be regarded as approximately constant. This constant may, however, have different values under different chemical conditions, such as widely different pH and NaCl concentrations as are investigated in the study.

1.2.8.2 Two-reaction sorption model with competing reactions
Let W be another aqueous species that potentially competes with X for adsorption sites S. This gives the two-equation conceptual model

\[ X[x] + S[s] \rightarrow XS[y] \]
\[ W[w] + S[s] \rightarrow WS[yz] \]

Again, reverse reactions have been neglected. These reactions will proceed to completion until either all of S is consumed, or all of both X and W are consumed. In the case that all of S is consumed then there will be some residual limiting presence of X or W, or both. In what follows W will be either NaOH or NaCl.
Corresponding to (2) above is the set of two rate equations
\[ \frac{dy}{dt} = k_1 x s \]
\[ \frac{dz}{dt} = k_2 ws \]

The same comments as above concerning the approximate constant value of \( k_1 \) apply here for \( k_1 \) and \( k_2 \) also. Supposing that both \( x \) and \( w \) are measured in units of M, the units of both \( k_1 \) and \( k_2 \) are M\(^{-1}\) s\(^{-1}\).

1.2.9 Desorption reaction model

Any reverse reaction has been neglected in the above two sorption models. Over the longer term, however, desorption may take place to some degree. This can be accounted for by allowing the above reaction to be reversible. For the two-reaction model this would give
\[ X[x] + S[s] \rightleftharpoons XS[y] \]
\[ W[w] + S[s] \rightleftharpoons WS[yz] \]

with corresponding rate equations
\[ \frac{dy}{dt} = k_1 x s - k_{1R} y \]
\[ \frac{dz}{dt} = k_2 ws - k_{2R} z \]

Here the reverse rate coefficients \( k_{1R} \) and \( k_{2R} \) have units of s\(^{-1}\) or equivalent. Just as for \( k_1 \) and \( k_2 \), they will depend upon the chemical environment but are assumed approximately constant in any given situation.

1.2.10 Stoichiometric ratio

In the experimental protocols described above the starting volume concentrations of the aqueous species \( X \) and \( W \) are known. However, the starting surface concentrations and the effective surface area of the coating will not usually be known. Indeed, to some degree, these are idealised or conceptual quantities anyway. To overcome this difficulty the starting stoichiometric ratio
\[ r_S = \frac{\text{number of available (free) adsorption sites } S}{\text{number of entities } X} \]

is introduced as another model parameter, in addition to the reaction rates.

1.3 RESULTS AND DISCUSSION

1.3.1 Synthesis of polysulfide sorbent

The direct reaction of limonene and elemental sulfur has been used to access polysulfide materials useful in mercury sorption,\(^{15, 27, 28}\) The starting materials are low cost, the synthesis is scalable to hundreds of grams, and at high concentrations of mercury the polysulfide changes colour—for potential use in mercury sensing.\(^{28}\) However, the original preparation of this has some shortcomings. For example, the reaction was carried out at 180 °C for several hours and distillation was required to remove low molecular weight byproducts such as \( p \)-cymene and malodorous thiols.\(^{28}\) Recently, it was discovered by Hasell and co-workers that the same material could be made at a lower temperature and with fewer byproducts through the use of an accelerator or catalyst such as zinc diethylthiocarbamate.\(^{29, 30}\) This advance eliminated the need for the distillation step, as less limonene is oxidised to cymene and fewer thiols are formed. The latter feature means the product—referred to here as poly(S-\( r \)-limonene)—is not as malodorous as in the original synthesis. Hasell and co-workers also showed that poly(S-\( r \)-limonene), as a soluble oligomer, can be used to coat silica gel for mercury sorption.\(^{29}\) The focus of this report, however, was primarily on the use of accelerators and catalysts in the
copolymerisation of sulfur with alkene-containing monomers, and not mercury sorption. Therefore, there is a need to characterise this sorbent in greater detail to understand its scope and limitations in mercury sorption. Accordingly, in this study we evaluated its use in mercury sorption and desorption at varying pH and salt concentrations, and we developed a model to account for the observed kinetics. Together, these results will help guide the deployment of this sorbent in mercury remediation.

The copolymerisation was run according to Hasell’s protocol, reacting equal masses of sulfur and limonene directly in the presence of zinc diethyldithiocarbamate at 140 °C (Fig. 1 and Figs S1-S5). The accelerator was used at a loading of 5 wt% relative to the total mass of sulfur and limonene. Within 30 minutes the reaction appeared to form a single phase. After 24 hours, 1H NMR analysis indicated complete consumption of the limonene alkenes (Fig. S6). To ensure that all sulfur was consumed, the reaction was continued for 7 days. The additional reaction time did not lead to detectable changes by 1H NMR or gel permeation chromatography (GPC) but the viscosity did appear to increase somewhat over this time (Fig. S3). The molecular weight of the target oligomers was approximately 800 g/mol, based on GPC analysis (Figs S7-S12). This result was consistent across accelerator loadings of 0, 1, and 5 wt%. The poly(S-r-limonene) with 5 wt% accelerator over 7 days was used for subsequent analysis and experiments. Simultaneous thermal analysis of this sample indicated a single mass loss starting at about 200 °C and no melting transition of elemental sulfur was observed, which is consistent with complete reaction of elemental sulfur (Figs S13-S14). Consistent with this result, powder X-ray diffraction analysis revealed an amorphous material, absent of crystalline S₈ (Figs. S15-S16). Even after prolonged storage of this product (10 months), no crystalline sulfur was observed to form—a common phenomenon observed for this class of polysulfide materials. Differential scanning calorimetry indicated a glass transition temperature of Tₛ = 8.6 °C (Fig. S14). Finally, combustible analysis was used to determine elemental composition, with C = 39.8%, H = 4.9%, S = 53.5%. For equal masses of limonene and sulfur, this is consistent with the expected values of C = 44.08%, H = 5.92%, S = 50.00%, suggesting good atom economy and negligible loss of mass through volatilisation of monomers or formation of H₂S. The elemental composition also remained consistent from day 3 to 7 of the synthesis (Fig. S18).

An advantage of the poly(S-r-limonene) oligomers is their solubility in organic solvents, facilitating use as a coating. To prepare poly(S-r-limonene) coated silica gel, 13.89 g of the oligomer was dissolved in 300 mL of dichloromethane and poured over 138.9 g of silica in a 1 L round bottom flask. The polymer was dry-loaded to the surface of the silica gel by rotary evaporation—rotating the sample while distilling the solvent at 40 °C and reduced pressure (800 mbar). The coated silica was then dried further under high vacuum to remove any remaining solvent. The product was a free-flowing orange powder (Figs 1 and S19). Examination under an electron microscope reveals no morphological differences between the coated and uncoated silica (Figs. S20-S21). Surface area of the coated silica was 273.40 m²/g, as determined by Brunauer-Emmett-Teller (BET) analysis of nitrogen adsorption isotherms. The uncoated silica had a measured surface area of 444.16 m²/g (Fig. S22). The reduction in surface area for the coating sample is likely the result of the coating filling in some pores of the silica. Figure 1 summarizes the synthesis of poly(S-r-limonene) and shows the final sorbent after dry-loading the oligomer on silica.
Figure 1: An oligomer was prepared by the direct reaction of equal masses of limonene and sulfur, catalyzed by zinc diethyldithiocarbamate. The oligomer, named poly(S-r-limonene), is fully soluble in organic solvents and can be coated onto silica gel using the dry loading method. Scanning electron microscopy (SEM) and energy-dispersive X-ray (EDX) imaging of the silica gel coated with poly(S-r-limonene). Additional characterization of the polymer and coated silica is provided in the Supporting Information.

1.3.2 Sorption of HgCl₂ in water

In the initial evaluation of mercury sorption for the poly(S-r-limonene)-coated silica, 100 mL of an aqueous solution containing 5 ppm HgCl₂ was added to 500 mg of the sorbent. After 1 minute, a 1 mL sample of the solution was obtained and any trace solids were removed by centrifugation before analysis by CVAAS. Remarkably, >90% of the mercury was removed from solution within this first minute (the average of triplicate experiments). No more than 20% of the mercury was removed by uncoated silica gel, indicating the key role of poly(S-r-limonene) in this rapid mercury uptake (Fig. S23). In order to monitor mercury sorption over smaller time intervals for kinetic analysis, a revised protocol was developed. Accordingly, 500 mg of the poly(S-r-limonene)-coated silica was added to a 250 mL beaker followed by 100 mL of a 5 ppm aqueous solution of HgCl₂ with stirring. Samples were drawn up into a syringe equipped with a syringe filter to separate the sampled water from the sorbent during the sampling. Samples were obtained every 10 seconds for the first minute, and then after 90, 120, 180, and 300 seconds. Sorption was again rapid: Over 90% of mercury was sequestered within the first minute and over 98% within 5 minutes. Kinetic modelling suggested that the maximum theoretical capacity of the sorbent for Hg²⁺ was 1 mg/g (equivalent to 11 mg/g of the poly(S-r-limonene coating) under these conditions (see more details below). This result was important in that the
sorbent proved to be fast and effective at low concentrations of inorganic mercury, such as those often encountered in the environment.\textsuperscript{3, 5}

Next, similar sorption experiments were carried out with varying pH and NaCl concentrations. These experiments were designed to help define under what conditions the sorbent is effective in aqueous media. The pH was adjusted to values of 3, 5, 9, and 11 (all 5 ppm in HgCl\textsubscript{2}) and compared to the unmodified 5 ppm solution of HgCl\textsubscript{2} in water (pH = 6.99). For the effects of sodium chloride on mercury sorption, concentrations of 6.85 mM and 599 mM NaCl were evaluated—designed to mimic the salt concentrations in tap water and seawater, respectively. Sampling was carried out as describe above, with mercury concentration monitored over 300 seconds by CVAAS. The results are plotted in Figure 2. Key findings were that mercury sorption was most effective at pH = 7 or lower. However, mercury sorption slowed significantly at higher pH and bound less total mercury (see kinetic section below for additional details). The most dramatic change in sorption was at high concentrations of NaCl, with 599 mM NaCl significantly inhibiting mercury sorption. At this concentration of NaCl, only 30% of mercury was removed by the poly(S-r-limonene)-coated silica over 5 minutes, while 86% of mercury was removed when the concentration of NaCl was 6.85 mM. When no exogenous NaCl was added, 99% of the mercury was removed by 500 mg of the sorbent from a 100 mL solution of 5 ppm mercury. These results are summarized in Figure 2. Additional data is provided in Figures S2-S29.

![A. Hg\textsuperscript{2+} sorption at varying pH](image)

![B. Hg\textsuperscript{2+} sorption at varying NaCl concentrations](image)

**Figure 2**: Silica coated with poly(S-r-limonene) (10:1 silica to polymer) was used in mercury sorption experiments at varying pH and sodium chloride concentrations. **A.** 500 mg of the sorbent was added to a 5000 ppb solution of HgCl\textsubscript{2} at pH = 3, 5, 7, 9, and 11. The sorption was slower at higher pH. **B.** 500 mg of the sorbent was added to a 5000 ppb solution of HgCl\textsubscript{2} in MilliQ purified water (no added NaCl), simulated tap water (6.85 mM NaCl), and simulated seawater (599 mM NaCl). The exogenous sodium chloride slowed mercury sorption.

### 1.3.3 Kinetic analysis of HgCl\textsubscript{2} sorption

A graphical fitting procedure was used to match the experimental results to the predictions of the kinetic models. The various model parameters ($k_1, k_2, r_5$) were varied until the predictions appeared to provide the best visual fit to the log-log representation of the experimental data (Fig. 3 and Figs S30-S33). There were two main reasons for using the log-log representation in this analysis.

1. A log-log scale is a non-linear scale that expands small values of time (horizontal axis) and concentration (vertical axis) relative to larger values. This focuses attention on the time and concentration regions that are most important: early times when there is a rapid initial decrease in concentration, and low values of concentration so as to better see the long term trend in concentration.
Sampling errors are probably the most significant source of experimental error. They are more likely to be of a constant relative error nature than of a constant absolute error kind. Such relative errors are better represented as constant errors on log scales. The measurement error itself in the CVAAS assay is likely to be quite small compared to the sampling error.

In most cases, varying $k_1$ from its best fitted value by $\pm 15\%$ or more produced a noticeably poorer fit. So $\pm 15\%$ can be thought of as an informal confidence interval for the parameter estimates. A summary of the results is given in the Table 1. A few explanatory notes are listed below:

1. For the sample in which the pH was not adjusted (neutral sample, pH = 6.99), $r_S$ was taken to be a model parameter to be estimated. Its estimated value was found to be $r_S \approx 1$, implying that the sorbent and aqueous HgCl$_2$ are close to stoichiometric balance (Fig. S30). Changing $r_S$ by as little as $\pm 2.5\%$, produced noticeable deviations from the experimental data. Since all other experimental cases used the same amounts of HgCl$_2$ and sorbent, it was supposed that $r_S$ would be unchanged, and it was given the fixed value $r_S = 1$ for these cases also.

2. For the neutral case (pH = 6.99) and the acidic cases (pH = 3 and pH = 5) the single reaction model was adequate to fit the experimental data well (Fig. S31). All these cases show a continuing decrease in Hg concentration with time over the sampling period. Indeed, the log-log plots asymptote to a slope of -1, indicating a $1/t$ asymptotic behaviour (in non-logarithmic units). The physical interpretation of these results is that ultimately all X will be adsorbed and all adsorption sites S will be utilised.

3. For the basic cases (pH = 9 and pH = 11) the two-reaction model was needed to fit the apparent steady state that was approached. Here the competing species W was taken to be NaOH. The physical interpretation of these results is that ultimately all adsorption sites S will be utilised, some by X, and the remainder by W. The ratio of the amounts of X and W adsorbed depends on the ratio $k_1/k_2$ as well as the starting stoichiometric ratios $r_S$ and $r_W$. Further details can be found in the Supplementary Material (Fig. S32).

4. For the tap and seawater cases, the two-reaction model was again needed to fit the apparent steady state that was approached (Fig. S33). Here the competing species W was taken to be NaCl. The physical interpretation is similar to that for the base cases but with NaCl as the competing species.

5. For the seawater case there was only a small reduction in Hg concentration during the 5 minutes of the experiment as the apparent steady state was approached. Because of this the model parameters, in particular $k_1$, could not be estimated with great accuracy. An informal confidence range for $k_1$ has been given in Table 1.
Figure 3: Kinetic models fit to mercury sorption data on log-log scales. A. Neutral pH, no added NaCl. B. pH 3. C. pH 5. D. pH 9. E. pH 11. F. Simulated tap water (6.85 mM NaCl) G. Simulated seawater (599 mM NaCl)
Table 1: Estimated values of the model parameters $k_1$ and $k_2$ based on fitting to experimental data. Unless specified otherwise, varying $k_1$ by more than ±15% produced predictions that are noticeably different from the experimental results. Based on the pH = 7 results, $r_5$ was estimated to be $r_5 = 1 ± 0.025$, and this value was used for all other cases.

| Case              | $k_1$ [M$^{-1}$s$^{-1}$] | $k_2$ [M$^{-1}$s$^{-1}$] | $k_2/k_1$ | Comment                  |
|-------------------|--------------------------|--------------------------|-----------|--------------------------|
| pH 3              | 2.2 x 10$^5$             |                          |           | single reaction model    |
| pH 5              | 3.6 x 10$^3$             |                          |           | single reaction model    |
| pH 7              | 6.7 x 10$^3$             |                          |           | single reaction model    |
| pH 9              | 4.9 x 10$^5$             | 2.2 x 10$^2$             | 4.6 x 10$^2$ | single reaction model, $r_5 = 1$ |
| pH 11             | 3.2 x 10$^5$             | 4.3                      | 1.4 x 10$^3$ |                          |
| Tap water 6.8 x 10$^{-3}$ M NaCl | 1.3 x 10$^3$             | 0.32                     | 2.5 x 10$^{-4}$ |                          |
| Saltwater 0.60 M NaCl | 6.7 x 10$^2$             | 4.9 x 10$^{-2}$          | 7.3 x 10$^{-5}$ | $k_1 ≈ 4 x 10^2 − 1 x 10^3$  |

Variation of $k_1$ across experimental conditions

It is instructive to plot $1/k_1$ against pH or the log of the chloride ion concentration (as appropriate for the case). This is shown in Figure 4. As all cases were prepared from HgCl$_2$, there was always a background Cl concentration arising from the 5 ppm HgCl$_2$. For this reason, the pH = 7 case has been treated as a (low level) reference for the acid, base and chloride cases. In the figure each of the acid (blue), base (red) and NaCl (black) segments are approximately straight lines over the ranges of concentration considered. This observation needs to be treated with caution as it is based on only three data points for each segment, and as noted above, there is considerable uncertainty in the estimates of $k_1$. Moreover, $1/k_1$ must have a strictly positive limit as the concentrations of H$^+$, HO$^-$ or Cl$^-$ tend to 0. Nonetheless, these plots give some indication that the presence in high concentrations of H$^+$, HO$^-$ or Cl$^-$ decreases $k_1$ (increases $1/k_1$) in a systematic way.

Figure 4: Plot of $1/k_1$ vs pH (for acid and base cases) and chloride concentration (for NaCl cases). There are different segments for each case. The estimated $k_1$ value for the pH = 7 experiment is plotted as a (reference) data point on both the acid and base segments, and also as the leftmost data point on the NaCl segment.
1.3.4 Influence of pH on mercury sorption kinetics

As seen in Table 1 and Figure 4, the poly(S-r-limonene) sorbent is fastest at neutral or near-neutral pH, with reduced rates of mercury uptake below or above pH 7. For example, within 5 minutes, 500 mg sorbent captured 95, 96, 94 and 90% of mercury from solutions at pH 3, 5, 9 and 11, respectively. In contrast, the sorbent removed 99% of the mercury in the same 5 minute period in the neutral sample under otherwise identical conditions. The pH might alter the mercury speciation, which could account for these differences in rates of mercury uptake. Note that for neutral and low pH the single reaction model fits the experimental data well, suggesting that the availability of sorption sites is not significantly affected by these pH values. This is also shown by Figure 3 where there is no apparent flattening of the curves for cases A, B and C. This contrasts with the high pH cases where a two-reaction model with a competing reaction was required. For high pH, hydroxide might potentially break S-S bonds in the polysulfide which could hinder both the measured rate of mercury sorption in addition to the capacity of the sorbent, which is reflected in the flattening of the model curve in Figures 3D and 3E. Indeed, prolonged exposure to high pH resulted in degradation of the polymer coating (see desorption experiments for additional discussion of this observation).

1.3.5 Influence of sodium chloride on mercury sorption kinetics

Sodium chloride clearly interfered with the mercury sorption of poly(S-r-limonene)—an important new finding. For a 100 mL sample of 5 ppm Hg\(^{2+}\) containing 6.85 mM NaCl, 500 mg of the poly(S-r-limonene)-coated silica gel only removed 86% of the mercury after 5 minutes. For a 100 mL sample of 5 ppm Hg\(^{2+}\) containing 599 mM NaCl, 500 mg of the poly(S-r-limonene)-coated silica gel only removed 30% of the mercury after 5 minutes. In contrast, with no exogenous NaCl added in an otherwise identical experiment, 99% of the mercury was removed.

The origin of the inhibition can be multi-faceted, but the models in Figure 3F and 3G clearly indicate that the NaCl reduces both the rate of mercury uptake and the amount of mercury that can be bound. The latter point is indicative of NaCl competing for or otherwise blocking binding sites. Indeed, NaCl particles could be detected directly on the sorbent. For instance, incubating the poly(S-r-limonene)-coated silica in water with 599 mM NaCl for one month resulted in substantial amounts of sodium chloride on the surface of the polymer, even after isolating by filtration and washing with water. It might also be important to consider chloride as a ligand for Hg\(^{2+}\), which may compete with sulfur in mercury binding. In any case, it is clear that the sorbent’s effectiveness is reduced with increasing NaCl. This suggests that its use in remediation of seawater may not be practical or require more sorbent than for samples with lower levels of NaCl, as discussed next.

To test if a higher concentration of sorbent can achieve more complete mercury binding in the presence of sodium chloride, 9 times the mass of sorbent was used in an identical sorption experiment. The kinetic model was used to predict that this amount of sorbent would provide sufficient binding sites to overcome the competing processes with NaCl. The qualitative concept here was that solely by a mass action effect, adding more sorbent would speed up both the desired mercury sorption and the undesired competing reaction. However, due to the relative rates of these reactions, \(k_1/k_2\), the sorption reaction is now able to proceed further before a steady state is achieved. Accordingly, 4.5 g poly(S-r-limonene) coated silica was mixed with 100 mL 5 ppm Hg\(^{2+}\) containing 599 mM NaCl for 5 minutes with regular sampling. Within the first 10 seconds the sorbent removed 83% of the mercury present—a greatly improved initial uptake than observed in the original experiment with 9-times less sorbent. Over 5 minutes, 91% of the mercury was removed (Fig S36). This experiment shows that additional sorbent can overcome the detrimental effect of sodium chloride on mercury binding performance as predicted from the kinetic model.

1.4 Desorption of mercury from sorbent

An important factor to consider in remediation is the stability of the spent sorbent. For instance, it is important that the mercury does not leach during the transport and storage of the spent...
sorbent. To evaluate desorption of mercury from the spent sorbent, 4.00 g of the poly(S-r-limonene)-coated silica was added to a 200 mL aqueous solution of HgCl₂ (20 ppm). The mixture was stirred for 10 minutes and then the sorbent was isolated by filtration and dried under vacuum. Based on the mercury remaining in the water (as determined by CVAAS), the spent sorbent bound 1.03 mg Hg²⁺ per gram of sorbent. To evaluate leaching, 500 mg portions of the spent sorbent were added to separate 50 mL aqueous solutions of varying pH and sodium chloride concentrations (pH = 3, 5, 7, 9, 11 and 6.85 mM or 599 mM NaCl). These seven samples were monitored over 28 days to determine how much mercury leached into the water. Minimal leaching was observed for all samples except for the sample at pH = 11 (Figs. 5A and S37). After 28 days, approximately 3% of the bound mercury had leached into the basic solution at pH = 11. All other samples had mercury levels at or below the limits of detection by CVAAS, indicating leaching was not significant for pH = 2, 5, 7, or 9. Leaching was also not significant for the samples with added sodium chloride. SEM analysis of the sorbent did reveal that the high pH degrades the polymer coating (Figs 5B and S37). This could lead to the release of mercury, perhaps bound to suspended polymer particles (or the products of polymer degradation). Further study is required to determine the mechanism and speciation of the mercury leaching at elevated pH. However, minimal leaching was observed for all other samples suggesting that the mercury remains strongly bound to the sorbent even in highly acidic media or brine.

**Figure 5:** A. The spent sorbent (saturated at 1 mg HgCl₂ per gram sorbent) was subjected to leaching experiments at varying pH and sodium chloride concentrations. A mass of 500 mg of the spent sorbent was added to 50 mL of the aqueous solution for these leaching experiments. The dotted line represents the limit of detection of 5 ppb. Only at pH 11 was significant leaching observed. After 28 days, this leaching still only corresponds to 3% of the bound mercury. B. SEM micrograph of the sorbent after the leaching experiment at pH 11. The coating appears to have been degraded at the high pH, as the surface was no longer smooth as seen in Figs 1 and in the other leaching experiments (Figs. S38-S47).

### 1.4 Conclusions
Poly(S-r-limonene)-coated silica was evaluated as a mercury sorbent over a range of pH values and also in the presence of sodium chloride. The sorbent rapidly removed HgCl₂ from water at or near neutral pH. Slightly reduced rates of uptake were observed at both low and high pH, but the sorbent was still effective across this wide pH range. One-reaction or two-reaction kinetic models were fitted to the experimental sorption results. These modes suggest that under low (acid) and neutral pH conditions, mercury sorption is a single reaction process which will ultimately proceed to completion until one or both of the mercury or the sorbent is consumed. In contrast, for basic pH and in the presence of NaCl, the models suggest that there are significant competing reactions whereby some of the sorbent becomes unavailable for mercury binding. It was also discovered that sodium chloride severely inhibits mercury binding, which could limit the use of the sorbent in salt water systems. The spent sorbent was found to be stable.
and did not leach significant mercury from pH = 3 to pH = 9. Aqueous sodium chloride also did not lead to leaching. However, at pH = 11 the coating degraded and mercury was released into the solution, possibly bound to suspended polymer particles or other polymer degradation products. Together, these results suggest that the sorbent is most effective at low and neutral pH and that elevated pH can lead to polymer degradation. This assessment of the scope and limitations of this sorbent will help define the conditions for which it is most effective in the field. While this study focussed on inorganic mercury, future studies will be carried out that evaluate the sorbent on a broader range of mercury species and field samples.

Author contributions
Max J. H. Worthington: Data curation, Formal analysis, Investigation, Methodology, Visualization, Writing – original draft, Writing – review and editing
Maximilian Mann: Data curation, Formal analysis, Methodology, Investigation, Writing – review and editing
Ismi Yusrina Muhti: Data curation, Formal analysis, Methodology, Investigation, Writing – review and editing
Zhongfan Jia: Formal analysis, Investigation, Supervision, Writing – review and editing
Anthony D. Miller: Conceptualization, Data curation, Formal analysis, Methodology, Project administration, supervision, Visualization, Writing – original draft, Writing – review and editing
Justin M. Chalker: Conceptualization, Formal analysis, Funding acquisition, Methodology, Project administration, Resources, Supervision, Writing – original draft, Writing – review and editing

Acknowledgements
This work was supported by the Australian Research Council (DP200100090). The authors also acknowledge the support of Flinders Microscopy and Microanalysis for access to polymer characterisation facilities.

References
1. Global Mercury Assessment 2018, UN Environment Programme, Chemicals and Health Branch, Geneva, Switzerland.
2. L. J. Esdaile and J. M. Chalker, Chem. Eur. J., 2018, 24, 6905-6916.
3. Q. Wang, D. Kim, D. D. Dionysiou, G. A. Sorial and D. Timberlake, Environ. Pollut., 2004, 131, 323-336.
4. P. B. Tchounwou, W. K. Ayensu, N. Ninashvili and D. Sutton, Eviren. Toxicol., 2003, 18, 149-175.
5. L. Wang, D. Hou, Y. Cao, Y. S. Ok, F. M. G. Tack, J. Rinklebe and D. O’Connor, Environ. Int., 2020, 134, 105281.
6. R. G. Pearson, J. Am. Chem. Soc., 1963, 85, 3533-3539.
7. J. M. Chalker, M. Mann, M. J. H. Worthington and L. J. Esdaile, Org. Mater., 2021, 3, 362-373.
8. M. Mann, X. Luo, A. D. Tikoalu, C. T. Gibson, Y. Yin, R. Al-Attabi, G. G. Andersson, C. L. Rason, L. C. Henderson, A. Pring, T. Hasell and J. M. Chalker, Chem. Commun., 2021, 57, 6296-6299.
9. T. Tian, R. Hu and B. Z. Tang, J. Am. Chem. Soc., 2018, 140, 6156-6163.
10. W. J. Chung, J. J. Griebel, E. T. Kim, H. Yoon, A. G. Simmonds, H. J. Ji, P. T. Dirlam, R. S. Glass, J. J. Wie, N. A. Nguyen, B. W. Guralnick, J. Park, A. Somogyi, P. Theato, M. E. Mackay, Y.-E. Sung, K. Char and J. Pyun, Nat. Chem., 2013, 5, 518-524.
11. Y. Zhang, R. S. Glass, K. Char and J. Pyun, Polym. Chem., 2019, 10, 4078-4105.
12. M. J. H. Worthington, R. L. Kucera and J. M. Chalker, Green Chem., 2017, 19, 2748-2761.
13. J. M. Chalker, M. J. H. Worthington, N. A. Lundquist and L. J. Esdaile, Top. Curr. Chem., 2019, 377, 16.
14. T. Lee, P. T. Dirlam, J. T. Njardarson, R. S. Glass and J. Pyun, *J. Am. Chem. Soc.*, 2022, **144**, 5-22.
15. T. Hasell, D. J. Parker, H. A. Jones, T. McAllister and S. M. Howdle, *Chem. Commun.*, 2016, **52**, 5383-5386.
16. M. W. Thielke, L. A. Bultema, D. D. Brauer, B. Richter, M. Fischer and P. Theato, *Polymers*, 2016, **8**, 266.
17. J. M. Scheiger, C. Direksilp, P. Falkenstein, A. Welle, M. Koenig, S. Heissler, J. Matysik, P. A. Levkin and P. Theato, *Angew. Chem. Int. Ed.*, 2020, **59**, 18639-18645.
18. D. J. Parker, H. A. Jones, S. Petcher, L. Cervini, J. M. Griffin, R. Akhtar and T. Hasell, *J. Mater. Chem. A*, 2017, **5**, 11682-11692.
19. L. A. Limjuco, G. M. Nisola, K. J. Parohinog, K. N. G. Valdehuesa, S.-P. Lee, H. Kim and W.-J. Chung, *Chem. Eng. J.*, 2019, **378**, 122216.
20. S. Akay, B. Kayan, D. Kalderis, M. Arslan, Y. Yagci and B. Kiskan, *J. Appl. Polym. Sci.*, 2017, **134**, 45306.
21. A. M. Abraham, S. V. Kumar and S. M. Alhassan, *Chem. Eng. J.*, 2018, **332**, 1-7.
22. V. S. Wadi, H. Mittal, E. Fosso-Kankeu, K. K. Jena and S. M. Alhassan, *Colloids Surf.*, **A**, 2020, **606**, 125333.
23. J. Lee, S. Lee, J. Kim, Z. Hanif, S. Han, S. Hong and M.-H. Yoon, *Bull. Korean Chem. Soc.*, 2018, **39**, 84-89.
24. L. A. Limjuco, H. T. Fissaha, H. Kim, G. M. Nisola and W.-J. Chung, *ACS Appl. Polym. Mater.*, 2020, **2**, 4677-4689.
25. M. Mann, B. Zhang, S. J. Tonkin, C. T. Gibson, Z. Jia, T. Hasell and J. M. Chalker, *Polym. Chem.*, 2022, in press, DOI: doi.org/10.1039/D1PY01416A.
26. M. L. Eder, C. B. Call and C. L. Jenkins, *ACS Appl. Polym. Mater.*, 2022, in press, DOI: 10.1021/acsapm.1c01536.
27. M. P. Crockett, A. M. Evans, M. J. H. Worthington, J. M. Chalker, Sulfur-Limonene Polysulfide. US Patent 10,590,012, 2020. Priority date 13 Oct 2015.
28. M. P. Crockett, A. M. Evans, M. J. H. Worthington, I. S. Albuquerque, A. D. Slattery, C. T. Gibson, J. A. Campbell, D. A. Lewis, G. J. L. Bernardes and J. M. Chalker, *Angew. Chem. Int. Ed.*, 2016, **55**, 1714-1718.
29. X. Wu, J. A. Smith, S. Petcher, B. Zhang, D. J. Parker, J. M. Griffin and T. Hasell, *Nat. Commun.*, 2019, **10**, 647.
30. L. James Dodd, Ö. Omar, X. Wu and T. Hasell, *ACS Catal.*, 2021, **11**, 4441-4455.