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Elevated temperature linked to signs associated with sea star wasting disease in a keystone European species, *Asterias rubens*

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1 TITLE

2 Elevated temperature linked to signs associated with sea star wasting disease in a keystone

3 European species, *Asterias rubens*

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6 RUNNING PAGE HEAD

7 Temperature-sensitive SSWD signs in *A. rubens*

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20 ABSTRACT

21 Sea Star Wasting Disease (SSWD) refers to a suite of gross signs affecting Asteroidea species.
22 These include epidermal lesions, everted viscera, arm autotomy, and ultimately, full body
23 disintegration leading to mortality. The common sea star *Asterias rubens* is a keystone
24 species in the coastal Northeast Atlantic and may be susceptible to the disease. While the
25 precise aetiology of SSWD remains poorly understood, environmental instability, including
26 rising sea temperatures, has been linked to SSWD outbreaks. To investigate this connection,
27 an experiment was conducted to quantify disease sign expression in *A. rubens* under
28 elevated temperature. We exposed sea stars to either elevated temperature (18°C) or a
29 control treatment (12°C) for a 14-day period. We quantified the presence of disease signs
30 associated with SSWD, the progression of signs, and survival of individuals. Elevated
31 temperature induced a greater number of signs consistent with SSWD, and also resulted in
32 mortality for some of those animals. Furthermore, larger individuals were more likely to
33 show increased disease signs. Our results provide evidence that signs associated with SSWD
34 increase with elevated temperature.

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37 KEYWORDS

38 Sea Star Wasting Disease, *Asterias rubens*, Asteroidea, Thermal Stress

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42 1. INTRODUCTION

43 Diseases are important drivers of change in the marine environment, with the potential to
44 affect community dynamics and trophic interactions (Burge et al. 2014). These changes can
45 have long-term consequences for ecosystem structuring and function, particularly if
46 affecting foundation, keystone, or ecosystem engineer species (Miner et al. 2018). Due to
47 global climate change, occurrences of large and fast temperature fluxes in marine
48 ecosystems may increase in frequency (IPCC 2021). Warmer oceans and increasingly
49 frequent temperature anomalies have been linked to disease-induced mortality in many
50 marine species (Harvell et al. 2002, Bruno et al. 2007, Ward et al. 2007, Eisenlord et al. 2016,
51 Harvell et al. 2019). Further, if the species affected by the mass-mortality event is a
52 keystone species (sensu Paine 1966), then there may be wider ecological implications
53 (Levitan et al. 2023).

54 The common starfish *Asterias rubens* is a highly abundant asteroid found on the Atlantic
55 coast of Northwest Europe, inhabiting both sub- and intertidal zones (Saier 2001, Gallagher
56 et al. 2008, Agüera et al. 2021). Sea stars in the genus *Asterias* are typical of many asteroids
57 in that their role as keystone predators provides an important control on prey populations.
58 *Asterias rubens* is an important predator for bivalve molluscs, notably the blue mussel,
59 *Mytilus edulis* (Agüera et al. 2020). While *Asterias rubens* is often considered a pest species
60 for mussel cultivators due to predation (Gallagher et al. 2008, Agüera et al. 2020), their
61 controlling effect on bivalves is also a factor in promoting benthic species diversity
62 (Wahlteitz et al. 2023). Thus, disease-induced mortality in *A. rubens* species could therefore
63 have cascading effects on coastal benthic ecology.

64 Sea star wasting disease (SSWD) refers to a suite of gross signs affecting Asteroids, and
65 often co-occurs with changes in behaviour (Miner et al. 2018, Hewson 2021). These signs
66 initially manifest to varying degrees as abnormal posture, reluctance to feed, bloating or
67 pinching of arms, and the appearance of epidermal lesions (Bucci et al. 2017, Hewson 2019,
68 Hewson 2021). Progression of these signs can lead to epidermal ulceration resulting in the
69 exposure of viscera, including pyloric glands and gonads, followed by disintegration of the
70 epidermis, and then death (Staehli et al. 2009, Eisenlord et al. 2016, Hewson et al. 2014).

71 SSWD was first used to describe disease outbreaks in Asteroids on the Pacific coast of the
72 United States in the 1980's & 1990's (Dungan et al. 1982, Eckert et al. 1999). It has since
73 been used to describe mass mortality events occurring on both Pacific and Atlantic coasts of
74 North America (Hewson et al. 2014, Bucci et al. 2017, Hewson et al. 2019). SSWD is now
75 understood to affect over 20 species (Hewson 2021). While a significant mass-mortality
76 event in the Northeast Pacific in 2013/14 brought greater attention to the phenomenon,
77 wasting-like signs of disease, or signs consistent with SSWD, have been observed in many
78 species in prior years. Staehli et al. (2009) reports on a mass die-off of the sea star
79 *Astropecten jonstoni* in the western Mediterranean, while Prestedge (1998) identified a
80 condition affecting *Patiriella vivipara* from Tasmania. Both these cases describe a condition
81 capable of causing severe tissue necrosis which may result in the animals' mortality. As far
82 back as the 19th century, Mead (1898) described a necrotic condition affecting the skin,
83 before progressing through the rest of the body.

84 Questions remain, however, on the precise drivers and cause of SSWD, and the exact
85 aetiology is yet to be fully resolved (Miner et al. 2018, Hewson et al. 2019). Researchers
86 identified a densovirus (*Parvoviridae*) as a candidate pathogen for the disease, however

87 subsequent work has called this into question (Hewson et al. 2014, Hewson et al. 2020,
88 Jackson et al. 2020a, Jackson et al. 2020b).

89 While not ruling out a pathogenic aetiology, the role of environmental stressors has also
90 been proposed (Burge et al. 2014, Hewson et al. 2019). SSWD events have coincided with
91 increased temperature (Dungan et al. 1982, Staehli et al. 2009, Eisenlord et al. 2016, Harvell
92 et al. 2019), although correlations with decreased temperature have also been reported
93 (Menge et al. 2016).

94 More environmental factors have also been investigated, such as an association with
95 precipitation (Hewson et al. 2018), demonstrating that the relationship between SSWD and
96 external stressors is complex. Indeed, it may be that no single aetiology can account for
97 SSWD across both its geographic extent and the species affected (Hewson et al. 2018).

98 SSWD as presently understood is not pathognomic (no uniquely identifying features).

99 Rather, it is a constellation of typically observed signs (Hewson et al. 2019), and it is possible
100 that certain signs associated with SSWD are also indicative of additional stressors or other
101 disease. Furthermore, microbial activity at the animal-water interface has been linked with
102 SSWD and, regardless of ultimate aetiology, these microbes are themselves influenced by
103 factors such as organic matter availability, precipitation, and temperature (Aquino et al.
104 2021).

105 *Asterias rubens* is known to manifest signs indicative of SSWD such as lesions, bloating and
106 loss of turgor (Menge 1979, Wahlteiz et al. 2023). Further, *A. rubens* shows reduced
107 feeding rates, energy uptake and growth under elevated temperature (Morón Lugo et al.
108 2020, Rühmkorff et al. 2023), highlighting sensitivity to thermal change. With SSWD linked
109 to environmental instability and thermal fluctuation, the susceptibility of *A. rubens* to SSWD

110 necessitates a deeper understanding of this phenomenon. With mass mortality events
111 reported in various asteroidea species, disease outbreak in this keystone benthic predator
112 may have cascading ecological implications (Harvell et al. 2019, Hewson et al. 2019).

113 The aim of this study was to measure the effect of elevated temperature on the prevalence
114 of signs corresponding to SSWD in *A. rubens*. We conducted a laboratory experiment to test
115 whether rapid temperature elevation could induce signs associated with SSWD. The
116 presence of disease signs was monitored, as well as survival rates to assess the role that
117 elevated temperature plays in inducing signs consistent with SSWD over time. Disease sign
118 progression at elevated temperature was predicted to increase over the course of the
119 experiment, with prolonged exposure resulting in a greater frequency on gross signs such as
120 lesions, bloating, loss of body turgor and arm autotomy when compared to the control
121 treatment. Further, we hypothesised that the abrupt increase in temperature experienced
122 by sea stars would decrease survival.

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131 2. MATERIALS & METHODS

132 2.1. Sea Star collection

133 Adult *Asterias rubens* were collected in October 2022 from Carlingford Lough, Irish Sea,
134 using sea star mops from cultivated mussel beds (*sensu* Calderwood et al. 2016). Annual sea
135 temperatures in Carlingford Lough range between 3°C and 20°C (AFBI 2015), and average
136 water temperature for October 2022 were 13.26°C (SeaTemperatures 2023). Animals were
137 transferred in aerated containers holding chilled seawater to the Queen’s University Belfast
138 Marine Laboratory in Portaferry within 90 minutes of collection. At the laboratory,
139 specimens were placed in a covered outdoor tank supplied by aerated UV and sand-filtered
140 seawater drawn from the marine Strangford Lough at a rate of approximately 250 Lhr⁻¹. Sea
141 star condition was monitored for one week, and deceased or damaged individuals were
142 removed. Animals were fed *Mytilus edulis ad libitum* during this stage.

143 Two weeks prior to trial start, sea stars with grossly normal appearance were transferred to
144 an acclimation tank in a temperature-controlled room under a 12hr/12hr light/dark cycle.
145 Grossly normal individuals were identified by the lack of typical gross SSWD signs, *i.e.* no
146 epidermal lesions, turgid body posture, and even taper of arms (Fig. 1). The acclimation tank
147 was supplied by the same flow-through seawater system, and temperature maintained at
148 12°C for the period of acclimation. Halfway through the acclimation phase, sea stars were
149 fed again *ad libitum* with *M. edulis*, with uneaten animals and empty shells removed the
150 following day.

151 2.2. Experimental set-up

152 Thirty-six five litre experimental units (opaque plastic container with translucent lid) were
153 evenly distributed between four tanks. Each large tank was treated with a 300-Watt

154 aquarium heater below a platform, allowing the large tanks to act as water baths. Prior to
155 the experiment, each unit was sterilised with 70 % ethanol solution and cleaned, before
156 being filled with fresh UV and sand-filtered seawater. Each water bath included a 250 Lhr⁻¹
157 pump to recirculate water, ensuring even heat distribution. Experimental units were placed
158 inside the water baths, and pressurised air was supplied to each to oxygenate the water
159 inside the units. Experimental units were self-contained in a closed system, therefore
160 limiting cross-contamination between sea stars. To maintain water quality, daily water
161 changes were performed from separate water baths maintained in temperature and salinity
162 equilibrium with both temperature treatments.

163 Two temperature treatments were assigned between the four water baths, with a target
164 temperature of 12°C representing a continuation of the acclimatisation temperature
165 regime, and an 18°C target temperature representing a 6°C increase to which sea stars were
166 exposed for a 14-day observation period. Although within the annual temperature range
167 experienced by *A. rubens* at the collection site, animals were subjected to an immediate
168 change in temperature to assess whether an abrupt increase would induce signs of SSWD.

169 2.3. Experimental Procedure

170 On day 0 of the experiment, 36 sea stars were randomly assigned to individual experimental
171 units. 18 *A. rubens* were exposed to each temperature regime, with these subdivided
172 between two baths for each treatment. Sea star radius was measured as the length from
173 the tip of the longest arm to the mouth at the centre of the oral disc. All animals presented
174 grossly normal appearance upon induction into the trial.

175 2.4. Data Collection

176 2.4.1. Disease Status and Severity

177 Sea stars were assessed every 24 hours, and gross signs of disease were recorded for each
178 animal. Overall appearance state for living animals was recorded as a binary assessment of
179 grossly normal/grossly abnormal i.e., those expressing zero signs or those expressing any
180 sign associated with SSWD disease (Fig. 1.).

181 Severity of disease was categorised across seven variables corresponding to commonly
182 reported SSWD signs (Bucci et al. 2017, Hewson 2019, Hewson 2021): loss of turgor, arm
183 autotomy, bloating/pinching on the arms or disc, presence of epidermal lesions, contortion
184 of the arms, eversion of viscera (pyloric glands or gonads), and gross disintegration of the
185 epidermis (Fig. 1b-d). Each of these signs was scored out of a maximum with respect to sign
186 characteristics. For signs that affected arms & central disc (bloating/pinching, epidermal
187 lesions, eversion of viscera, gross disintegration) this score ranged from 0 to a maximum of
188 6; 1 point for sign present on each arm, plus the central disc. For signs that only affected
189 arms (contortion, arm autotomy), the maximum score was 5 (Fig. 1e.). Loss of turgor was a
190 binary score with 0 being absent, and 1 being present. After each 24-hour period, a sum
191 score was calculated for each sea star based on the presence and extent of each of the signs
192 associated with SSWD, thus approximating disease severity for an individual. To track signs
193 through each day of observation, arms were ordered in a clockwise direction around the
194 central disc, with the madreporite allowing for orientation of the start-point at arm no. 1
195 (Fig. 1a.).

196 Daily temperature and salinity measurements were recorded for each experimental unit.

197 For the control treatment, the mean temperature for the duration of the study was 12.40°C
198 (s.d. = 0.14), while for the elevated temperature treatment a mean of 18.24°C (s.d. = 0.13)

199 was recorded. Mean salinity for all experimental units for the duration of the trial was
200 36.11ppt (s.d. = 0.14).

201 2.4.2. Disease Category

202 With principal focus on SSWD outbreaks centred on the Pacific coast of North America,
203 several resources have been made available for identification of disease in species local to
204 the region. Guides have been produced for both the ochre sea star (*Pisaster ochraceus*) and
205 the mottled star (*Evasterias troschelii*). The disease guides (MARINe 2018) utilise a
206 categorisation system with five stages of disease severity applied to both species (Table 1.).
207 Both these species belong to the family Asteroiidae and have a similar body plan and
208 ecological niche to *Asterias rubens*. Assessing disease severity can therefore be conducted
209 utilising disease categorisation as an alternative framework to the effect of temperature on
210 total disease sign expression.

211 A notable difference between the categorical assessment disease and the sum of all
212 recorded disease signs is that the former does not factor in several signs that are commonly
213 recorded. It does not account for the bloating/pinching of arms, nor does it account for loss
214 of body turgor or arm contortion. Further, the categorical approach was modified such that
215 dead animals were recorded separately.

216 2.4.3. Survival

217 Animals were removed from the trial upon death. Death was determined by total cessation
218 of movement and adhesion of the tube-feet to the surface of the experimental units. In
219 cases where arms had autotomised from the central disc following autotomy, assessment of
220 death was conducted based solely upon the central disc and arms that remained intact.

221 2.5. Statistical analysis

222 Statistical analysis was conducted using R (v4.2.2; R Core Team 2022). For each analysis, a
223 significance level of $p \leq 0.05$ was used. A Kolmogorov-Smirnov test was used to test for
224 normality, and in the case Poisson-distributed data, dispersion tests and tests for zero-
225 inflation were applied.

226 The analyses of SSWD in the two temperature treatments utilised a generalised linear mixed
227 effects model using the *lme4* package (Bates et al. 2015). Initial exploration of data
228 displayed a non-gaussian distribution, necessitating the generalised approach.

229 To test the relationship between healthy and diseased sea stars over time, a generalised
230 linear mixed effects model was produced using a binomial distribution family. This model
231 also incorporated sea star size (radius/mm) as a predictor in the model outcome. For this
232 analysis, data for day 0 was removed, as all sea stars inducted into the trial were grossly
233 normal in appearance. All data associated with the four sea stars that died were also
234 removed from this analysis.

235 To analyse the relationship between disease severity and temperature treatments, a
236 generalised linear mixed effects model was used with a Poisson distribution. To account for
237 repeated measures of individuals over the 14-day trial, a random effect of sea star ID and
238 day was incorporated into the model. For this model, data related to day 0 was omitted as
239 all sea stars presented grossly normal appearance upon induction into the trial. Data
240 associated with the four sea stars that died was also omitted.

241 To test for the effect of temperature on the disease category, the analysis was performed
242 using a Poisson distribution. Although dead sea stars were incorporated into the categorical

243 assessment of disease severity, the four animals that died in the trial were excluded from
244 this model.

245 To compare the survival of sea stars for the duration of the trial a Kaplan-Meier survival
246 curve was calculated. To compare survival between the two treatments we used a Cox
247 Proportional Hazards model.

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262 3. RESULTS

263 3.1. Disease Status

264 Sea star radii did not differ significantly between treatments (Control: 92.00 ± 14.63 mm,
265 Elevated temperature treatment: 89.39 ± 12.24 mm; *t*-test: $t = -0.581$, $df = 32.969$, $p =$
266 0.565). The proportion of animals expressing any sign associated with SSWD was
267 significantly higher in the elevated temperature treatment than in the control, and the
268 proportion of SSWD signs increased significantly over the duration of the trial (Fig. 2; Table
269 2). Furthermore, signs associated with SSWD were significantly more prevalent among
270 larger individuals (Fig. 3; Table 2).

271 3.2. Disease Severity

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273 In an initial interactive model temperature treatment showed a significant effect on total
274 disease sign score ($\beta = -0.667$, *s.e.* = 0.187 , $z = -3.574$, $p < 0.001$), with animals exposed to
275 the elevated temperature treatment displaying more SSWD signs than those in the control
276 (Fig. 4). The effect of day was significant ($\beta = 0.037$, *s.e.* = 0.012 , $z = 3.019$, $p = 0.003$) and
277 disease signs increased over the duration of the trial. However, there was no significant
278 interaction of treatment and temperature exposure time ($\beta = -0.015$, *s.e.* = 0.017 , $z = -0.877$,
279 $p = 0.381$), showing that the rate of disease sign progression did not differ between
280 treatments. As this interaction was non-significant, subsequent models used an additive
281 effect (Table 3).

282 Further, model fit was improved significantly ($p < 0.001$) when size was incorporated into the
283 model as a fixed effect (Table 3). The resulting effects of temperature treatment was
284 significant, as was temperature exposure time. A significant effect of sea star radius was

285 found (Table 3), with greater sea star size associated with increased expression of key SSWD
286 signs.

287 3.3. Disease category

288 Sea stars in the elevated temperature treatment showed higher category disease signs than
289 in the control treatment, and disease category increased over the course of the experiment
290 (Fig. 5). However, the effect of sea star radius was not significant (Table 4), therefore
291 differing from the analysis of sum disease signs score (Table 3).

292 3.4. Survival

293 In total, four animals died during the experiment (Fig. 6). All these individuals were in the
294 elevated temperature treatment and occurred between Day 6 and Day 8. All displayed signs
295 consistent with wasting prior to death including loss of body turgor, bloating in the arms and
296 the appearance of white lesions. In three of these animals, multiple arm autotomies were
297 recorded prior to death, while one displayed disintegration of the epidermis at the joining
298 between one arm and the central disc. Viscera was clearly visible protruding through the
299 epidermis, and death occurred within a day of first observation of these signs.

300 No mortalities were recorded for the control treatment, thus the Cox Proportional Hazards
301 model resulted in a degenerate estimate. In this case, a log-rank test was applied across the
302 whole distribution, resulting in a significant difference between the two treatments ($\chi^2 =$
303 4.4, $df = 1$, $p < 0.04$). Survival probability in the elevated temperature treatment after 14
304 days of exposure was 0.778 (s.d. = 0.098, 95% CI = 0.608 - 0.996).

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307 4. DISCUSSION

308 We found that exposure to immediate temperature increase is linked to increased SSWD-
309 associated signs and their severity. Our experiment shows that rapid change in temperature
310 exacerbates disease expression and survival in *A. rubens*. Gross signs typically associated
311 with SSWD were found to increase upon exposure to sustained elevated temperature.
312 Mortality only occurred in the raised temperature treatment, highlighting the detrimental
313 effects of rapid temperature increase. Further, size was found to be a significant factor in
314 explaining disease expression in our sea stars, with a relationship between increased disease
315 signs and larger individuals. We discuss these results in light to current assumptions on
316 SSWD progression and aetiology.

317 4.1. Disease Severity

318 With elevated temperature influencing SSWD sign expression in *A. rubens*, our work
319 supports the idea that there is a link between SSWD signs and environmental stressors. The
320 first SSWD signs visually occurred after 24 hours, and sea stars in the elevated temperature
321 treatment maintained a higher disease sign score for the duration of the experiment (Fig. 4).
322 Prolonged exposure also acted to increase total disease sign score. Although temperature
323 treatments used were within the range reported at the collection site (AFBI 2015), the
324 abrupt and sustained exposure to elevated temperature may not be reflective of field
325 conditions. Therefore, conclusions of wider ecological consequences must be treated with
326 caution. It is apparent, however, that rapid and sustained exposure to elevated temperature
327 did induce signs consistent with wasting, therefore supporting the link between
328 environmental stressors and the condition.

329 Elevated temperatures result in high cellular oxygen demand due to increased metabolic
330 activity (Rühmkorff et al. 2023). This, combined with lower solubility of oxygen in warmer
331 seawater, leads to reduced concentrations of oxygen in the coelomic fluid and tissues,
332 followed by acute stress, tissue damage and mortality (Rühmkorff et al. 2023, Wahltinez et
333 al. 2023). Elevated temperatures endured for longer durations may enhance this effect.

334 Furthermore, metabolic stress may limit immune response to opportunistic pathogens
335 (Harvell et al. 2019, Hewson 2021). Coelomocyte count in *A. rubens*, an indicator of immune
336 system function in echinoderms, was found to increase with higher temperatures
337 (Wahltinez et al. 2023), and may indicate that an inflammatory response was occurring. A
338 similar effect was reported for *P. ochraceus* with active wasting signs (Work et al. 2021),
339 showing an energetic cost.

340 Echinoderms have mutable collagenous tissue (MCT) which allows passive changes to the
341 mechanical properties of the body structure (Wilkie 2005). MCT is under direct neural
342 control and allows rapid changes to body rigidity, while also being integral to the process of
343 arm autotomy (Wilkie 2005, Work et al. 2021). Inflammation of the MCT has been
344 associated with epidermal ulceration in sea stars (Núñez-Pons et al. 2018, Smith et al. 2022),
345 and disruption to control of MCT may also be responsible for general lack of body turgor,
346 the observations of bloating/pinching of arms, and abnormal body posture (Work et al.
347 2021).

348 Inflammation markers at higher temperatures may relate to disruption of MCT (Wahltinez et
349 al. 2023), leading to SSWD signs that were observed in *A. rubens*. While questions remain as
350 to the mechanism behind this inflammatory response, the process appears temperature

351 sensitive. These results also mirror experimental temperature elevation on in *A. johnstoni*
352 (Staehli et al. 2009), and *P. ochraceus* (Bates et al. 2009).

353 The addition of size into the model for total disease signs improved the fit, with larger
354 animals trending towards enhanced disease sign expression (Table 3). Elevated temperature
355 is related to greater oxygen stress placed on animals with smaller surface-to-volume ratios
356 (Staehli et al. 2009, Eisenlord et al. 2016). Smaller size may act as a refuge, therefore, during
357 periods of environmental instability. An energetically costly inflammatory response as is
358 observed under elevated temperatures (Wahlteiz et al. 2023) is better sustained by smaller
359 individuals. This may then lead to recovery of early disease signs, for example loss of turgor,
360 as the response is brought under control.

361 Dissolved oxygen (DO) was not monitored in this study as constant airflow was maintained
362 in all experimental units to maximise oxygen saturation. However, fine-scale hypoxic
363 conditions may be experienced by some sea stars due to their rugose body structure
364 (Aquino et al. 2021, Hewson 2021).

365 Microorganisms inhabiting the epidermal tissue may drive hyper-localised hypoxic
366 conditions at the interface between the sea star and the water column, thus reducing DO
367 availability (Aquino et al. 2021, Hewson 2021). Localised hypoxic conditions may exacerbate
368 existing disease in animals by microbial colonisation of necrotic tissue at lesion sites
369 (Hewson 2021). The combination of temperature associated inflammation weakened
370 immune function, MCT dysfunction and progression to lesions and tissue necrosis may all be
371 further confounded by opportunistic microorganisms inducing hypoxia at the boundary
372 layer. Our observations support this mechanism of action, with greater susceptibility to
373 SSWD at higher temperatures, and effects felt more greatly by larger individuals.

374 4.2. Disease Category

375 Overall disease signs were more frequent in sea stars exposed to elevated temperature, but
376 we also found a comparable pattern when assessing disease category based on gross body
377 characteristics (Fig. 5). Animals exhibiting Category 1 disease appeared after 24 hours in the
378 elevated temperature treatment, with Category 2 first appearing 48 hours after exposure.
379 Category 3 and Category 4 disease appeared on Day 4 and 5 respectively, and first mortality
380 was observed on Day 6. In contrast, disease in the control treatment never increased above
381 Category 1, with animals showing lesions on a single arm or disc at most after five days of
382 exposure.

383 The categorical assessment diverges from disease sign score due to the lack of consideration
384 for abnormal posture i.e., bloating/ pinching; contortion of arms; loss of body turgor. This
385 difference is reflected by the fact that all animals in the control treatment were categorised
386 as healthy until Day 5, while disease sign observations began on Day 1 when all signs are
387 counted (Fig. 4-5). This discrepancy is a result of bloating and loss of turgor being observed
388 in some individuals in the control treatment. This may also explain the non-significant effect
389 of size in the categorical assessment of disease, where bloating loss of turgor are not
390 considered (Table 4).

391 As abnormal posture, including bloating, is consistently associated with SSWD in various
392 species (Hewson et al. 2019, Work et al. 2021, Wahltinez et al. 2023), it is questionable as to
393 why a categorisation of disease severity would not account for it. Signs of abnormal posture
394 do, however, appear to be recoverable, placing them clearly at the lower end of disease
395 severity. As the pathway leading to lesion formation begins with inflammation in the MCT
396 (Work et al. 2021), it is possible that early onset of SSWD will cause disruption to MCT

397 control resulting abnormal posturing of sea stars. The recoverability of varying stages of
398 disease would therefore be worthwhile investigating, especially given the conflicting results
399 regarding sea star size.

400 In contrast, asteroids appear to present a limited suite of disease signs, and these may not
401 be consistent with a single aetiology (Hewson 2021). It is therefore possible that a
402 proportion of disease signs observed are driven by other factors. Moving animals from a
403 flow-through system to a closed system with daily water changes may have induced a stress
404 response unrelated to temperature. This may account for observations of disease signs in
405 the control treatment (Fig. 2), and that these signs are indicative of stress associated with
406 handling and water quality deteriorations over the intermediary 24 hours between water
407 changes.

408 Limited observations of lesion formation did occur in the control treatment, and may
409 suggest that coelomocyte infiltration and inflammation can progress at temperatures often
410 experienced by free-ranging *A. rubens* (Rühmkorff et al. 2023). Uncertainty remains
411 regarding the precise aetiology of SSWD. Although disease signs and progression were
412 elevated in the high temperature treatment, their presence in the control treatment
413 suggests additional stressors. Reduced immune system functioning derived from alternative
414 factors could be the cause of MCT inflammation, and further investigation is needed to
415 disentangle competing environmental stressors, as well as from a potential pathogenic
416 aetiology.

417 4.3. Survival

418 Exposure to elevated temperature caused greater than 20% mortality in the sea star *A.*
419 *rubens* (Fig. 6). Although a deeper analysis was limited by the small number of cases (4) and

420 no mortality in the control treatment, deaths were all associated with SSWD signs on
421 previous days' recordings.

422 Under simulated heatwave scenarios with peaks of 26°C, Rühmkorff et al. (2023) reported
423 100% mortality of *A. rubens*. Similarly, Staehli et al. (2009) reported a significant lethal effect
424 of elevated temperature in the Mediterranean burrowing sea star (*Astropecten johnstoni*),
425 with sea stars developing wasting-like signs prior to death. In the latter study, size was also
426 found to have a significant effect on survival, with larger animals suffering greater mortality
427 rates.

428 In the Pacific, elevated temperature was linked to increased mortality rates in *P. ochraceus*
429 (Eisenlord et al. 2016). Laboratory studies conducted by Eisenlord et al. (2016) were paired
430 with field observations, and showed reduced abundance of larger individuals following a
431 2014 outbreak of SSWD. As our mortality rates were comparatively low, we were unable to
432 perform an analysis of the effect of size on survival in *A. rubens*. However, due to the
433 combined effect of increased metabolic activity and reduced oxygen solubility at higher
434 temperatures, smaller animals may have an advantage under warming scenarios due to
435 higher surface-to-volume ratio for diffusion (Rühmkorff et al. 2023).

436 Further, it is notable that first mortality in Rühmkorff et al. (2023) occurred on day 8 after
437 exposure to 26°C, whereas at 18°C in the present study, first mortality occurred at day 6. It
438 is possible that sea star size could explain this difference. In Rühmkorff et al. (2023) mean
439 arm length was 5.5 ± 0.3 cm; smaller than the 8.9 ± 1.2 cm arm length of those inducted into
440 this experiment. As discussed, current ideas regarding SSWD progression suggest a
441 connection between larger individuals and enhanced susceptibility (Staehli et al. 2009,
442 Eisenlord et al. 2016). However, while the present study found support for size influencing

443 SSWD sign expression, further targeted research is needed to demonstrate the effect of size
444 on overall survival and mortality.

445 4.4. Ecological Implications

446 In the face of global climate change, marine disease outbreaks are thought to be becoming
447 more prevalent (Harvell et al. 2002, Eisenlord et al. 2016). Our results support a link
448 environmental stressors and SSWD-associated signs and mortality in *A. rubens*. As with
449 many sea stars, this abundant North Atlantic species acts as a keystone predator, with a
450 controlling effect on the region's bivalve molluscs (Hancock 1955, Saier 2001, Gallagher et
451 al. 2008). The possibility of thermal association of SSWD in *A. rubens* is concerning given
452 projected climate change scenarios. The increasing frequency of extreme climatic events,
453 such as marine heatwaves, is predicted to have widespread and significant impacts on
454 marine ecosystems (Oliver et al. 2019, Rühmkorff et al. 2023). Through predation on
455 bivalves, sea stars increase the possibility for settlement and attachment of other
456 organisms. Declines in *A. rubens* populations would remove this important aspect of benthic
457 ecology and encourage overgrowth of mussels in naturally biodiverse habitat. Further,
458 SSWD has been observed in the common sunstar, *Crossaster papposus*, itself a reported
459 predator of *A. rubens*, adding greater uncertainty to the potential consequences of SSWD
460 outbreaks in North Atlantic sea stars (Smith et al. 2022).

461 4.5. Limitations

462 While we found that signs associated with SSWD in *A. rubens* are temperature-sensitive, the
463 underlying mechanism remains elusive, and a pathogenic aetiology is not ruled out.
464 Regarding hypoxia, enrichment events may themselves be associated with thermal fluxes,
465 and thus disentangling different environmental stressors as a potential trigger for SSWD

466 onset may be challenging (Aquino et al. 2021, Hewson 2021). Further, changing thermal
467 regimes may push natural pathogens towards their thermal optima, thus making a single
468 pathogenic trigger challenging to disentangle from environmental stressors (Eisenlord et al.
469 2016, Hewson et al. 2018, Byers 2021). Furthermore, given that some signs of disease were
470 present in the control treatment, it is possible that additional drivers in disease
471 manifestation were present.

472 Additionally, the temperature increase to which the sea stars were exposed was abrupt,
473 rather than a gradual warming. However, temperature increases on the intertidal can be
474 rapid and severe (Legrand et al. 2018, Gilson et al. 2021), and *A. rubens* may experience
475 such extremes at local scales. Projected warming may further increase the magnitude and
476 spatial scale of such thermal fluxes, and further targeted research is needed to assess the
477 potential for SSWD outbreaks at the population level.

478 Finally, the number of sea stars inducted into the study was subject to logistical constraints,
479 and a greater sample size may add greater validity to the results obtained. Similarly, this
480 study lacks validation from field observation of wasting *A. rubens*. Anecdotal reports have
481 provided information about the presence of wasting individuals on the Irish Sea coast,
482 however this has not been systematically observed to date.

483 4.6. Conclusions

484 Signs of SSWD in *A. rubens* were induced when exposed to abrupt temperature increase.
485 Disease signs were more frequent at elevated temperatures and persisted for the duration
486 of exposure. Further, our work suggests an influence of body size on disease severity, which
487 aligns with current ideas regarding SSWD aetiology. Although the exact aetiology is yet to be
488 fully resolved, this work develops our understanding of the condition, and may support the

489 connection of SSWD to environmental stressors. The possible thermal association of SSWD
490 raises concerns for ecological stability and resilience in the face of global climate change.
491 Many sea stars are keystone species, with *A. rubens* having a significant controlling effect on
492 bivalve molluscs. If the possibility of disease outbreak is influenced by global climate change,
493 it could have cascading ecosystem effects detrimental to benthic diversity in the North
494 Atlantic.

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513 Mánuis Cunningham provided helpful comments.

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528 LITERATURE CITED

529 AFBI (Agri-Food and Biosciences institute) (2015) Carlingford Lough information - SMILE
530 project. <https://www.afbini.gov.uk/publications/carlingford-lough-information-smile->
531 [project](https://www.afbini.gov.uk/publications/carlingford-lough-information-smile-project) (accessed 07 August 2023)

532 Agüera A, Jansen JM, Smaal AC (2020) Blue mussel (*Mytilus edulis* L.) association with
533 conspecifics affects mussel size selection by the common seastar (*Asterias rubens* L.). J Sea
534 Res 164:101935.

535 Agüera A, Saurel C, Møller LF, Fitridge I, Petersen JK (2021) Bioenergetics of the common
536 seastar *Asterias rubens*: a keystone predator and pest for European bivalve culture. Mar Biol
537 168:1–14.

538 IPCC (Intergovernmental Panel on Climate Change) (2021) Climate Change 2021 – The
539 physical science basis: Working Group I Contribution to the Sixth Assessment Report of the
540 Intergovernmental Panel on Climate Change. Cambridge University Press, Cambridge

541 Aquino CA, Besemer RM, DeRito CM, Kocian J, Porter IR, Raimondi PT, Rede JE, Schiebelhut
542 LM, Sparks JP, Wares JP, Hewson I (2021) Evidence that microorganisms at the animal-water
543 interface drive sea star wasting disease. Front Microbiol 11:3278.

544 Bates AE, Hilton BJ, Harley CDG (2009) Effects of temperature, season and locality on
545 wasting disease in the keystone predatory sea star *Pisaster ochraceus*. Dis Aquat Org
546 86:245–251.

547 Bates D, Mächler M, Bolker B, Walker S (2015) Fitting linear mixed-effects models using
548 lme4. J Stat Softw 67:1–48.

549 Bruno JF, Selig ER, Casey KS, Page CA, Willis BL, Harvell CD, Sweatman H, Melendy AM
550 (2007) Thermal stress and coral cover as drivers of coral disease outbreaks. PLOS Biol
551 5:e124.

552 Bucci C, Francoeur M, McGreal J, Smolowitz R, Zazueta-Novoa V, Wessel GM, Gomez-Chiarri
553 M (2017) Sea star wasting disease in *Asterias forbesi* along the Atlantic coast of North
554 America. PLOS ONE 12:e0188523.

555 Burge CA, Mark Eakin C, Friedman CS, Froelich B, Hershberger PK, Hofmann EE, Petes LE,
556 Prager KC, Weil E, Willis BL, Ford SE, Harvell CD (2014) Climate change influences on marine
557 infectious diseases: implications for management and society. Annu Rev Mar Sci 6:249–277.

558 Byers JE (2020) Marine parasites and disease in the era of global climate change. Annu Rev
559 Mar Sci 13:397-420

560 Calderwood J, O'Connor NE, Roberts D (2016) Efficiency of starfish mopping in reducing
561 predation on cultivated benthic mussels (*Mytilus edulis* Linnaeus). Aquaculture 452:88–96.

562 Dungan ML, Miller TE, Thomson DA (1982) Catastrophic decline of a top carnivore in the
563 Gulf of California rocky intertidal zone. Science 216:989–991.

564 Eckert GL, Engle JM, Kushner DJ (2000) Sea star disease and population declines at the
565 Channel Islands. In: Browne DR, Mitchell KL, Chaney HW (eds). Proceedings of the fifth
566 California Islands Symposium. Minerals Management Service, Camarillo, CA. p 435–441.

567 Eisenlord ME, Groner ML, Yoshioka RM, Elliott J, Maynard J, Fradkin S, Turner M, Pyne K,
568 Rivlin N, van Hooidek R, Harvell CD (2016) Ochre star mortality during the 2014 wasting
569 disease epizootic: role of population size structure and temperature. Philos Trans R Soc B
570 371:20150212.

571 Gallagher T, Richardson CA, Seed R, Jones T (2008) The seasonal movement and abundance
572 of the starfish, *Asterias rubens* in relation to mussel farming practice: A case study from the
573 Menai Strait, UK. J Shellfish Res 27:1209–1215.

574 Gilson AR, Coughlan NE, Dick JTA, Kregting L (2021) Marine heat waves differentially affect
575 functioning of native (*Ostrea edulis*) and invasive (*Crassostrea [Magallana] gigas*) oysters in
576 tidal pools. Mar Environ Res 172:105497.

577 Hancock DA (1955) The feeding behaviour of starfish on Essex oyster beds. J Mar Biol Assoc
578 UK 34:313–331.

579 Harvell CD, Mitchell CE, Ward JR, Altizer S, Dobson AP, Ostfeld RS, Samuel MD (2002)
580 Climate warming and disease risks for terrestrial and marine biota. Science 296:2158–2162.

581 Harvell CD, Montecino-Latorre D, Caldwell JM, Burt JM, Bosley K, Keller A, Heron SF,
582 Salomon AK, Lee L, Pontier O, Pattengill-Semmens C, Gaydos JK (2019) Disease epidemic and
583 a marine heat wave are associated with the continental-scale collapse of a pivotal predator
584 (*Pycnopodia helianthoides*). Sci Adv 5:eaau7042.

585 Hewson I (2021) Microbial respiration in the asteroid diffusive boundary layer influenced
586 sea star wasting disease during the 2013-2014 northeast Pacific Ocean mass mortality
587 event. Mar Ecol Prog Ser 668:231–237.

588 Hewson I, Aquino CA, Derito CM (2020) Virome variation during sea star wasting disease
589 progression in *Pisaster ochraceus* (Asteroidea, echinodermata). Viruses 12:1332

590 Hewson I, Button JB, Gudenkauf BM, Miner B, Newton AL, Gaydos JK, Wynne J, Groves CL,
591 Hendler G, Murray M, Fradkin S, Breitbart M, Fahsbender E, Lafferty KD, Kilpatrick AM,
592 Miner CM, Raimondi P, Lahner L, Friedman CS, Daniels S, Haulena M, Marliave J, Burge CA,

593 Eisenlord ME, Harvell CD (2014) Densovirus associated with sea-star wasting disease and
594 mass mortality. *Proc Natl Acad Sci USA* 111:17278–17283.

595 Hewson I, Bistolas KSI, Quijano Cardé EM, Button JB, Foster PJ, Flanzenbaum JM, Kocian J,
596 Lewis CK (2018) Investigating the complex association between viral ecology, environment,
597 and northeast Pacific Sea Star Wasting. *Front Mar Sci* 5:77.

598 Hewson I, Sullivan B, Jackson EW, Xu Q, Long H, Lin C, Quijano Cardé EM, Seymour J, Siboni
599 N, Jones MRL, Sewell MA (2019) Perspective: Something old, something new? Review of
600 wasting and other mortality in Asteroidea (Echinodermata). *Front Mar Sci* 6:406.

601 Jackson EW, Pepe-Ranney C, Johnson MR, Distel DL, Hewson I (2020a) A highly prevalent
602 and pervasive densovirus discovered among sea stars from the North American Atlantic
603 coast. *Appl Environ Microbiol* 86:e02723-19

604 Jackson EW, Wilhelm RC, Johnson MR, Lutz HL, Danforth I, Gaydos JK, Hart MW, Hewson I,
605 Pfeiffer JK (2020b) Diversity of sea star-associated densoviruses and transcribed
606 endogenous viral elements of densovirus origin. *J Virol* 95:e01594-20

607 Legrand E, Riera P, Pouliquen L, Bohner O, Cariou T, Martin S (2018) Ecological
608 characterization of intertidal rockpools: seasonal and diurnal monitoring of physico-
609 chemical parameters. *Regional Studies in Marine Science* 17:1–10.

610 Levitan DR, Best RM, Edmunds PJ (2023) Sea urchin mass mortalities 40 y apart further
611 threaten Caribbean coral reefs. *Proc Natl Acad Sci USA* 120:e2218901120.

612 MARINe (Multi-agency Rocky Intertidal Network) (2018). Sea Star Wasting Syndrome |
613 MARINe. <https://marine.ucsc.edu/data-products/sea-star-wasting/index.html#id-guides>
614 (accessed 15 November 2022).

615 Mead AD (1898) Twenty-eighth annual report of the commissioners of inland fisheries made
616 to the general assembly at its January session, 1898. In: Southwick JMK, Root HT, Willard
617 CW, Morton WMP, Roberts AD, Bumpus HC (eds). Freeman and Sons, Printers to the State,
618 p112.

619 Menge BA (1979) Coexistence between the seastars *Asterias vulgaris* and *A. forbesi* in a
620 heterogeneous environment: a non-equilibrium explanation. *Oecologia* 41:245–272.

621 Menge BA, Cerny-Chipman EB, Johnson A, Sullivan J, Gravem S, Chan F (2016) Sea star
622 wasting disease in the keystone predator *Pisaster ochraceus* in Oregon: insights into
623 differential population impacts, recovery, predation rate, and temperature effects from
624 long-term research. *PLOS ONE* 11:e0153994.

625 Miner CM, Burnaford JL, Ambrose RF, Antrim L, Bohlmann H, Blanchette CA, Engle JM,
626 Fradkin SC, Gaddam R, Harley CDG, Miner BG, Murray SN, Smith JR, Whitaker SG, Raimondi
627 PT (2018) Large-scale impacts of sea star wasting disease (SSWD) on intertidal sea stars and
628 implications for recovery. *PLOS ONE* 13:e0192870.

629 Morón Lugo SC, Baumeister M, Nour OM, Wolf F, Stumpp M, Pansch C (2020) Warming and
630 temperature variability determine the performance of two invertebrate predators. *Sci Rep*
631 10:6780

632 Núñez-Pons L, Work TM, Angulo-Preckler C, Moles J, Avila C (2018) Exploring the pathology
633 of an epidermal disease affecting a circum-Antarctic sea star. *Sci Rep* 8:11353.

634 Oliver ECJ, Burrows MT, Donat MG, Sen Gupta A, Alexander LV, Perkins-Kirkpatrick SE,
635 Benthuisen JA, Hobday AJ, Holbrook NJ, Moore PJ, Thomsen MS, Wernberg T, Smale DA

636 (2019) Projected marine heatwaves in the 21st century and the potential for ecological
637 impact. *Front Mar Sci* 6:734

638 Paine RT (1966) Food web complexity and species diversity. *Am Nat* 100:65–75.

639 Prestedge GK (1998) The distribution and biology of *Patiriella vivipara* (Echinodermata:
640 Asteroidea: Asterinidae) a sea star endemic to Southeast Tasmania. *Rec Aust Mus* 50:161–
641 170.

642 R Core Team (2022) R: A language and environment for statistical
643 computing. R Foundation for Statistical Computing, Vienna, Austria.

644 Rühmkorff S, Wolf F, Vajedsamiei J, Barboza FR, Hiebenthal C, Pansch C (2023) Marine
645 heatwaves and upwelling shape stress responses in a keystone predator. *Proc R Soc B*
646 290:20222262.

647 Saier B (2001) Direct and indirect effects of seastars *Asterias rubens* on mussel beds (*Mytilus*
648 *edulis*) in the Wadden Sea. *J Sea Res* 46:29–42.

649 SeaTemperatures (2023). Carlingford Sea Temperature in October - SeaTemperatures.net.
650 <http://seatemperatures.net/europe/ireland/carlingford-october-temperature/> (accessed 07
651 August 2023).

652 Smith, L.C., Byrne, M., Gedan, K.B., Lipscomb, D.L., Majeske, A.J. and Tafesh-Edwards, G.,
653 2022. Echinoderm diseases and pathologies. In: Rowley AF, Coates CJ, Whitten MM (eds)
654 Invertebrate pathology. Oxford University Press pp. 563–588.

655 Smith S, Hewson I, Collins P (2022) The first records of sea star wasting disease in *Crossaster*
656 *papposus* in Europe. *Biol Lett* 18:20220197

657 Staehli A, Schaerer R, Hoelzle K, Ribi G (2009) Temperature induced disease in the starfish
658 *Astropecten jonstoni*. Mar Biodivers Rec 2:e78

659 Wahltinez SJ, Kroll KJ, Behringer DC, Arnold JE, Whitaker B, Newton AL, Edmiston K, Hewson
660 I, Stacy NI (2023) Common sea star (*Asterias rubens*) coelomic fluid changes in response to
661 short-term exposure to environmental stressors. Fishes 8:51.

662 Ward JR, Kim K, Harvell CD (2007) Temperature affects coral disease resistance and
663 pathogen growth. Mar Ecol Prog Ser 329:115–121.

664 Wilkie IC (2005) Mutable collagenous tissue: overview and biotechnological perspective.
665 Prog Mol Subcell Biol 39:221–250.

666 Work TM, Weatherby TM, DeRito CM, Besemer RM, Hewson I (2021) Sea star wasting
667 disease pathology in *Pisaster ochraceus* shows a basal-to-surface process affecting color
668 phenotypes differently. Dis Aquat Org 145:21–33.

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678 TABLES

679 **Table 1. Disease Categories and identification characteristics for *Pisaster ochraceus* &**
680 ***Evasterias troschellii* as defined by Pacific Rocky Intertidal.org.**

Identification	Disease Category
Animal shows grossly normal appearance. Animals above tide line may have deflated appearance due to water loss, yet this is not indicative of SSWD.	0
Lesions present on one arm or the central disc.	1
Lesions present on two arms/one arm and the central disc.	2
Lesions present on most of body. One or Two missing arms.	3
Severe Tissue deterioration/ death. Three or more missing arms.	4

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689 **Table 2. The effect of elevated temperature on disease status in *Asterias rubens*. A)**
 690 Displays terms, coefficients, p values for the best fitting model. B) Displays Akaike
 691 Information Criterion (AIC) and Log Likelihood for all models testing the effect of
 692 temperature treatment on disease signs. Day was included as a random effect, and nested
 693 within sea star ID, allowing individual slopes to vary according to individual through time.
 694 Significant results ($p < 0.05$) highlighted in bold.

(a) -Term	Estimate	s.e.	z-value	p-value
Intercept	8.718	2.798	3.116	0.002
Day	-0.238	0.068	-3.483	< 0.001
Sea Star Radius	-0.116	0.032	5.185	< 0.001
Treatment (Control)	4.891	0.943	-3.668	< 0.001

(b) -Model	AIC	Log. Likelihood
Day + Sea Star Radius + Treatment	360.74	-173.37
Day * Treatment	367.61	-176.81
Day + Treatment	371.83	-179.91

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700 **Table 3. The effect of elevated temperature SSWD disease signs score in *Asterias rubens*.**

701 A) Displays terms, coefficients, p values for the best fitting model. B) Displays Akaike
702 Information Criterion (AIC) and Log Likelihood for all models testing the effect of
703 temperature treatment on disease signs. Day was included as a random effect, and nested
704 within sea star ID, allowing individual slopes to vary according to individual through time.
705 Significant results ($p < 0.05$) highlighted in bold.

(a) -Term	Estimate	s.e.	z-value	p-value
Intercept	-5.747	1.702	-3.376	< 0.001
Day	0.030	0.008	3.506	< 0.001
Log (Sea Star Radius)	1.545	0.380	4.061	< 0.001
Treatment (Control)	-0.854	0.115	-7.438	< 0.001

(b) -Model	AIC	Log. Likelihood
Day + Log (Sea Star Radius) + Treatment	1050.5	-518.27
Day + Treatment	1062.0	-525.03
Day * Treatment	1063.3	-524.65

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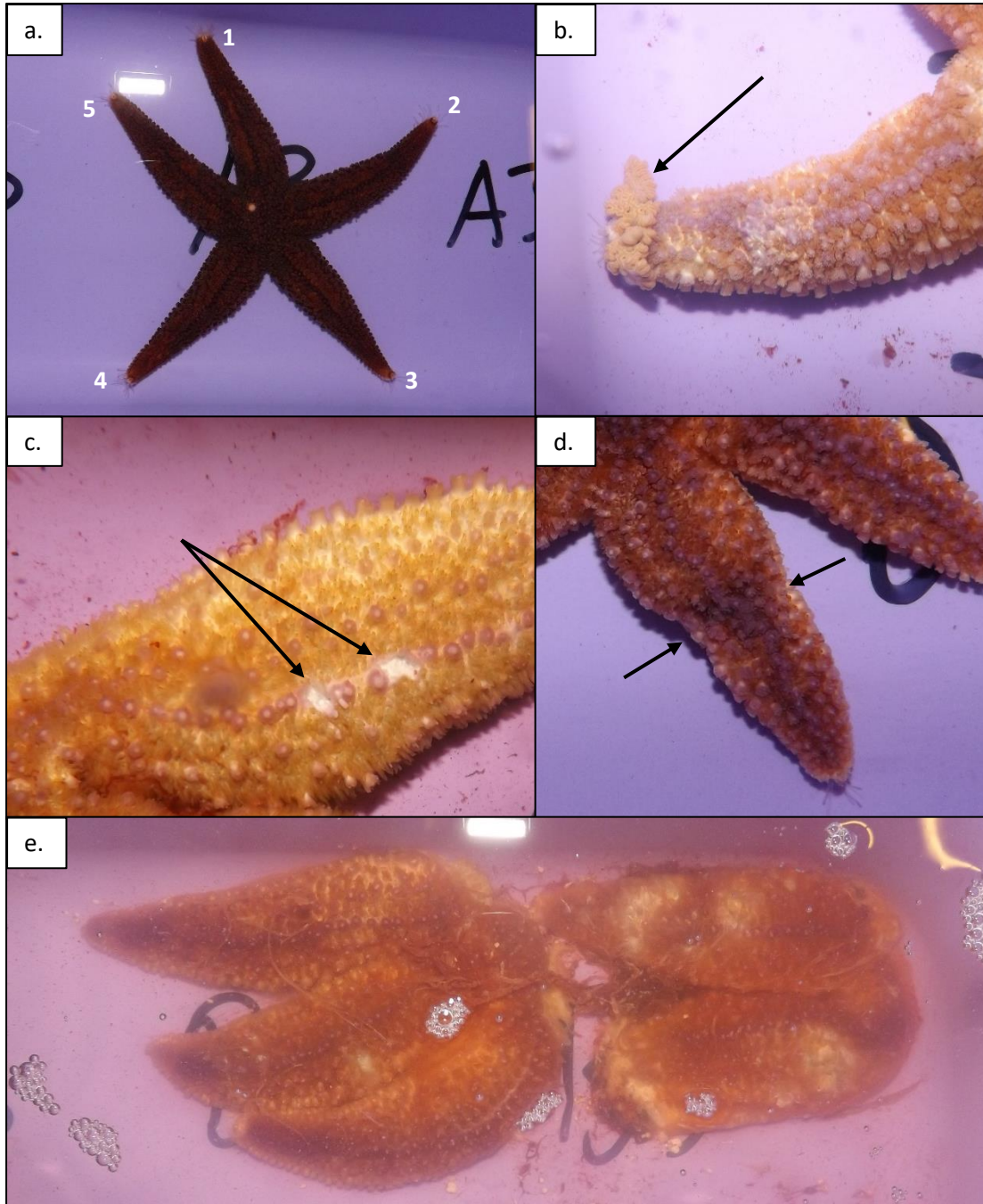
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710 **TABLE 4. The effect of elevated temperature on SSWD category in *Asterias rubens*.** Table
 711 displays terms, coefficients, p values for the best fitting model. Day was included as a
 712 random effect, and nested within sea star ID, allowing individual slopes to vary according to
 713 individual through time. Significant results ($p < 0.05$) highlighted in bold.

(a) -Term	Estimate	s.e.	z-value	p-value
Intercept	-25.152	14.329	-1.755	0.079
Day	0.126	0.043	2.928	0.003
Log (Sea Star Radius)	5.131	3.190	1.608	0.108
Treatment (Control)	-3.862	1.018	-3.794	< 0.001

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727 **Figure 1. Specimens of the common starfish *Asterias rubens* showing key gross signs**
728 **associated with sea star wasting disease.** Nominally healthy/asymptomatic individual
729 displaying no gross abnormalities; no epidermal lesions or ulceration, even arm tapering.

730 Arms numbered for tracking of disease signs (a) viscera everted from arm tip (b) epidermal
731 lesion on arm median presenting as bright white patches on epidermis (c) arm displaying
732 bloated/pinched appearance leading to abnormal taper (d) deceased individual displaying
733 multiple arms autotomised from central disc, with associated gross disintegration of
734 epidermal tissue and no movement or adhesion of tube feet (e).

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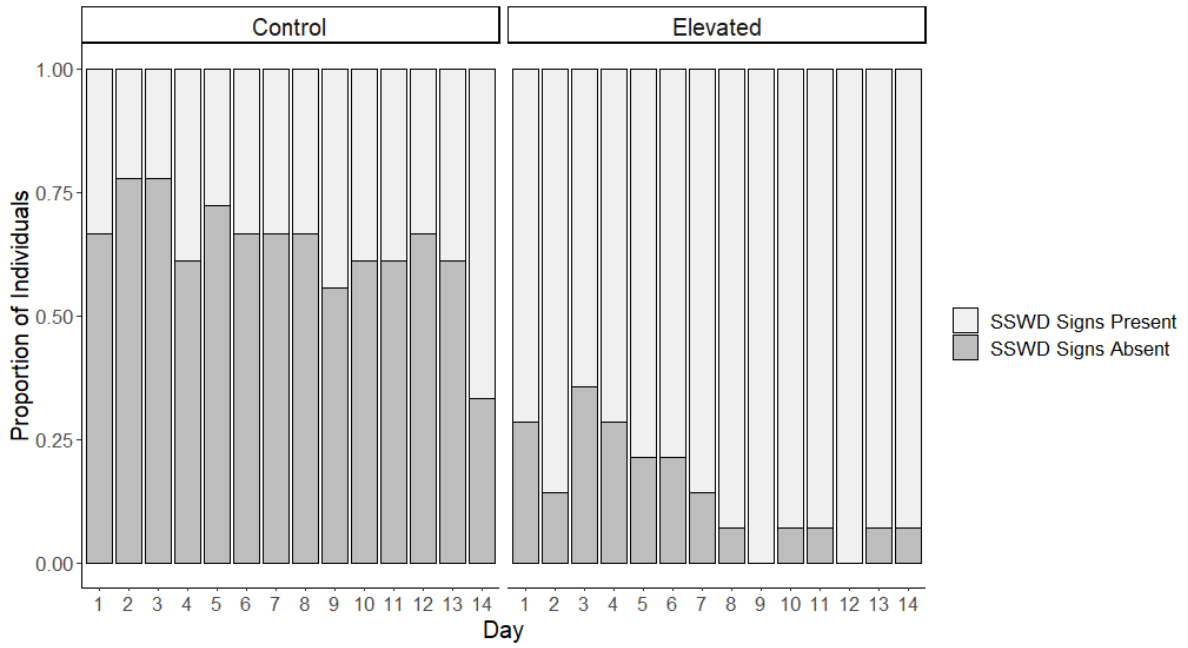
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751 **Figure 2. Stacked bar plot displaying the proportion of animals that were grossly normal**
 752 **for each temperature treatment.** Plot compares grossly normal animals (dark) against
 753 animals expressing any sign of disease (light), with proportions displayed for each day of
 754 temperature treatment exposure. Left panel shows control treatment; right panel shows
 755 elevated treatment.

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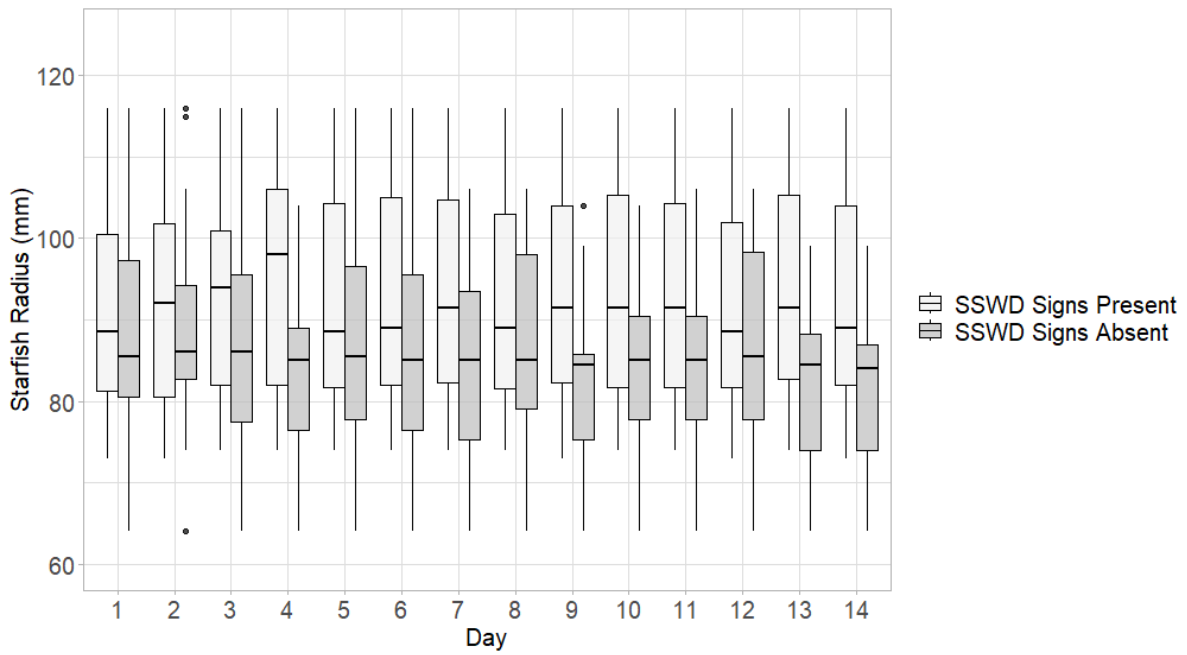
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764 **Figure 3. Boxplot of sea star radius and appearance.** Grossly normal animals (dark)

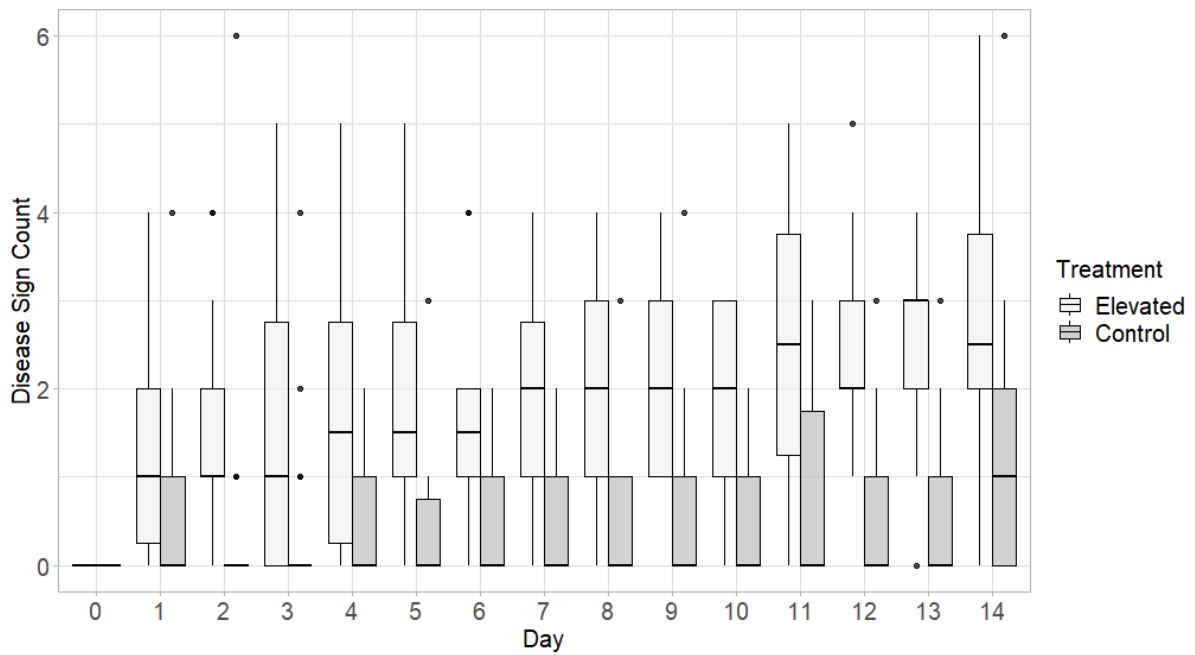
765 compared to animals with any disease sign present (light). Boxplots for each day of

766 temperature treatment exposure display median and bind 25th and 75th percentiles.

767 Whiskers mark upper and lower data bounds, points denote outliers.

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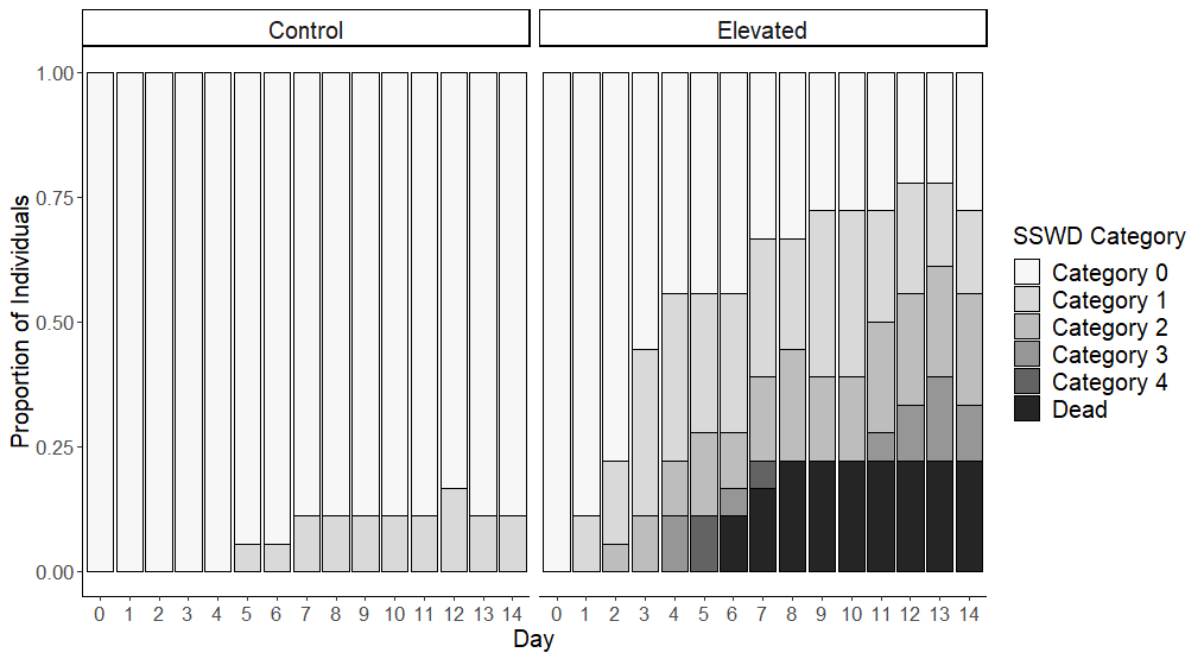
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771 **Figure 4. The effect of increasing temperature on disease signs of SSWD.** Boxplot of total
772 number of observed signs for each sea star exposed to differing temperature treatments
773 over the 14-day exposure period, including day 0. Boxplots describe median values and bind
774 25th and 75th percentiles. Whiskers mark upper and lower data bounds, points denote
775 outliers. Animals that died were removed from data set ($n = 4$). A target of 12°C represented
776 the control treatment ($n = 18$), while a target of 18°C was used for the elevated temperature
777 treatment ($n = 14$).

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782 **Figure 5. Stacked bar plot showing the proportion of animals assigned to discrete disease**

783 **categories when exposed to different temperature treatments over 14 days. Left panel**

784 **displays results for the control treatment ($n = 18$); right panel shows the elevated**

785 **temperature treatment ($n = 18$).**

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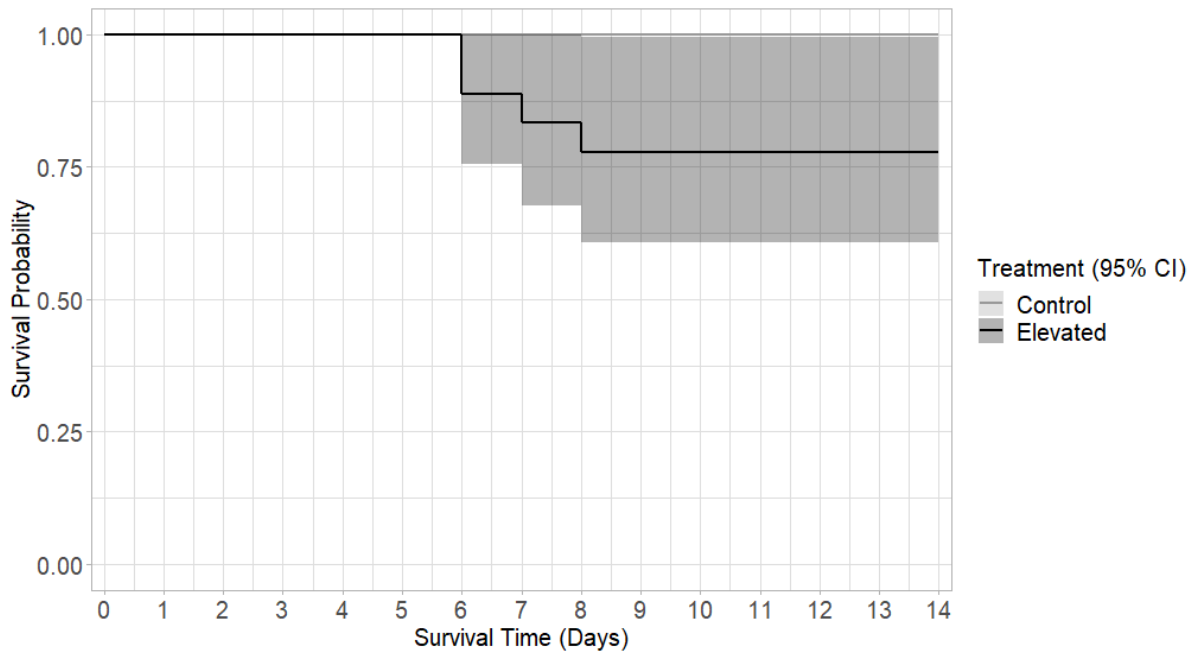
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794 **Figure 6. Kaplan-Meier Survival Plot of *Asterias rubens* subjected to different temperature**

795 **treatments over a two week period. A target of 12°C represented the control treatment (n**

796 **=18), while a target of 18°C was used for the elevated temperature treatment (n =18).**

797 Shaded area represents 95% CI.

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