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1 Computational studies on the Reactions of Thiols,
2 Sulfides and Disulfides with Hydroperoxides.
3 Relevance for Jet Fuel Autoxidation.

4
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7
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15
16 **Abstract**

17
18 Density Functional Theory calculations (DFT) are reported on the reactions of hydroperoxides
19 with different classes of sulfur: thiols (RSH), sulfides (RSR) and disulfides (RSSR), all of
20 which are important trace species in the auto-oxidation of jet fuel. It is shown that thiols can
21 react under auto-oxidation conditions with hydroperoxides to form sulfonic acids and alcohols.
22 In contrast, it is shown that disulfide species are more likely to form thiyl radicals, which are

23 less likely to be important for the direct autoxidation of fuels due to prohibitive reaction
24 barriers. The reaction mechanisms reported here for sulfur oxidation and the associated
25 calculated thermodynamic data can be used to extend the applicability of current chemical
26 kinetic models for fuel autoxidation, which are currently treated as a single elementary reaction
27 despite the range of sulfur species found in fuels.

28

29 **Keywords:** DFT, Fuel autoxidation, Sulfur oxidation, Reaction Mechanisms.

30

31 **Introduction**

32

33 Jet fuel is primarily comprised of a blend of aliphatic and aromatic hydrocarbons as well as
34 low levels of hydroperoxides and trace heteroatomic species (the latter generally in ppm
35 quantities). The hydrocarbons afford the fuel its bulk properties such as viscosity, density and
36 volatility which are not only important when it is used as a propellant, but also when exploited
37 as a coolant as happens in modern aircraft. In the latter use, at temperatures of approximately
38 140 °C a process called autoxidation is initiated. Here a series of chemical reactions can occur
39 that can lead to both bulk and surface deposits. Experimentally, it has been shown that many
40 species can have an effect on this process including sulfur compounds¹⁻³, polar species⁴⁻⁹,
41 dissolved metals^{10, 11} and hydroperoxides.^{12, 13} A number of studies have reported on the
42 thermal stability of jet fuel, with a focus on the oxidative mechanisms involved therein.¹⁴⁻²¹

43

44 In the current paper, we focus on the reactions between sulfur species and hydroperoxides. A
45 number of different classes of sulfur compounds can be present in jet fuels as shown in **Figure**



65

66 **Figure 2:** Reaction of hydroperoxides and sulfides to form alcohols and sulfones. (R =
 67 aliphatic, aromatic)

68

69 A number of literature studies have reported on the effects of sulfur species on jet fuel
 70 autoxidation. In 1976, Taylor showed that thiophenes and disulfide species contribute
 71 significantly to deposition.⁸ Moreover, dibenzothiophenes were found to have little effect on
 72 deposition. These observations were further corroborated by Mushrush et al. who showed that
 73 thiophenes and thiophenols have a destabilizing effect on jet fuel.²⁹ Further work by
 74 Offenhauer and Hiley showed that both sulfonic acids and thiophenols increased both
 75 deposition and gum formation whereas diaryl sulfides have little effect.^{31, 32} The proposed dual
 76 roles that sulfur compounds can play was reported by both Denison et al. and Thompson et
 77 al..^{33, 34} They observed that disulfides decreased the thermal stability of the fuel. In contrast,
 78 aliphatic sulfides had a small stabilizing effect. In a series of experimental studies, Daniel and
 79 Henemen doped jet fuel with different sulfur species at a range of concentrations.³⁵ Thiols and
 80 thiophenes decreased the thermal stability whilst disulfides and sulfides increased the thermal
 81 stability. The largest amount of deposits were observed by Rawson et al. for model fuels doped
 82 with benzylosulfonic acid, diphenyl disulfide and elemental sulfur.³⁶ The contrasting results in
 83 many of these studies highlight the complex nature of the underlying mechanisms.

84

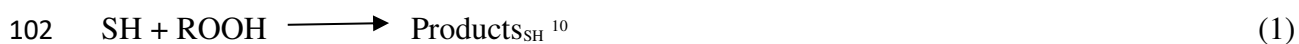
85 Despite the importance of these trace sulfur species in jet fuel, there has only been a limited
 86 number of computational studies into the energetics of the reactions involving these species.
 87 Bach and co-workers investigated the activation energy for the reaction of Me₂S with

88 methylperoxide (MeOOH) and tert-butyl peroxide (^tBuOOH), which were calculated to be 32.4
89 and 32.2 kcal mol⁻¹ respectively.³⁷ In these calculations it was shown for the first time that the
90 transition state is concerted. In particular, the hydroxyl oxygen atom in the hydroperoxide is
91 transferred to the sulfur as the hydrogen moves to form the alcohol. More recently, Zabarnick
92 et al. modelled the reactions of Et₂S and EtSSEt with ⁿBuOOH.³⁰ The activation energies for
93 these reactions were 26.1 and 28.7 kcal mol⁻¹.

94

95 The lack of detailed investigations into the reactions of sulfur compounds and hydroperoxides
96 prompted our investigation. Current chemical kinetic models, which aim to predict the rate of
97 fuel autoxidation describe sulfur chemistry using one elementary reaction (**equation 1**).¹⁰
98 Given the wide range of sulfur species that could be present in jet fuels, it is likely that the
99 applicability of such kinetic models could be improved by a more thorough understanding of the
100 individual reactions each sulfur species could undergo.

101



103

104 In this paper, we report on a series of DFT calculations on a large range of sulfur compounds,
105 which are each in turn reacting with multiple hydroperoxides. Further reactions beyond the
106 initial oxidation are considered. The different routes to the formation of sulfones and sulfonic
107 acids are also probed. The propensity of disulfide species to undergo fission reactions and
108 form thiyl radicals are also investigated as are the potential implications of the formation of
109 such species. In its entirety, our work provides an extensive thermochemical library for sulfur–
110 peroxide reactions to further improve current kinetic models for aviation fuel degradation.

111

112

113 2 Experimental and computational Details

114

115 An analysis of a sample of Jet A-1 fuel was conducted in order to determine which sulfur
116 species were the most appropriate to model. Jet A-1 fuel samples were analysed for sulfur
117 content using an in-house method. This method identifies the sulfur species in middle
118 distillates using an Agilent 7890N Gas chromatogram combined with a Zoex thermal
119 modulation and chemiluminescence detector. Quantification of the types of sulfur species
120 present was achieved through normalization against the total sulfur content of the fuel sample
121 as determined by combustion followed by UV fluorescence analysis. This analytical method
122 separates all the sulfur-containing compounds according to their boiling point and polarity.
123 This afforded the ability to differentiate between benzothiophenes and dibenzothiophenes
124 observed as two well-defined bands, which were well separated from the lower polarity
125 thiophenes, mercaptans and sulfides.³⁸⁻⁴⁰

126

127 DFT calculations were performed using Gaussian 09 software⁴¹, version D.01, using Gaussian-
128 supplied versions of BLAS and ATLAS.^{42, 43} All calculations used the B3LYP functional.⁴⁴⁻⁴⁶
129 The cc-pVTZ basis set was used for all elements.⁴⁷ Benchmarking studies show this setup to
130 be acceptable.^{16, 19} In all calculations solvent was accounted for by the polarizable continuum
131 model (PCM) method using solvent parameters for dodecane as implemented in Gaussian.^{48, 49}
132 Calculations were carried out at 298.15 K. Geometry optimizations were confirmed to be local
133 minima by the absence of imaginary frequencies in the vibrational spectra. Transition states
134 were optimized using the QST3 method as implemented in Gaussian.⁵⁰ All transition states
135 were confirmed both visually *via* the presence of one large imaginary frequency corresponding
136 to the saddle point and *via* intrinsic reaction coordinate (IRC) scans. An ultrafine grid was
137 employed for all calculations with no symmetry constraints. Radical species were calculated

138 as singlets with the HOMO and LUMO orbitals mixed (guess=mix option) in order to break
139 the symmetry of the system. Free energies were calculated using the Grimme quasiharmonic
140 entropy correction using the GoodVibes script.⁵¹ Selected stationary points were improved with
141 coupled cluster calculations (CCSD(T)) using MolPro.⁵² Quoted values in the manuscript are
142 subject to an error of ± 2.5 kcalmol⁻¹ based upon benchmarking studies.^{16, 19}

143

144 Activation energies are calculated as the energy difference between the transition state and both
145 reactants at infinite separation. Arrhenius pre-exponential factors were calculated according
146 to **equation 2**.

147

$$148 \quad A = \frac{k_b T}{h} \exp\left(\frac{\Delta S}{R}\right) \quad (2)$$

149

150 **3 Results and Discussion**

151 **3.1 Sulfur selection**

152 **Table 1:** Results from the analysis of sulfur content in a Jet A-1 fuel sample

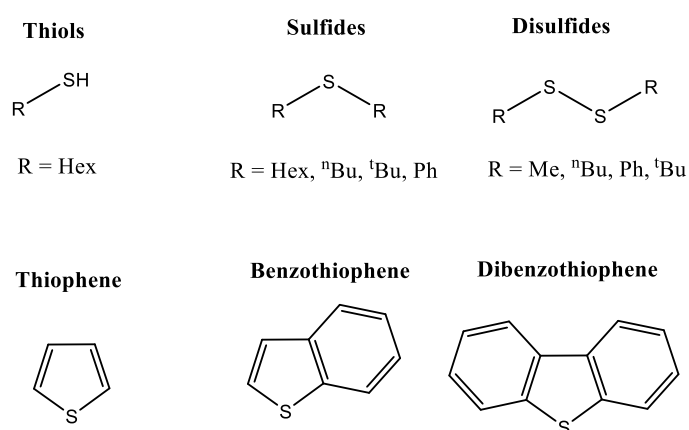
Sulfur species	Amount detected / %
Thiols, sulfides and disulfides	57
Dibenzothiophenes	26
Unknown	12
Dibenzothiophenes	2
Benzothiophenes	2
Substituted dibenzothiophenes	1

153

154 In order to guide our computational investigations, we initially conducted a speciation analysis
155 of a JetA-1 fuel using GC x GC as described in the experimental section. The results of this
156 analysis are shown in **Table 1**. By far, the most abundant sulfur species detected were thiols,
157 disulfides and sulfides (57 %). These were followed by substituted benzothiophenes (26.0 %)

158 and dibenzothiophenes (12.0 %). A small quantity of non-substituted benzothiophenes were
159 also detected. Based upon these observations, the species shown in **Figure 3** were selected for
160 computational analysis. For thiols, sulfides and disulfides a mixture of aromatic and aliphatic
161 R groups were selected. These were chosen to provide as wide a selection of structures as
162 possible within the range of compounds detected in the jet fuel sample.

163



164

165 **Figure 3:** Structures of sulfur species selected for computational analysis

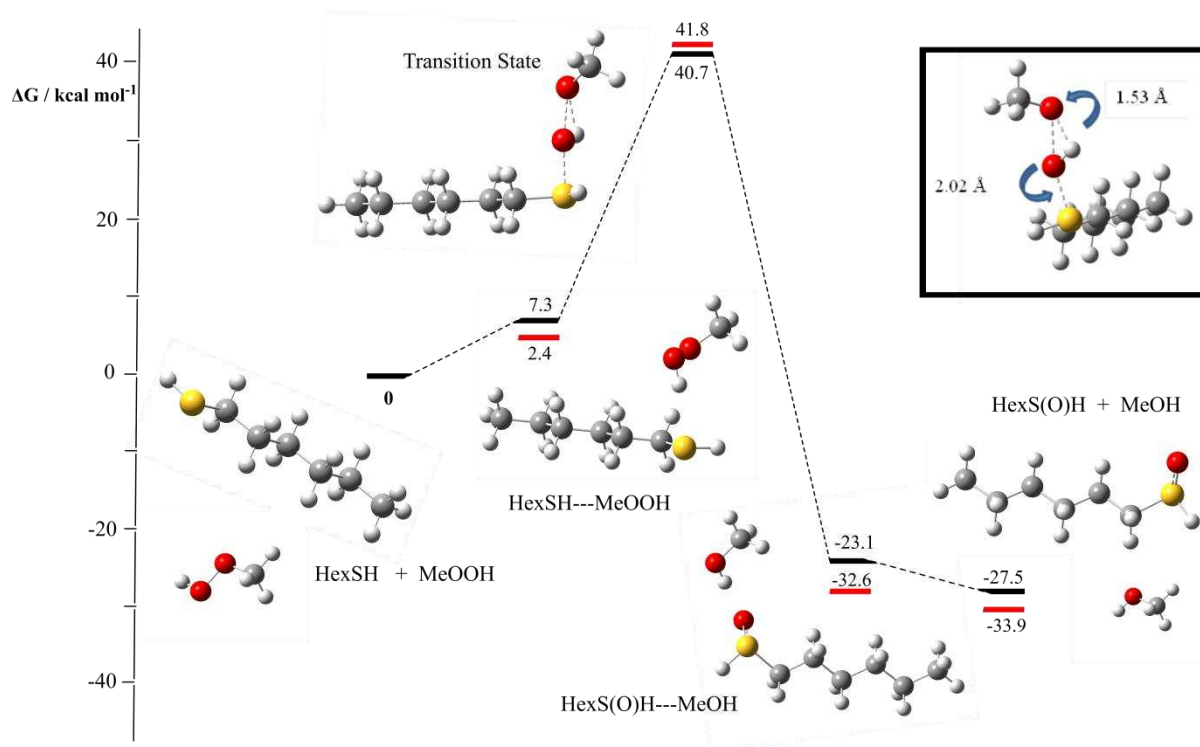
166

167 3.2 Reactions of thiols and sulfides with hydroperoxides

168

169 Our computational studies were initially focused on how the aforementioned sulfur species
170 might affect autoxidation by investigating how they react with hydroperoxides. It has
171 previously been reported that sulfur species react with hydroperoxides to form alcohols and
172 oxidized sulfur species through a non-radical pathway.²⁹ Our initial studies focused on the
173 reactions of thiols (general formula RSH). The energy profile for the reaction of hexane thiol
174 and methyl hydroperoxide is shown in **Figure 4**.

175



176

177 **Figure 4:** Gibbs free energy profile for the reaction of hexane thiol and MeOOH. Inset:

178 Transition state structure. Results from CCSD(T) calculations shown in red.

179

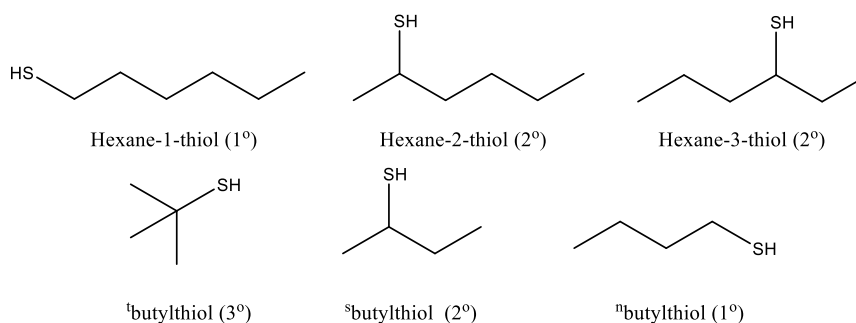
180 The pre-reaction complex, (HexSH---MeOOH in **Figure 4**) is defined as the structure where
 181 the two reactants are in close proximity immediately prior to the reaction taking place. Here,
 182 the pre-reaction complex is 7.3 kcal mol^{-1} less stable than the separated reactants. The
 183 transition state geometry is shown inset in **Figure 4**. The hydroxyl oxygen of the incoming
 184 hydroperoxide is transferred to the sulfur atom of the thiol. Concurrently, the hydroxyl
 185 hydrogen is transferred back to form an alcohol. The activation free energy for this reaction is
 186 40.7 kcal mol^{-1} and the reaction free energy is -27.5 kcal mol^{-1} .

187

188 Next, we considered the effect of the chosen hydroperoxide on the activation energy. It is
 189 computationally efficient to conduct these calculations using MeOOH as a model for the
 190 hydroperoxides found in jet fuel. However, it is not a representative structure. Fuel
 191 hydroperoxides will contain a much longer carbon chain or could be derived from branched

192 paraffinic species. Therefore, we investigated the reactions with different hydroperoxides to
193 ascertain whether MeOOH was an appropriate model. To address this, we calculated the same
194 energy profile as shown in **Figure 4**, replacing MeOOH with a range of linear alkyl peroxides
195 (Carbon numbers 1 to 12) and two branched hydroperoxides – cumene hydroperoxide (CHP)
196 and ^tBuOOH. The effect on the activation energy and reaction energy of increasing chain
197 length is presented in **Figure S3** in the supporting information.

198
199 As can be seen from **Figure S3**, the effect of increasing the hydroperoxide chain length on both
200 the activation free energy and reaction free energy is small. In particular, activation free
201 energies range from 40.7 - 42.4 kcal mol⁻¹. Reaction free energies range from -33.6 to -34.3
202 kcal mol⁻¹. A reduction in the activation free energy to 39.95 kcal mol⁻¹ is noted when CHP is
203 used as the model hydroperoxide. The corresponding value for ^tBuOOH is 41.5 kcal mol⁻¹. In
204 light of these results, further studies hereafter are carried out using both MeOOH, ^tBuOOH and
205 CHP. MeOOH was chosen as it clearly is an acceptable model for the larger linear-chained
206 hydroperoxides. Moreover, it can be modelled in a timely manner. CHP and ^tBuOOH were
207 chosen as more realistic fuel peroxides for aromatic and branched hydroperoxides respectively.
208 Data for the reactions of sulfur species and both ^tBuOOH and CHP can be found in the
209 supporting information and is collated in **Table 3**.



210
211 **Figure 5:** Primary (1°), secondary (2°) and tertiary (3°) thiol species selected for
212 investigation

213

214 Consideration was also given as to how branching in the sulfur species affects the reactivity
 215 with hydroperoxides. Several sulfur species were selected for investigation as shown in **Figure**
 216 **5**, containing primary, secondary and tertiary thiols. In each case, the reaction with MeOOH
 217 was modelled. The Gibbs free activation and reaction energies are collated in **Table 2**. As can
 218 be seen, branching of the carbon chain has a minimal effect on both the Gibbs free activation
 219 and reaction energies for the first oxidation reaction. This suggests that, at least for thiols
 220 containing only carbon chains and no further functional groups, linear thiols can be investigated
 221 as model complexes.

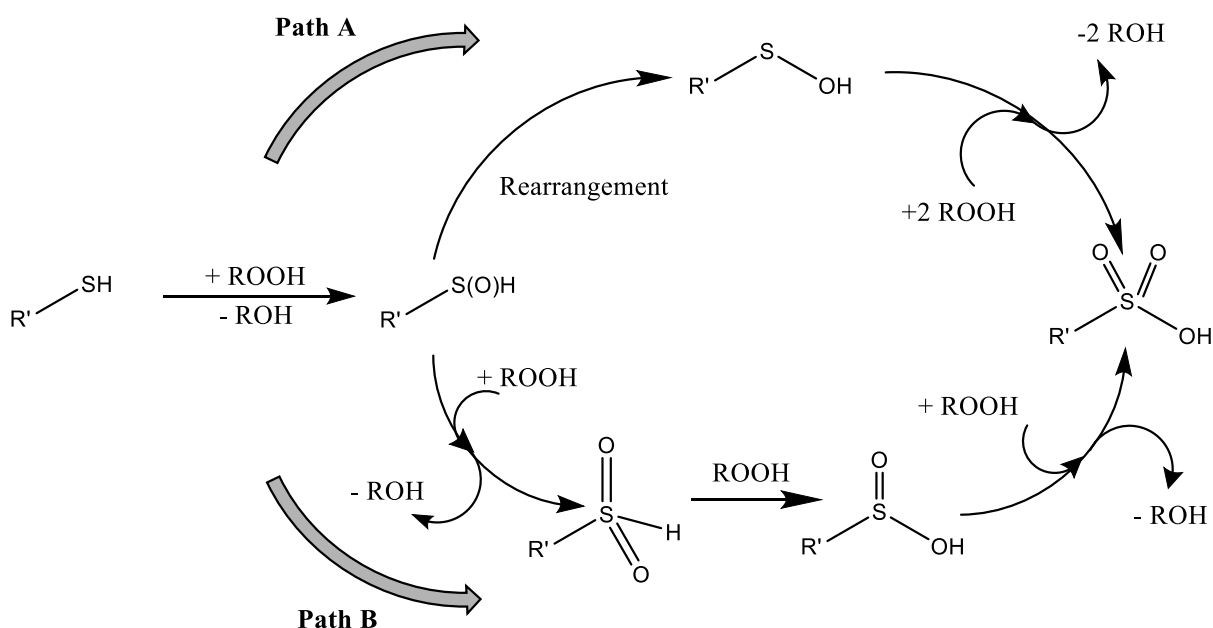
222

223 **Table 2:** Collated Gibbs free activation and reaction energies for the reaction of primary
 224 (1°), secondary(2°) and tertiary thiols (3°) and MeOOH.

Sulfur species (type)	Gibbs free activation energy / kcal mol ⁻¹	Gibbs reaction energy
Hexane-1 thiol (1°)	41.8	-27.5
Hexane-2 thiol (2°)	40.9	-27.0
Hexane-3 thiol (2°)	42.6	-25.7
ⁿ butylthiol (1°)	41.8	-25.7
^s butylthiol (2°)	40.8	-28.0
^t butylthiol (3°)	40.4	-29.1

225

226



227

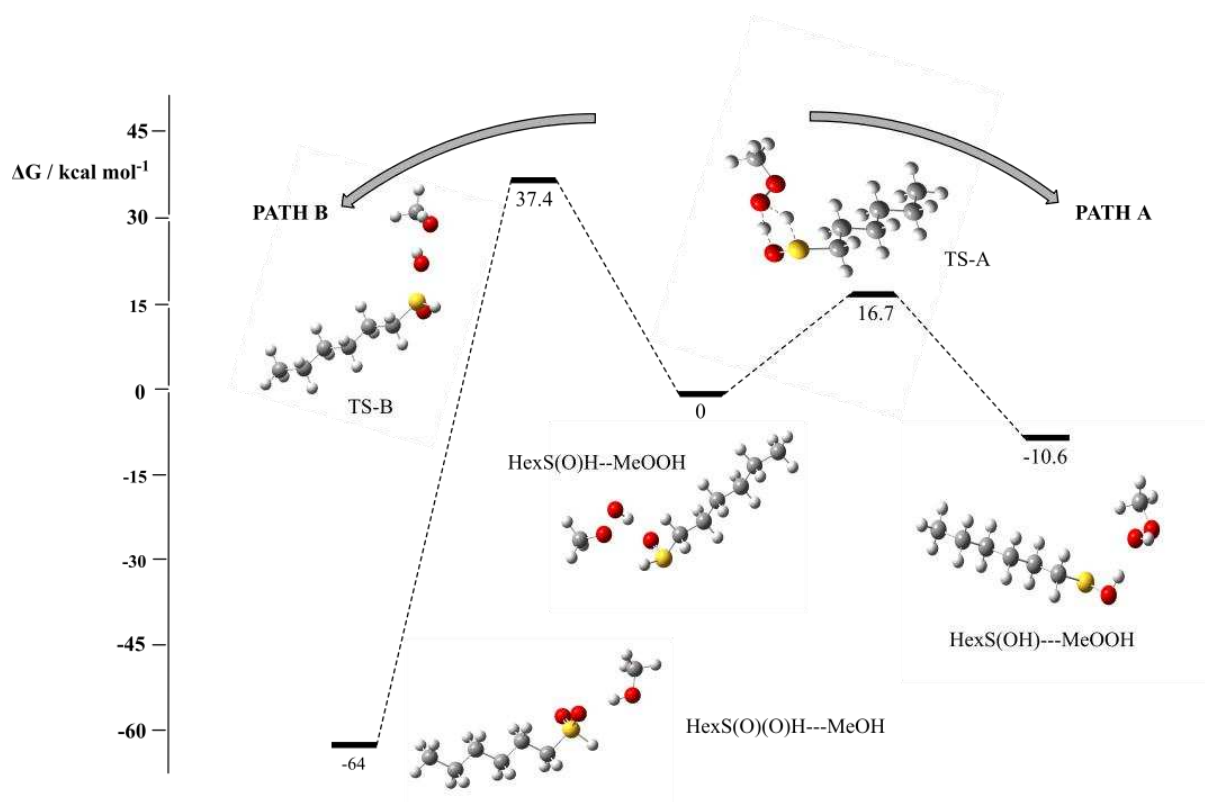
228 **Scheme 1:** Two potential mechanistic pathways leading to formation of $RS(O)(O)OH$.

229

230 Having considered the effect of both branching in the sulfur species and how the hydroperoxide
 231 chain length affects the reactivity, we turned our attention to the subsequent reactions that the
 232 oxidized thiol can undergo. The reaction of hexanethiol and $MeOOH$ from **Figure 4** initially
 233 forms an oxidized sulfur species ($RS(O)H$) and an alcohol. Through successive further
 234 oxidations it can form a sulfonic acid ($RS(O)(O)OH$). This can occur through two distinct
 235 routes as illustrated in **Scheme 1**. The oxidized sulfur species can first rearrange to form a
 236 sulfenic acid ($RSOH$) which can subsequently undergo two further oxidation reactions with
 237 hydroperoxides to form the sulfonic acid (**path A** in **Scheme 1**). Alternatively, two further
 238 oxidation reactions can occur either side of a rearrangement to form $RS(O)(O)OH$ (**path B** in
 239 **Scheme 1**). The energy profile illustrating the two divergent pathways starting from $RS(O)H$ -
 240 $-ROOH$ is presented in **Figure 6**.

241

242



243

244 **Figure 6:** Gibbs free energy profile illustrating the two reactions that oxidized hexanethiol
 245 can undergo with MeOOH.

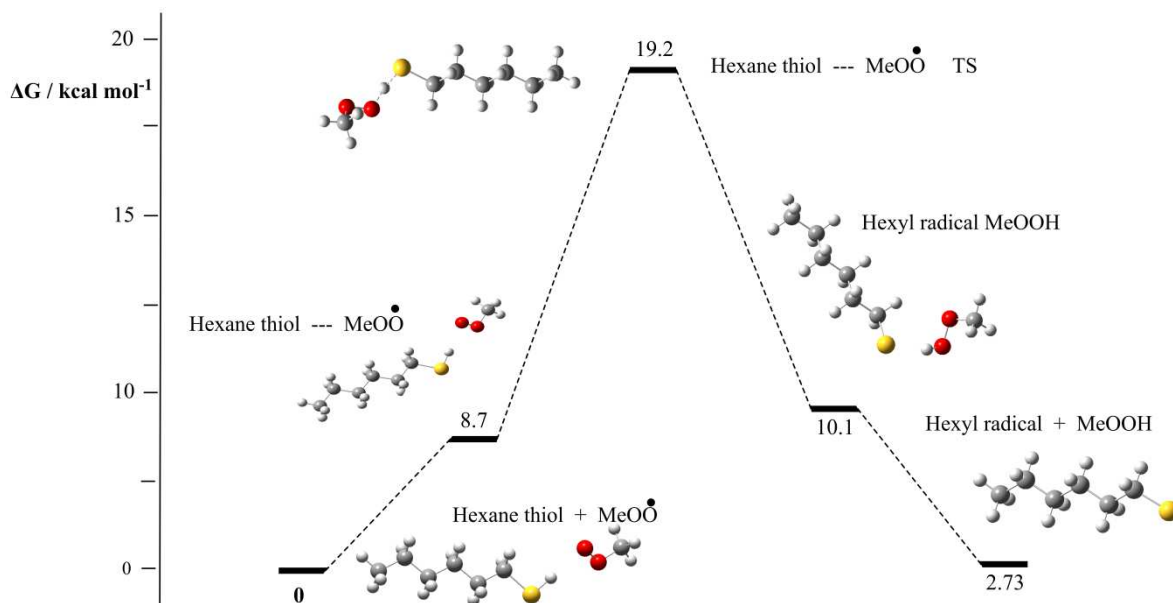
246

247 The activation free energy for the oxidation of HexS(O)H to HexS(O)(O)H is $37.4 \text{ kcal mol}^{-1}$.
 248 (path B in Figure 6) This is significantly larger than the activation free energy for re-
 249 arrangement to form a sulfenic acid, HexS(OH) ($16.7 \text{ kcal mol}^{-1}$). Thus, the calculation
 250 suggests that re-arrangement occurs prior to further oxidation. Activation free energies for
 251 subsequent oxidations starting from a sulfenic acid are $23.1 \text{ kcal mol}^{-1}$ and $33.1 \text{ kcal mol}^{-1}$ for
 252 the transformation of HexSOH to HexS(O)(O)H and HexS(O)(O)H to HexS(O)(O)OH
 253 respectively. This data is provided in the supporting information.

254

255 We note that in a recent publication, we showed that the concentration of hydroperoxides
 256 decreases significantly with each successive oxidation.^{53,54} As a consequence, each successive
 257 oxidized sulfur species will be present in a much lower concentration than the initial non-

258 oxidized sulfur compound. Thus, it is likely that only the initial oxidation reaction will be
259 important in jet fuel given that further oxidation reactions will occur with a low frequency. Of
260 course, the significance of each successive reaction may increase for fuels containing a higher
261 concentration of hydroperoxides.



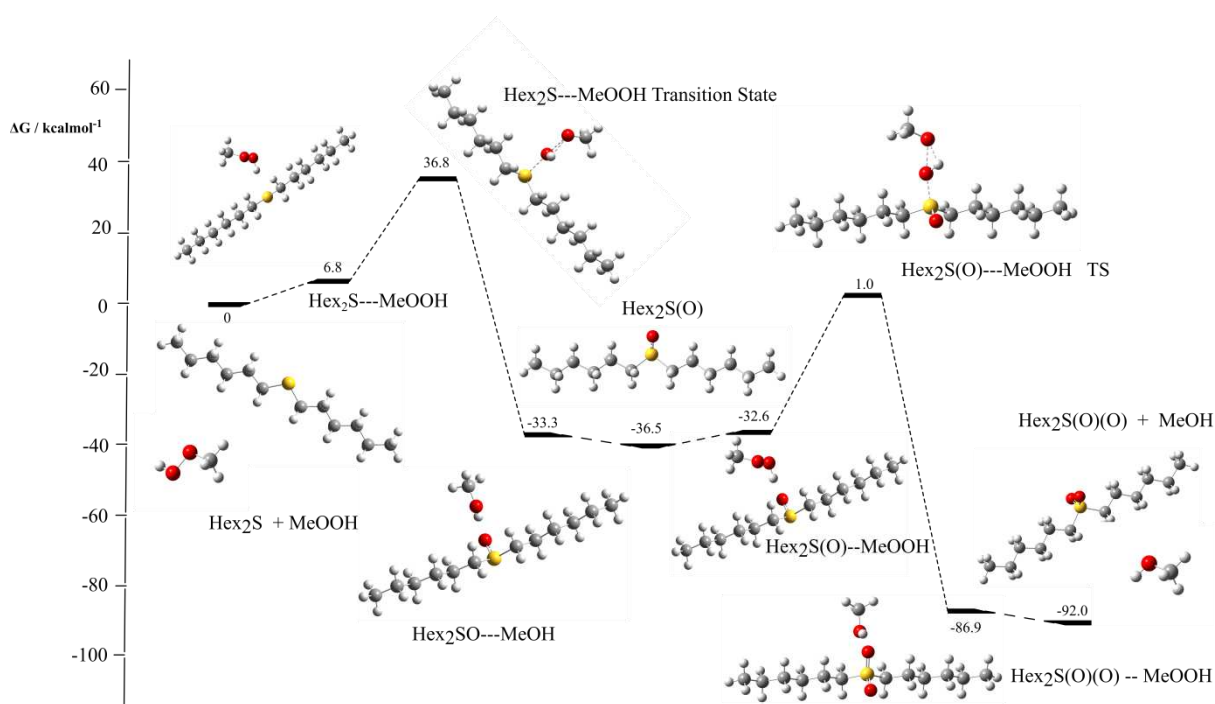
262
263 **Figure 7:** Gibbs free energy profile for the reaction of hexanethiol and MeOO•.

264
265 Our final consideration regarding thiol reactivity concerned the possibility of thiols reacting
266 with peroxy radicals. Given that sulfur species have been shown to produce widely varying
267 effects on jet fuel autoxidation and deposition, it is likely that they react with more than just
268 hydroperoxides. Consequently, we investigated the reaction of hexane thiol and MeOO•. The
269 Gibbs free energy profile is shown in **Figure 7**. This reaction has a free activation energy of
270 19.2 kcal mol^{-1} and the overall reaction is only marginally uphill. This data suggests that thiols
271 can react with peroxy radicals and generate hydroperoxides and thiyl radicals. Further
272 investigations of the reactions that thiyl radicals can undergo will be detailed in **section 3.4**.

273

274 Sulfides were the next class of compounds to be investigated. Sulfides have the general formula
 275 RSR. They can undergo similar reactions with hydroperoxides as observed for thiols.
 276 However, in this case, there is a maximum of two oxidation steps. Moreover, sulfenic acid
 277 cannot be formed because there is no terminal –SH bond. The free energy profile for the
 278 reaction of hexyl sulfide and MeOOH is shown in **Figure 8**.

279



280

281 **Figure 8:** Gibbs free energy profile for the reaction of hexylsulfide and MeOOH.

282

283 The activation free energy for the oxidation of Hex₂S to Hex₂S(O) is 36.8 kcal mol⁻¹, which is
 284 similar to the activation free energy for the oxidation of hexane thiol. The second oxidation
 285 reaction has an activation free energy of 37.5 kcal mol⁻¹. Both of these reactions are permissible
 286 given the standard conditions experienced during autoxidation (temperatures of between 140
 287 and 300 °C). Corresponding activation energies for the reaction of CHP and Hex-S-Hex are
 288 36.7 kcal mol⁻¹ and 37.4 kcal mol⁻¹ for the conversion of Hex₂S to Hex₂S(O) and Hex₂S(O) to
 289 Hex₂S(O)(O) respectively.

290 **Table 3:** Collated Gibbs free activation energies for the first and second oxidation reactions
291 between three sulfides and MeOOH.

Sulfur species	Gibbs free activation energy / kcal mol ⁻¹	
	First oxidation	Second oxidation
phenylsulfide	39.3	39.3
ⁿ butylsulfide	36.6	37.4
^t butylsulfide	35.3	35.0

292
293 **Table 3** contains calculated Gibbs free activation energies for the reaction of three further
294 sulfides with MeOOH. The chosen sulfides contained phenyl, ⁿbutyl and ^tbutyl substituents.
295 As can be seen in **Table 3**, the first and second oxidation reactions of ⁿbutylsulfide and MeOOH
296 are very similar to those calculated in **Figure 8**. As the only change between ⁿbutylsulfide and
297 hexylsulfide is a reduction in the carbon chain length, this result is to be expected. The first
298 and second oxidation reactions of phenylsulfide and MeOOH are higher than those reported
299 for hexylsulfide

300

301 **3.3 Reactions of thiophenes, benzothiophenes and dibenzothiophenes**

302

303 The speciation analysis that we carried out on jet fuel indicated that substituted benzo- and
304 dibenzothiophenes were the most observed species after thiols, sulfides and disulphides.
305 However, the analysis did not provide information as to the exact nature and location of the
306 substitutions. Due to this, we next investigated non-substituted analogues (ie, thiophene,
307 benzothiophene and dibenzothiophenes) as models for these species.

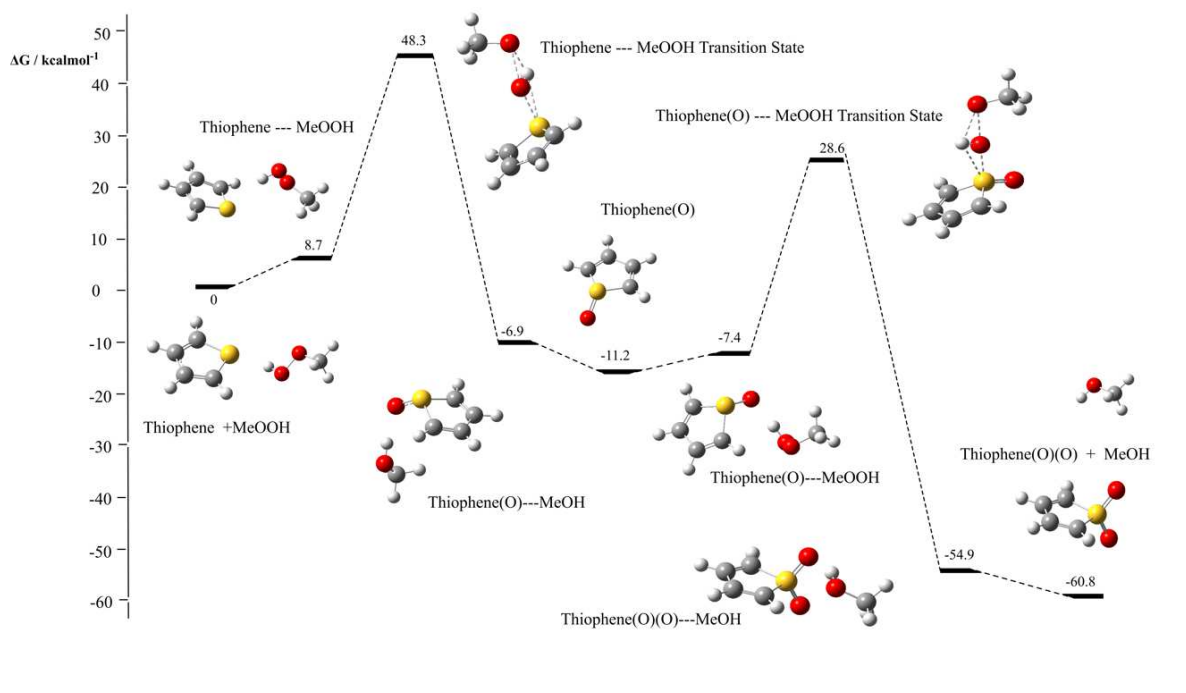
308

309 **Figure 9** shows the calculated energy profile for the reaction of thiophene and MeOOH.

310 Thiophene can undergo two successive oxidations with hydroperoxides as was the case for

311 sulfides (RSR). The activation free energy for the oxidation of thiophene to thiophene(O) is
312 48.3 kcal mol⁻¹. The subsequent oxidation reaction to thiophene(O)(O) has an activation free
313 energy of 39.7 kcal mol⁻¹.

314



315

316 **Figure 9:** Gibbs free energy profile for the reaction of thiophene and MeOOH.

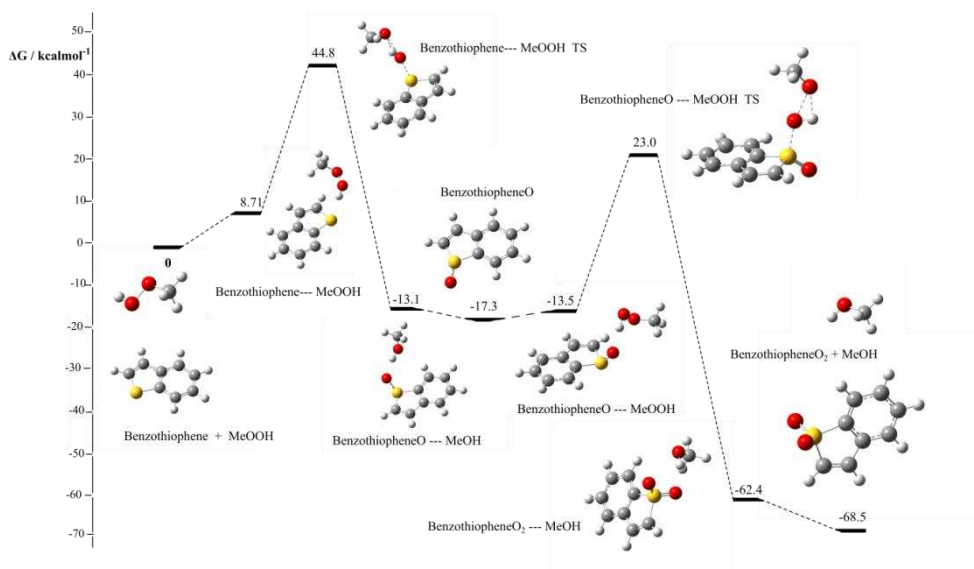
317

318 As can be seen in **Figures 10** and **11**, similar activation free energies were calculated for the
319 reactions of both benzothiophene and dibenzothiophene with MeOOH. The calculated
320 activation free energies for the first and second oxidation of benzothiophene are 44.8 and 40.3
321 kcal mol⁻¹. The first and second oxidation reactions of dibenzothiophenes have activation free
322 energies of 43.2 and 40.7 kcal mol⁻¹.

323

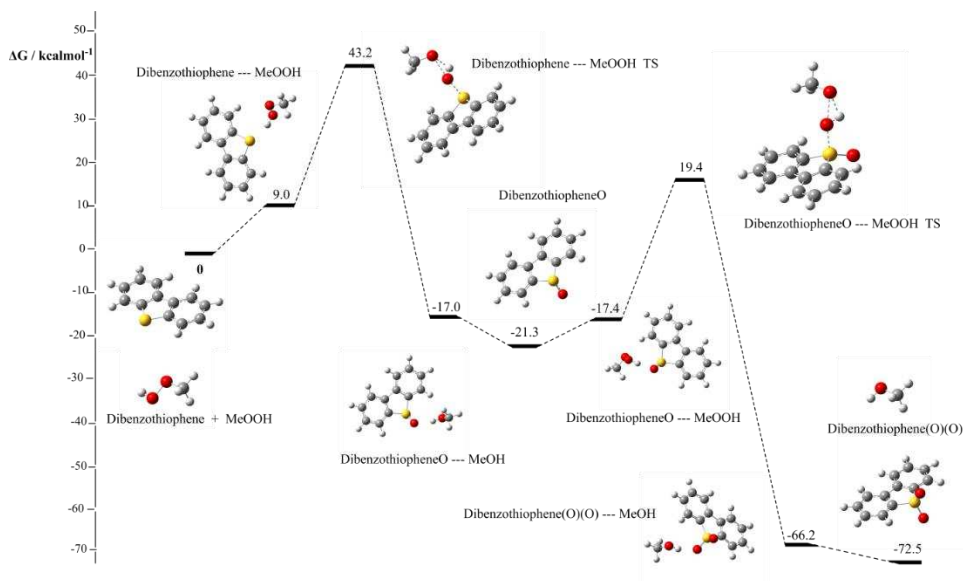
324 The calculated activation free energies for the first reaction between hydroperoxides and each
325 of thiophene, benzothiophene and dibenzothiophene are higher than those calculated for
326 hexylsulfide. This is attributed to the loss in aromaticity during the oxidation.⁵⁵

327



328

329 **Figure 10:** Gibbs free energy profile for the reaction of benzothiophene and MeOOH.



330

331 **Figure 11:** Gibbs free energy profile for the reaction of dibenzothiophene and MeOOH.

332

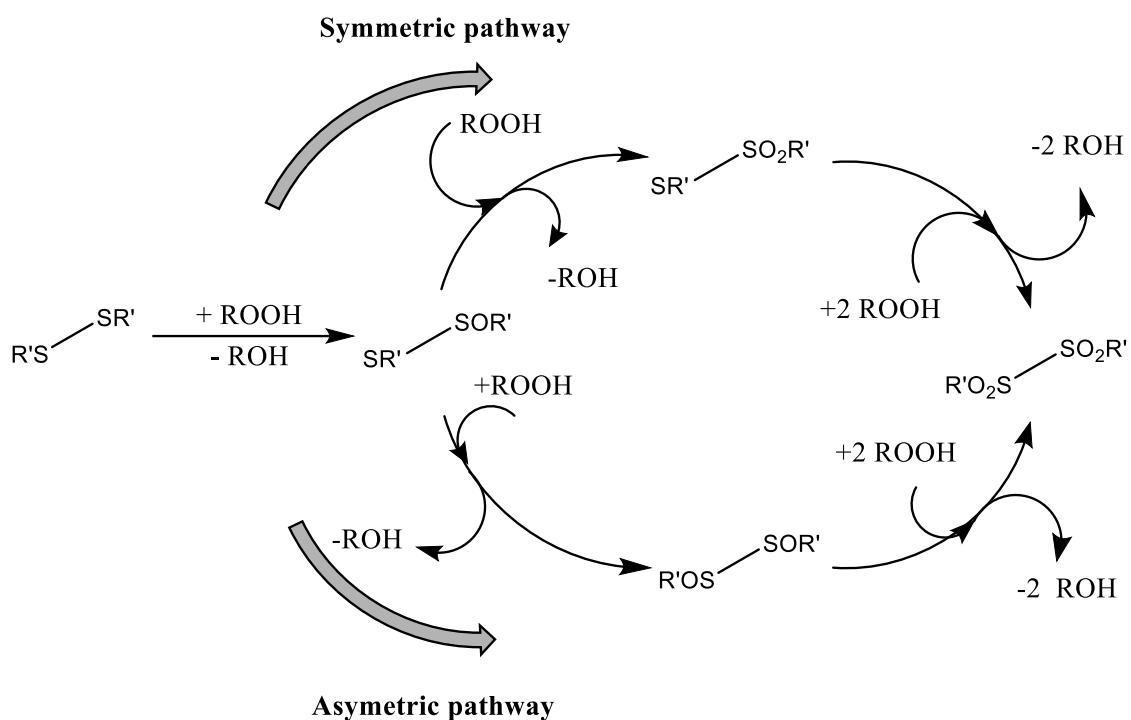
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334

335

336

337



339

340 **Scheme 2:** Two potential mechanistic pathways leading to formation of $\text{RSO}_2\text{SO}_2\text{R}$.

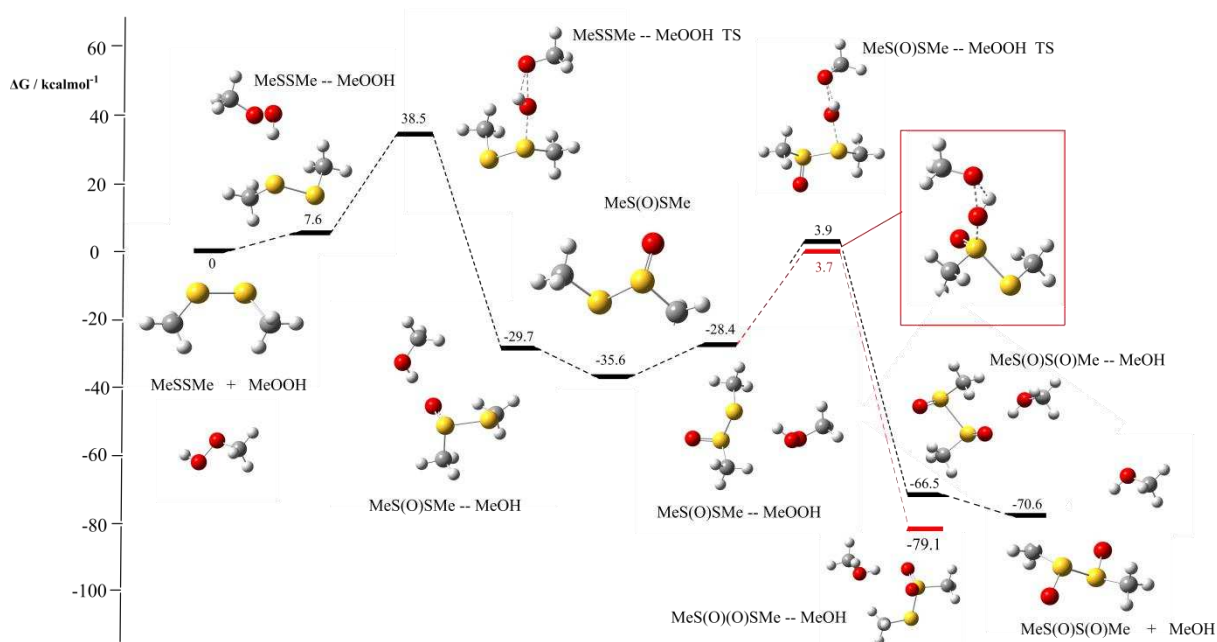
341

342 Disulfides, with the general formula RSSR can potentially undergo as many as four oxidations
 343 with hydroperoxides. After the first oxidation, the second can occur at either the non-oxidized
 344 or oxidized sulfur atom as summarized in **Scheme 2**, leading to a second oxidation species that
 345 is either symmetric or asymmetric. However further oxidations will lead to the same end
 346 product.

347

348 The free energy profile for the reaction of dimethyldisulfide and MeOOH up to and including
 349 the second oxidation *via* the two different routes is shown in **Figure 12**. The initial oxidation
 350 has an activation energy of $38.5 \text{ kcal mol}^{-1}$, which is comparable to the activation free energy
 351 calculated for RSH oxidation in **Figure 8**. The activation free energies for the 2nd oxidation
 352 are 39.3 and $39.5 \text{ kcal mol}^{-1}$ for symmetric and asymmetric oxidation respectively. This

353 indicates that there is almost no kinetic preference as to which sulfur atom will be oxidized
 354 after the initial oxidation
 355



356
 357 **Figure 13:** Gibbs free energy profile for the reaction of MeSSMe and MeOOH up to and
 358 including the second oxidation.

359
 360 Corresponding activation free energies for the reaction of CHP and MeSSMe are 38.2 kcal mol⁻¹
 361 and 39.2 kcal mol⁻¹ for the conversion of MeSSMe to MeS(O)SMe and MeS(O)SMe to
 362 MeS(O)S(O)Me respectively.

363
 364 Oxidation is not the only reaction that disulfides can undergo. In particular, disulfides contain
 365 a diheteroatomic bond in much the same way as hydroperoxides and could potentially undergo
 366 a fission reaction to form two thiyl radical species which are known to be very reactive.
 367 Breaking of the disulfide bond will be easier if there is a significant weakening of the bond
 368 during the autoxidation process, as indicated by a lengthening of the S-S bond and
 369 consequentially change in the bond enthalpy.

370

371 The calculated bond enthalpies and corresponding S-S bond lengths for all of the potential
372 species along the reaction coordinate are given in **Table 4**. The calculated S-S bond enthalpies
373 for RSSR are 54 and 53 kcal mol⁻¹ for R=Me and ⁿBu respectively. These values are
374 comparable to literature values for disulfide bonds that are generally around 50-60 kcal mol⁻¹
375 ^{1,56} **Table 4** shows that the first and second oxidations appreciably weaken the disulfide bond.
376 Therefore fission of the S-S bond under autoxidation conditions should be considered in more
377 detail.

378 **Table 4:** S-S bond lengths and enthalpies for a series of disulfide species.

Disulfide species	Bond enthalpy / kcal mol ⁻¹		Bond Length / Å	
	R = Me	R = ⁿ Bu	R = Me	R = ⁿ Bu
RSSR	54	53	2.068	2.092
RSOSR	32	32	2.167	2.235
RSOSOR	9	11	2.329	2.387
RSO ₂ SOR	18	18	2.310	2.389
RSO ₂ SO ₂ R	25	24	2.285	2.343

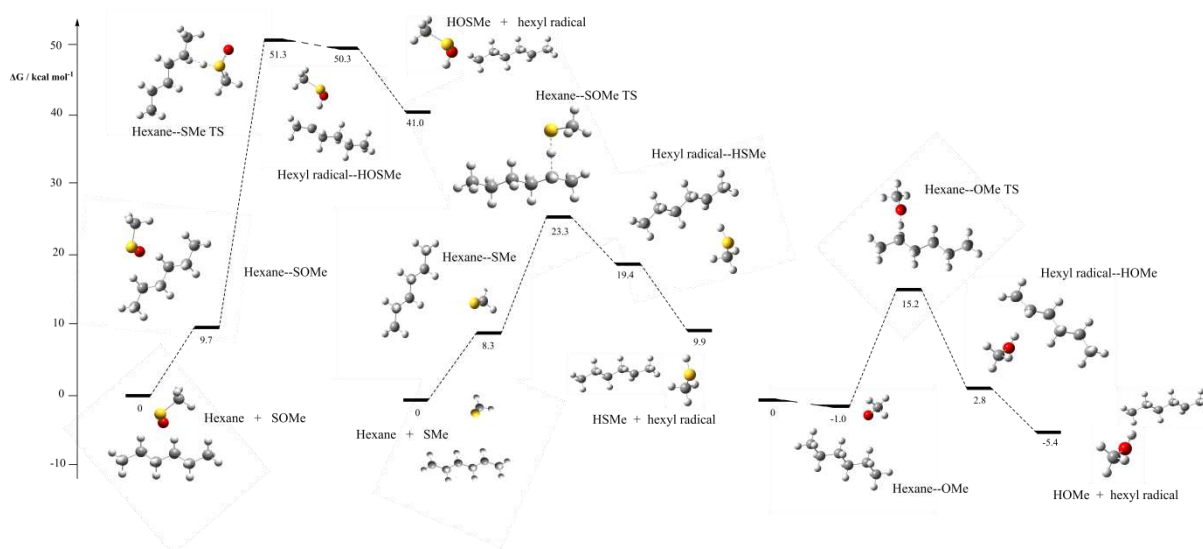
379

380 Our calculations show that the S-S bond significantly weakens upon successive oxidations.
381 Even after a single oxidation, the activation free energy for the cleavage of the S-S bond
382 becomes competitive with subsequent oxidations. Thus, the likely products from bond fission
383 in these species are RSO● and RS●. Indeed, comparison with the activation free energy for
384 RS(O)S(O)R formation, it is as likely to undergo bond fission than react with hydroperoxide.
385 This is further substantiated by comparison of the frequency factors for both reactions (1.2E+22
386 for bond fission and 1.8E+06 for the oxidation reaction). Whilst RSO₂● can potentially form,
387 the weakness of the S-S bond in RSOSOR makes this unlikely. We note that RSO● and RS●
388 are relatable to the radical species formed during hydroperoxide fission, RO● and HO●, which
389 are known to have critical roles in the autoxidation mechanism in fuels. Thus, it was

390 investigated whether any of the radicals that could potentially originate from disulfide fission
391 could react with the bulk fuel in a similar way to peroxy radicals.

392

393 The free energy profile for the reaction of $\text{MeSO}\bullet$ and $\text{MeS}\bullet$ with bulk fuel is shown in **Figure**
394 **13**. For comparison, an equivalent profile was calculated for the reaction of $\text{MeO}\bullet$ and bulk
395 fuel. In these reactions, hexane was used as a model for the hydrocarbons typically found in
396 jet fuel to reduce the computational cost of the calculations. This approach can be justified as
397 the reduction in chain length is not expected to have a significant effect on the reaction at the
398 C2 carbon in the chain based upon previous studies.⁵³

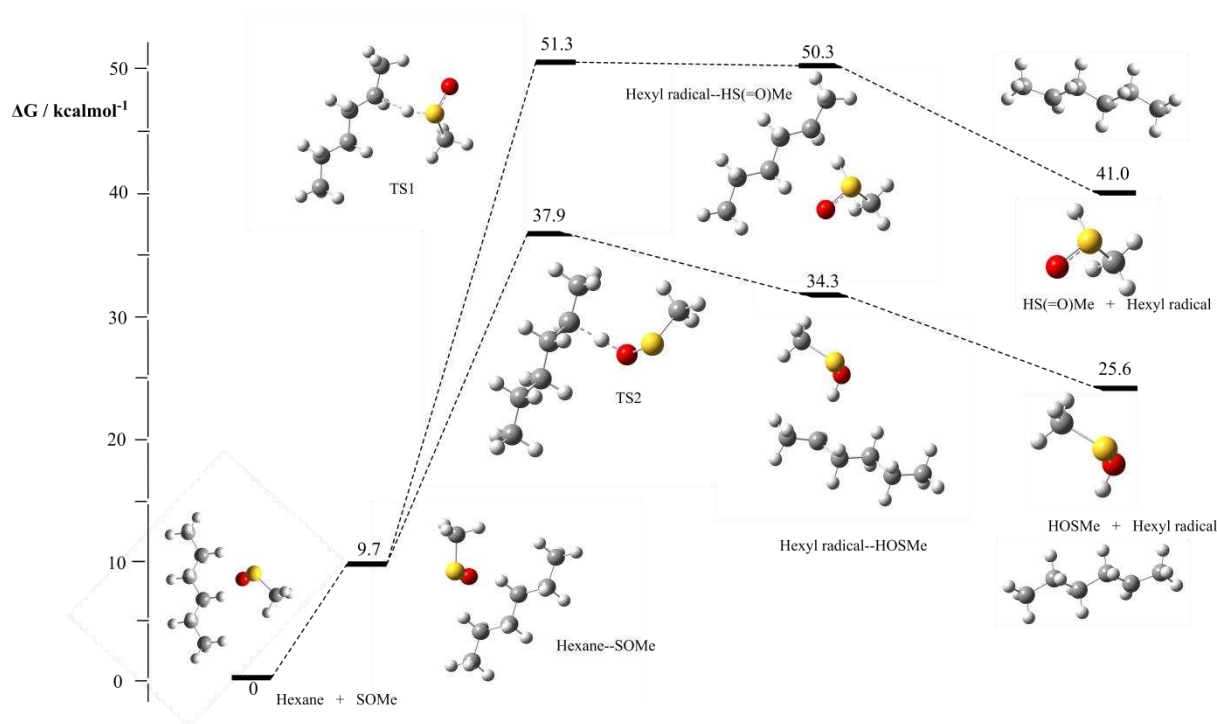


399

400 **Figure 13:** Gibbs free energy profile for the reaction of (left to right) $\text{MeS(O)}\bullet$, $\text{MeS}\bullet$ and
401 $\text{MeO}\bullet$ with hexane.

402 The activation free energies for the reaction of $\text{MeS}\bullet$, $\text{MeS(O)}\bullet$ and $\text{MeO}\bullet$ with bulk fuel are
403 calculated to be 51.3, 23.3 and 15.2 kcal mol^{-1} , respectively. Thus, the calculated activation
404 free energy is appreciably larger for any of the sulfur species than for $\text{MeO}\bullet$. Moreover, the
405 reactions involving sulfur radicals are significantly endothermic. This indicates that the sulfur
406 radical is better stabilized than a carbon radical.

407 In the above it was assumed that hydrogen abstraction happens through sulfur, i.e. that the
408 radical character is localized there. However, a reaction could also happen through the oxygen
409 atom. A comparison of the two energy profiles is shown in **Figure 14**.



410

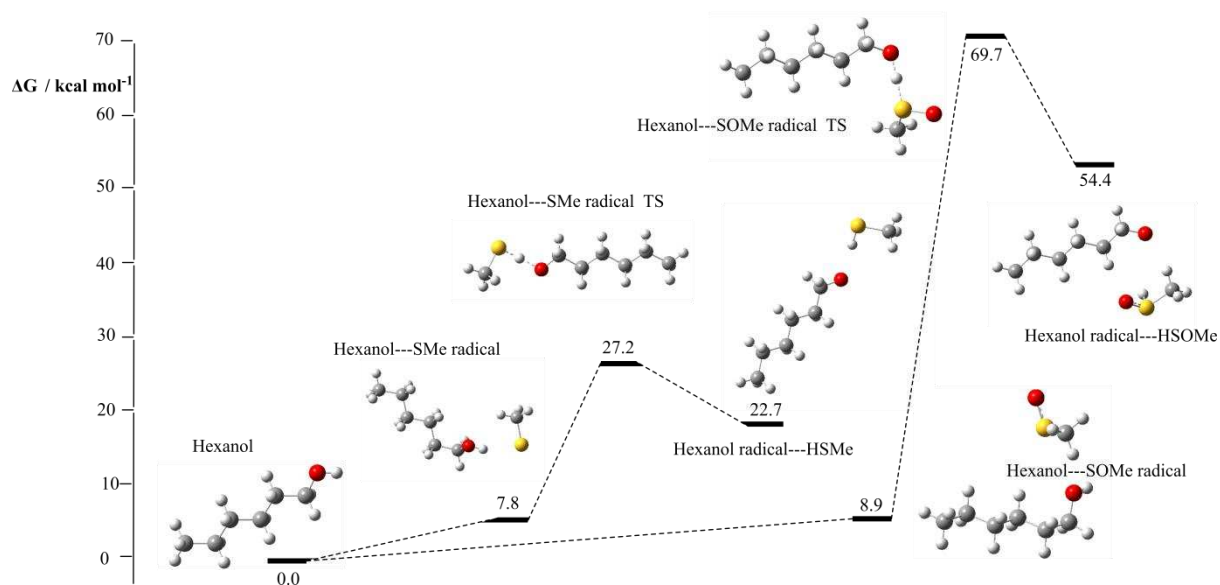
411 **Figure 14:** Gibbs free energy profile for the reaction of $\text{MeS(O)}\bullet$ and $\text{MeS}\bullet(\text{O})$ with
412 hexane.

413 As can be seen, abstraction by oxygen is preferred over sulfur (activation free energies 37.9
414 and 51.3 kcal mol⁻¹, respectively). However, even hydrogen abstraction by oxygen is still
415 endothermic. Overall, the data in **Figures 13** and **14** suggest that any radical sulfur species
416 formed from disulfide fission react less efficiently than radicals resulting from hydroperoxide
417 fission.

418

419 Our final consideration was whether the thiyl radicals, $\text{MeS}\bullet$ and $\text{MeS(O)}\bullet$, could potentially
420 react with alcoholic species in the fuel. **Figure 15** shows the gibbs free energy profile for both
421 reactions. The reaction of hexanol and $\text{MeS}\bullet$ has a low kinetic barrier and will be

422 surmountable given typical autoxidation conditions. In contrast, the reaction of hexanol and
423 $\text{MeS(O)}\bullet$ has a significantly higher Gibbs free activation energy, which is likely to be
424 prohibitive. In both cases, the reaction is endothermic, which suggests that the radical is better
425 stabilized when localized on the sulfur of the respective thiyl radical than the oxygen on
426 hexanol.



428 **Figure 15:** Gibbs free energy profile for the reaction of both $\text{MeS(O)}\bullet$ and $\text{MeS}\bullet(\text{O})$ with
429 hexanol.

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437 3.5 Mechanistic implications

438

439 The work undertaken herein has provided greater insight into the elementary reactions involved
440 in the oxidation of sulfur species. Oxidation of sulfur species can lead to the formation of
441 alcohols, sulfones, sulfoxides and sulfonic acids. Moreover, cleavage of disulfides can form
442 thiyl radicals.

443

444 The presence of various sulfur species in jet fuel has the potential to retard the rate of
445 autoxidation by reacting with hydroperoxides. The larger data set, consisting of a range of
446 sulfur species reacting with different hydroperoxides, provides a more robust framework for
447 the improvement of current chemical kinetic models. As mentioned previously, these models
448 are of great importance for predicting the rate of autoxidation in jet fuels. In the widely used
449 Kuprowicz mechanism there is a single reaction where sulfur is involved, which does not
450 differentiate between the different classes of sulfur species in the fuel (equation 1).¹⁰ This work
451 would allow for the expansion of this single reaction into a series of reactions that not only
452 differentiate between the classes of sulfur compounds but also the specific individual reactions
453 that each one might undergo.

454

455 To more accurately represent the reactions that potentially could occur, we would propose
456 supplementing equation 1 in the current kinetic mechanism with the reactions outlined in **Table**
457 **5**, using the lumped data in **Table 6**. Reactions 1 to 4 describe the reactions of thiols and
458 hydroperoxides. Reactions 5 and 6 describe the reactions of both thiophenes and sulfides with
459 hydroperoxides. The reactions of disulfides and hydroperoxides are described by reactions 7-
460 10.

461

Table 5: Proposed elementary steps to improve current chemical kinetic mechanisms

Elementary reaction step		Label
RSH + ROOH	\longrightarrow	RSHO + ROH Reaction 1
RSHO + ROOH	\longrightarrow	RSOH + ROOH Reaction 2
RSOH + ROOH	\longrightarrow	RSO ₂ H + ROH Reaction 3
RSO ₂ H + ROOH	\longrightarrow	RSO ₃ H + ROH Reaction 4
RSR + ROOH	\longrightarrow	RSOR + ROH Reaction 5
RSOR + ROOH	\longrightarrow	RSO ₂ R + ROH Reaction 6
RSSR + ROOH	\longrightarrow	RSOSR + ROH Reaction 7
RSOSR + ROOH	\longrightarrow	RSOSOR + ROH Reaction 8
RSOSOR + ROOH	\longrightarrow	RSO ₂ SOR + ROH Reaction 9
RSO ₂ SOR + ROOH	\longrightarrow	RSO ₃ SO ₃ R + ROH Reaction 10

463

464 The reactions, lumped activation energies and Arrhenius data for the elementary reactions
 465 investigated herein are summarized in **Table 6**. With appropriate validation against
 466 experimental data and testing to deduce the importance of each individual reaction, it is hoped
 467 that this data will lead to a more accurate chemical kinetic model for fuel autoxidation.

468

469 **Table 6.** Lumped data for proposed steps to improve current chemical kinetic mechanisms.

470

(COH = Cumene hydroxide)

Sulfur species (Hydroperoxide)	Elementary reaction step		E _a kcal mol ⁻¹	A mol ⁻¹ s ⁻¹	Source	
	SH + R'OOH	\longrightarrow	Products _{SH}	18	3E+09	Ref 10
Thiols (MeOOH)	RSH + MeOOH	\longrightarrow	RSHO + MeOH	41.6	7.8E+05	This work
	RSHO + MeOOH	\longrightarrow	RSOH + MeOOH	19.2	4.0E+09	This work
	RSOH + MeOOH	\longrightarrow	RSO ₂ H + MeOH	31.0	5.1E+09	This work
	RSO ₂ H + MeOOH	\longrightarrow	RSO ₃ H + MeOH	38.9	1.9E+11	This work
Thiols (CHP)	RSH + CHP	\longrightarrow	RSHO + COH	40.9	2.9E+05	This work
	RSHO + CHP	\longrightarrow	RSOH + CHP	23.9	1.9E+07	This work
	RSOH + CHP	\longrightarrow	RSO ₂ H + COH	33.9	6.0E+08	This work
	RSO ₂ H + CHP	\longrightarrow	RSO ₃ H + COH	41.1	1.9E+09	This work
Thiols (^tBuOOH)	RSH + ^t BuOOH	\longrightarrow	RSHO + ^t BuOH	42.4	3.6E+05	This work
	RSHO + ^t BuOOH	\longrightarrow	RSOH + ^t BuOOH	20.2	3.5E+10	This work
	RSOH + ^t BuOOH	\longrightarrow	RSO ₂ H + ^t BuOH	33.7	2.9E+10	This work
	RSO ₂ H + ^t BuOOH	\longrightarrow	RSO ₃ H + ^t BuOH	42.3	3.0E+11	This work
Sulfides (MeOOH)	RSR + MeOOH	\longrightarrow	RSOR + MeOH	36.9	9.6E+06	This work
	RSOR + MeOOH	\longrightarrow	RSO ₂ R + MeOH	37.3	2.3E+09	This work
Sulfides	RSR + CHP	\longrightarrow	RSOR + COH	36.7	6.8E+05	This work

(CHP)	RSOR + CHP	—————>	RSO ₂ R + COH	37.4	E+08	This work
Sulfides	RSR + 'BuOOH	—————>	RSOR + 'BuOH	37.9	6.8E+05	This work
('BuOOH)	RSOR + 'BuOOH	—————>	RSO ₂ R + 'BuOH	39.6	2.2E+08	This work
Disulfides	RSSR + MeOOH	—————>	RSOSR + MeOH	39.9	6.5E+04	This work
(MeOOH)	RSOSR + MeOOH	—————>	RSOSOR + MeOH	39.3	1.3E+06	This work
	RSOSOR + MeOOH	—————>	RSO ₂ SOR + MeOH	34.3	2.0E+09	This work
	RSO ₂ SOR + MeOOH	—————>	RSO ₃ SO ₃ R + MeOH	38.3	1.1E+08	This work
Disulfides	RSSR + CHP	—————>	RSOSR + COH	41.9	1.3E+05	This work
(CHP)	RSOSR + CHP	—————>	RSOSOR + COH	41.6	3.8E+06	This work
	RSOSOR + CHP	—————>	RSO ₂ SOR + COH	36.6	8.0E+06	This work
	RSO ₂ SOR + CHP	—————>	RSO ₃ SO ₃ R + COH	38.9	1.4E+06	This work
Disulfides	RSSR + 'BuOOH	—————>	RSOSR + 'BuOH	39.8	5.8E+04	This work
('BuOOH)	RSOSR + 'BuOOH	—————>	RSOSOR + 'BuOH	40.6	5.4E+06	This work
	RSOSOR + 'BuOOH	—————>	RSO ₂ SOR + 'BuOH	35.2	8.5E+06	This work
	RSO ₂ SOR + 'BuOOH	—————>	RSO ₃ SO ₃ R + 'BuOH	38.9	2.2E+06	This work
Thiophenes and Benzothiophenes	RSR + MeOOH	—————>	RSOR + MeOH	45.4	8.3E+04	This work
(MeOOH)	RSOR + MeOOH	—————>	RSO ₂ R + MeOH	40.8	4.6E+09	This work
Thiophenes and Benzothiophenes	RSR + CHP	—————>	RSOR + COH	47.6	1.3E+04	This work
(CHP)	RSOR + CHP	—————>	RSO ₂ R + COH	39.9	7.6E+08	This work
Thiophenes and Benzothiophenes	RSR + 'BuOOH	—————>	RSOR + 'BuOH	46.3	1.2E+04	This work
('BuOOH)	RSOR + 'BuOOH	—————>	RSO ₂ R + 'BuOH	41.6	4.3E+08	This work

471

472 4 Conclusions

473

474 In this work, we have reported on the reactions of a series of sulfur compounds with model
475 hydroperoxides for those found in jet fuel. Thiols can react with up to four equivalents of
476 peroxides *via* a sulfenic acid to form sulfonic acids and alcohols. In contrast, sulfides can only
477 react with two equivalents of peroxides and form sulfones as opposed to sulfonic acids.

478

479 The reaction of disulfide species with hydroperoxides generally have a higher activation energy
480 compared to thiols and sulfides. These sulfur species can potentially react with multiple
481 equivalents of peroxides to form sulfones. Each successive oxidation acts to weaken the sulfur-
482 sulfur bond, which facilitates the homolytic fission reaction to form thiyl radicals. Our results
483 indicate that these radicals do not favourably react with bulk fuel components

484

485 Once appropriately validated, current chemical kinetic models for the autoxidation of jet fuel
486 will be improved with the inclusion of the more detailed individual reactions investigated here.
487 This flexibility will also allow tailoring of the kinetic scheme to account for specific species of
488 sulfur present in specific fuels.

489

490 **Supporting Information**

491

492 Cartesian coordinates for DFT-optimized structures. Raw GC data resulting from the
493 speciation analysis. Collated thermochemical data for all species investigated.

494

495

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511

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518

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