

1 **Population structure and reproduction of the alvinocaridid shrimp *Rimicaris exoculata* on**
2 **the Mid-Atlantic Ridge: variations between habitats and vent fields**

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21 **Abstract**

22 *Rimicaris exoculata* is a dominant species of deep Mid-Atlantic Ridge (MAR) vent fields and
23 inhabits areas close to vent emissions at depths below 2000 m. Its high abundance and strong
24 genetic connectivity along the MAR point at a remarkable ability to produce dispersing larval
25 stages. However, the reproduction of this species long remained enigmatic because brooding
26 females were rarely observed. Here, we describe the population structure and reproduction of *R.*
27 *exoculata* at the Snake Pit and TAG vent fields (3600m depth) for the months of January-
28 February. We observed major differences in population structure between habitats within a single
29 vent field: females widely dominate the large swarms near active venting while inactive
30 peripheries are inhabited by large males. Low temperature diffusion zones are mainly colonized
31 by small juveniles of *R. chacei* instead of *R. exoculata*. Size structure of populations from dense
32 active areas is polymodal at both fields, suggesting discontinuous recruitment. Male and female
33 sizes did not vary across habitats and vent fields, with sexually mature female being slightly larger
34 than males. In contrast to previous studies, hundreds of ovigerous females were observed at both
35 vent fields, suggesting seasonal reproduction. Proportion of ovigerous females among sexually
36 mature females were similar between vent fields (36.7 %). However, reproductive output was
37 lower at TAG, where ovigerous females had smaller size-specific fecundity and egg size, and
38 more aborted broods. Broods were colonized by the dirivultid copepod *Stygiopontius pectinatus*
39 at both vent fields, apparently without deleterious effect on egg development. In the light of the
40 observed variability in *R. exoculata* population structure, we propose a hypothetical scenario
41 depicting its mating system and brooding behavior, and discuss more generally intraspecific
42 interactions during its benthic life stages.

43 **Key words:** Life cycle, population structure, reproduction, *Rimicaris exoculata*, habitat
44 variability, hydrothermal vents

45 **1. Introduction**

46 The fragmentary and ephemeral nature of deep-sea hydrothermal vent ecosystems is
47 challenging for animal populations connectivity and resilience. Visually dominant species are
48 usually endemic to these ecosystems, and large scale distribution and connectivity between
49 populations were observed for some of them (Thaler et al. 2011, Teixeira et al. 2012, Beedessee
50 et al. 2013). Advances in understanding mechanisms and processes related to reproduction,
51 dispersal, recruitment and structure of vent populations are still hampered by the difficulties to
52 upscale our observations temporally and/or spatially due to the cost and technical challenges of
53 deep-sea studies. Remaining gaps in our understanding of vent species life histories may hide
54 complex population structures that have not yet been acknowledged, even in some well-studied
55 iconic species such as vent shrimps. Indeed, a recent study in another vent crustacean, the
56 anomuran crab *Kiwa tyleri*, revealed complex life stages distribution affecting local population
57 structures and reflecting life history of the species (Marsh et al. 2015).

58 The alvinocarid shrimp *Rimicaris exoculata* is a visually dominant species at hydrothermal
59 vents of the Mid-Atlantic Ridge (MAR), especially in fields located at depths below 2000 m. This
60 species lives close to the vent emission in dense aggregations of thousands of individuals per
61 square meter (Desbruyères et al. 2000, Copley et al. 2007, Gebruk et al. 2010) and many aspects
62 of its biology have been studied since the discovery of MAR hydrothermal vents in 1985 (Zbinden
63 & Cambon-Bonavita 2020). The relationship of this shrimp with the dense and diverse symbiotic
64 microbial communities hosted in its cephalothoracic cavity and gut have been extensively studied,
65 demonstrating a clear trophic dependency of the shrimp on its symbionts, and pointing at tight
66 regulation along its life cycle (Corbari et al. 2008, Ponsard et al. 2013, Jan et al. 2014, Le Bloa et
67 al. 2020). However, the information available about reproduction and population biology is still
68 scattered and sometimes contradictory. In terms of population structure, major differences in
69 shrimp densities have been found at the Broken Spur vent field depending on the levels of

70 hydrothermal activity, including the decrease of adult densities in vent areas not directly exposed
71 to vent emission (Copley et al. 1997). Habitats at the base of vent edifices were suggested to
72 serve as nurseries for juvenile recruitment due to their high densities in these areas (Komai and
73 Segonzac 2008). However, it remained unclear whether variations in adult density between
74 habitats also involved variations in population structures. Samples collected at various MAR vent
75 fields by Shank et al. (1998) showed a female-biased sex ratio, but the association of sex ratio
76 with a particular habitat or temporal variation was not tested by additional sampling. In addition,
77 patches of juveniles have been reported in adult aggregations (Shank et al. 1998, Copley et al.
78 2007) but their distribution and proportions in the populations remain undetermined. Other
79 aspects of population structure, such as size structure, are based on single sample, pooled
80 samples or preliminary analyses (Gebruk et al. 1997, Vereshchaka 1997).

81 The reproduction of *R. exoculata* also has many intriguing gaps. Although oocyte size
82 frequency suggests a lack of seasonality in the reproduction (Ramirez-Llodra et al. 2000, Copley
83 et al. 2007), very few brooding specimens have been collected since the species description
84 (Williams & Rona 1986, Ramirez-Llodra et al. 2000, Gebruk 2010, Guri et al. 2012). The lack of
85 brooding females also contrasts with the large densities and strong genetic connectivity observed
86 in MAR vent fields (Teixeira et al. 2012), which must be supported by a large larval pool. Egg size
87 has been estimated in only two studies (Williams & Rona 1986, Ramirez-Llodra et al. 2000) and
88 this size estimation was used to infer that these shrimp should have planktotrophic larvae with
89 long planktonic duration. However, recent morphological evidence on hatching larval stages
90 lacking functional mouth structures led to the revision of the former hypothesis and proposition
91 that these larvae are lecithotrophic early in their dispersive planktonic phase (Hernandez-Avila et
92 al. 2015). In addition, the realized fecundity (988 eggs per brood) was estimated on only one
93 specimen due the lack of samples (Ramirez-Llodra et al. 2000).

94 Very few data concerning the distribution of *R. exoculata* populations at different spatial
95 scales or their interactions in vent ecosystems are available in the literature. At vents, several
96 studies have shown that variations of physical and chemical conditions of vent emission shape
97 the structure of communities dominated by bathymodiolin mussels or siboglinid tube worms at
98 various spatial and temporal scales (Sarrazin et al. 1997, 2015, Cuvelier et al. 2009, 2014).
99 Variations in community structure are observed between edifices at the field-scale (Desbruyères
100 et al. 2000, 2001, Sarrazin et al. 2020) as well as between habitats within the same edifice
101 (Cuvelier et al. 2009, Sarrazin et al. 2015) highlighting the high level of complexity of these
102 ecosystems.

103 In addition to variations at community level, vent fauna also show variations in their
104 population structures. These variations are linked to differences in habitats occupied by the
105 different life stages for motile organisms (Shank et al. 1998, Marsh et al. 2015), changes in
106 population structure between vents (Nye et al. 2013) or habitats (Copley & Young 2006, Marsh et
107 al. 2015) and various scales of temporal variations (Copley et al. 1997, 1999, 2007, Gebruk et al.
108 2010, Cuvelier et al. 2011). Although less studied, there is evidence that populations of dominant
109 species exhibit spatial and temporal variations at small scales. For instance, Copley et al. (1999)
110 proposed that short-term changes in *R. exoculata* population density close to vent emission could
111 be associated with tidal variation, as observed in other mobile vent species (Lelièvre et al. 2017).
112 Variations in population structure most likely reflect the complex life cycles of vent species with
113 different reproductive strategies, variations in physiological tolerance and resource use by the
114 different developmental stages, and their interactions with environmental variations.

115 As one of the dominant species at MAR vent fields below 2000 m depth, *R. exoculata*
116 plays a major role at ecosystem level, and its population biology could have important implications
117 for the resilience, structure and biomass of these vent communities (Desbruyères et al. 2000,
118 2001). In January-February 2014, large numbers of ovigerous females were observed at two vent

119 fields of the MAR, TAG and Snake Pit, during the BICOSE cruise. In addition, striking variations
120 in the densities of adult and juvenile shrimps locally led us to hypothesize local variations in
121 population structure. In this study, we tested such hypothesis and analyzed both population
122 structure and reproductive features of ovigerous *R. exoculata* females in a series of samples
123 collected in visually distinct shrimp assemblages. We untangled different levels of spatial variation
124 in reproductive parameters and population structure and used this information to propose a
125 scenario depicting interactions of shrimps with their conspecifics and their environment that would
126 explain the observed distribution patterns, and provide clues on some aspects of their
127 reproductive behavior and life cycle.

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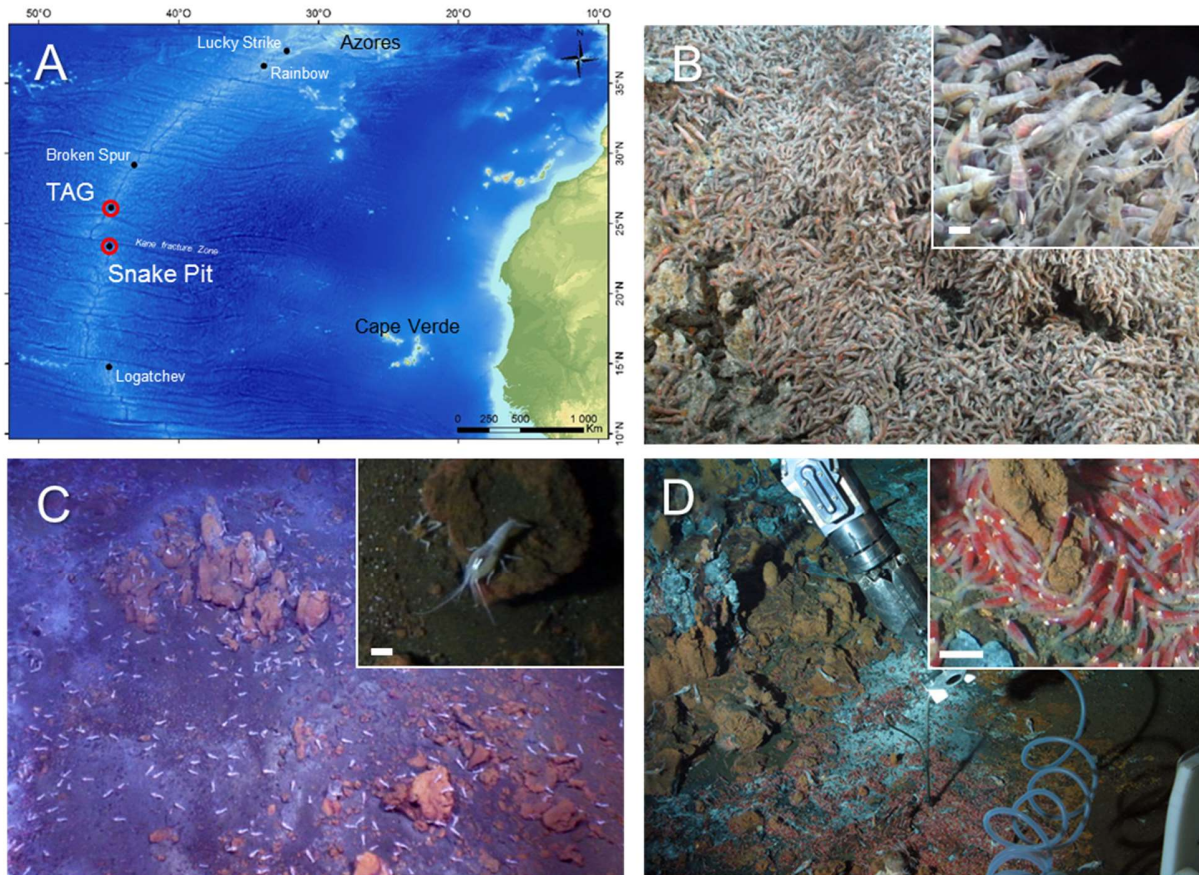
129 **2. Material and Methods**

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131 **2.1. Sampling**

132 *Rimicaris exoculata* were collected at the Snake Pit (SP, 23°22.1'N 44°57.1'W, 3470 m
133 depth) and TAG (26°08.2'N, 44°49.5'W, 3620 m depth) vent fields on the MAR (Fig. 1a) during
134 the BICOSE cruise (DOI: 10.17600/14000100) from January 10th to February 11th, 2014. The two
135 vent fields are approximately 310 km away, and separated by the Kane Fracture Zone. At Snake
136 Pit, six samples were collected in dense shrimp aggregations on the walls of active chimneys of
137 the Beehive site, close to fluid emissions (therein Active Emission Habitat/AEH) (Table 1, Suppl.
138 Fig. 1). At TAG, three samples were collected in the AEH of Active Mound (Fig. 1b, Suppl. Fig. 1)
139 and two samples were collected in dense aggregations of small alvinocaridid juveniles settled in
140 flat areas of diffuse flow (Diffuse Emission Habitat/DEH) herein termed “nurseries” (Fig. 1d, Suppl.
141 Fig. 2). Additionally, three other samples were collected at the base of the TAG mound, where no
142 active emission was visible (Inactive Emission Habitat/IEH) and with adult shrimps scattered over

143 large areas (Fig. 1c, Suppl. Fig. 2). These different types of habitats occurred within meters or
144 tens of meters from each other.



145
146 Figure 1. Sampled vent fields and habitats. A) North Atlantic regional map with TAG and Snake
147 Pit vent fields on the Mid Atlantic Ridge (Amante, C. and B.W. Eakins, 2009.
148 doi:10.7289/V5C8276M). B) *R. exoculata* swarms in the active emission habitat (AEH). C)
149 Inactive emission habitat (IEH) with scattered *R. exoculata* adults (white spots). D) Diffuse
150 emission habitat (DEH), showing red aggregation of small juveniles. B-D : Pictures from the TAG
151 vent field, Ifemer/ROV Victor 6000/BICOSE2014. Scale bars in close-up views in insets: 1 cm.

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155 Table 1. Shrimp sample details, including temperature ranges measured within centimeters from
156 the faunal sampling point when available. ND: no data available.

Vent Field	Dive	Sample	Sample size (ind.)	Juveniles for genetics	Habitat	Depth (m)	T (°C)
SP	PI01-564-Aspi3	S1	143		AEH	3463	3.5-22
SP	PI01-564-Aspi4	S2	390		AEH	3463	3.5-22
SP	PI01-564-Aspi5	S3	808		AEH	3463	3.5-22
SP	PI05-568-Aspi1	S4	391		AEH	3465	10-21
SP	PI05-568-Aspi4	S5	110		AEH	3468	4-18
SP	PI05-568-Aspi6	S6	364	1	AEH	3472	ND
TAG	PI08-571-Aspi2	S7	600	4	AEH	3626	8-14
TAG	PI10-573-Aspi1	S8	207		AEH	3624	6-33
TAG	PI10-573-Aspi2	S9	161		AEH	3627	3-30
TAG	PI10-573-Aspi5	S10	18		IEH	3635	2.4-2.8
TAG	PI10-573-Aspi6	S11	38		IEH	3635	2.4-2.8
TAG	PI12-575-Nasse 2	S12	37		IEH	3634	2.4-2.8
TAG	PI08-571-Aspi1	S13	77	36	DEH	3637	2.8-5.3
TAG	PI12-575-Aspi2	S14	101	36	DEH	3637	2.8-5.3

157

158 Shrimps were collected with the suction sampler of the Remotely Operated Vehicle (ROV)
159 Victor6000. In AEH and DEH, the tip of the sampler was pointed as close as possible to the
160 individuals and maintained immobile during sampling to avoid as much as possible disturbing the
161 aggregation. The suction was activated for a few seconds in order to collect individuals from a
162 restricted area. Due to the low shrimp density in IEH, samples were obtained from larger areas
163 covering a few m². In addition, 37 specimens caught with a shrimp-trap deployed in IEH were
164 included in the analyses. In total 3 445 specimens of *R. exoculata* were examined (Table 1).

165 Temperature measurements were conducted along with shrimp collections. Records were
166 obtained either from discrete measurements with the submersible temperature probe prior to
167 sampling, or from time-series measurements with autonomous temperature probes (WHOI-MISO
168 low temp-ONSET®) deployed within the shrimp aggregation for a few days prior to sampling.

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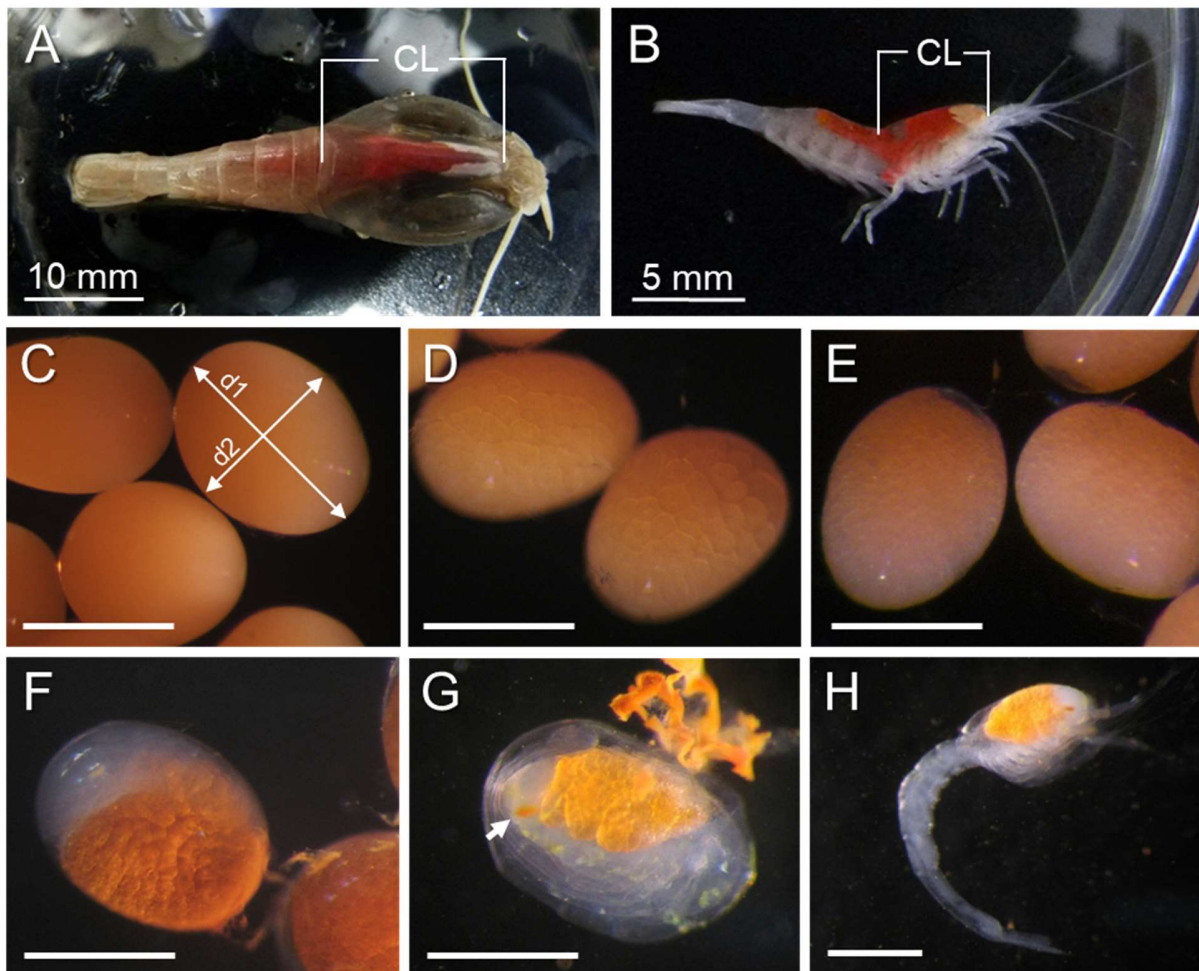
170 **2.2. Identification and measurements**

171 Specimens were initially identified and classified as juveniles, subadults or adults, in
172 accordance with Komai & Segonzac (2008). The identification of the smaller juveniles from DEH
173 was further assessed by DNA analyses because their morphology did not fit completely the
174 description of early juvenile stages of *R. exoculata* available at the time of this study (Komai &
175 Segonzac, 2008). Thirty-six juveniles from each DEH sample and 5 juveniles from AEH samples
176 were used for molecular identifications (Table 1 & Suppl. Table 1).

177 Sex was identified in adults by the occurrence of the “appendix masculina” on the second
178 pleopod in males, and the shape of the endopod of the first pleopod (Komai & Segonzac, 2008).
179 Since these sexual characters appear during the transition from subadult to adult stage, sex of
180 juvenile and subadult specimens could not be determined. Brooding females are characterized
181 by the presence of embryos held between their pleopods under the abdomen, and by
182 modifications of their pleopods (addition of setae to maintain the brood). Hatched females
183 (females just after larval release but prior to molt) were identified by their modified pleopods.
184 Brooding and hatched females are referred to as ovigerous females (following Nye et al. 2013).
185 Young small adults and subadults resemble females (*i.e.* lack of appendix masculina) and also in
186 some cases lack gonadal tissue that could be used for sex determination. In order to estimate the
187 minimal size for confident determination of sex, a subsample of adult and subadult shrimps in
188 small size classes (7 to 15 mm carapace lengths) were dissected to verify macroscopic evidence
189 of gonadal tissue and estimate the onset size of sexual differentiation (OSD, see section 2.4,
190 equivalent to the “minimum sexable size” of Anger & Moreira, 1998).

191 Carapace length (CL) was measured from the posterior margin of the ocular shield to the
192 mid-posterior margin of the carapace in adults and subadults (Fig. 2a), with a precision of 0.1 mm.

193 In juvenile stages, CL was measured from the anterior tip of the rostrum to the posterior margin
194 of the carapace (Fig. 2b). Morphological changes between juvenile and adult stages (rostrum
195 reduction and development of the ocular shield) may introduce a bias in our measurements but
196 this was small compared to the total length of the carapace, and had little impact on size frequency
197 distributions and size comparisons.



198
199 Figure 2. Size measurements of different life stages of *Rimicaris exoculata*. A) Adults and
200 subadults. B) Juveniles. C-H) Embryos and hatched larvae. Early developmental stage: C)
201 Fertilized egg (with position of measurements for maximum (d1) and minimum (d2) diameter), D)
202 Blastula stage; Mid developmental stage: E) Early gastrulation, F) Nauplius stage; Late
203 developmental stage: G) Pre-hatch embryo (arrow indicates eye spot). H) Hatching Zoea larva.
204 CL: carapace length. Scale bars C-H: 500 μ m.

205 Embryos were removed from the abdomen of brooding females, counted and staged. We
206 classified embryonic developmental stages into three categories, similar to those defined by Nye
207 et al (2013) for *R. hybisae*. Early stage embryos encompass freshly laid fertilized eggs without
208 cellular division, and dividing eggs until the blastula stage (Figs. 2C, D). Mid stage starts with
209 gastrulation when a clear region differentiates at one pole of the embryo and extends to the end
210 of the naupliar development (Figs. 2E, F). Late stage includes post-naupliar development, when
211 eye pigmentation becomes visible, abdomen appears clearly separated from the rest of the body
212 with yolk in the cephalothorax, and appendages are fully lengthened and appear curled around
213 the cephalothorax (Fig. 2G). For each brood, ten embryos were randomly selected and both their
214 maximum and minimum diameters were measured at a precision of 0.03 mm under a
215 stereomicroscope with a graduated ocular. The volume of embryos was estimated according to
216 Oh & Hartnoll (2004), considering a spheroid volume as $v = (4/3) \pi r_1 r_2^2$, where r_1 and r_2 are half
217 of the maximum and minimum axis, respectively. This estimation has a precision of 1.6×10^{-5}
218 mm³.

219 During examination of female broods, we found dirivultid copepods between the embryos
220 or attached at the base of the pleopods. They were identified to the species level through
221 barcoding (Suppl. Table 1).

222

223 **2.3. Genetic identifications**

224 DNA was extracted from shrimp juveniles and copepods using the CTAB method (Doyle
225 1990) on muscle tissue or on whole specimens for copepods. A section of the cytochrome oxidase
226 I gene (COI) was amplified in a 50 μ L solution of 1X reaction buffer, 2 mM MgCl₂, 0.25 mM dNTP,
227 1.2 units of Taq polymerase and 0.6 mM of each primer (LCOI1490 and HCOI2198, Folmer et al.
228 1994). Amplifications were performed as following: initial denaturation (5 min at 95°C), 40 cycles

229 including denaturation (1 min at 94°C), annealing (1 min at 52 °C) and extension (2 min at 72 °C),
230 followed by a final extension of 7 min at 72°C. All PCR amplifications were conducted on a
231 GeneAmp PCR system 9700 (Applied Biosystems). PCR products were purified and sequenced
232 by Macrogen, Inc. (Netherlands) using the amplification primers. We aligned our sequences using
233 MUSCLE (Edgar 2004), along with a set of alvinocaridid and other shrimps sequences. For our
234 dirivultid copepod sequence, a subset of the sequences obtained by Gollner et al. (2011) was
235 used for comparison. Neighbor-joining trees were constructed using Geneious R8 software
236 (Kearse et al. 2012) using a HKY evolutionary model of nucleotide substitution. Robustness of
237 the inferred trees was evaluated using bootstrap method with 1000 replicates.

238

239 **2.4. Data analysis**

240 Onset of sexual differentiation (OSD) was estimated using a similar procedure as what
241 was used for determination of the size of sexual maturity in Crustacea (Wenner et al. 1974).
242 Proportions of specimens with gonad tissue were estimated for size classes between 7-15 mm
243 CL (larger specimens with clear sex differentiation were not included). The proportion of
244 specimens with gonad differentiation were plotted against size classes and fitted to the logistic
245 equation:

$$246 \quad P_{sd} = \frac{1}{1 + e^{(a - b \cdot CL)}}$$

247 where P_{sd} is the proportion of shrimps with gonad differentiation, CL is the carapace
248 length, and a and b are constants. The size at which 50 % of the individuals exhibit sexual
249 differentiation denoted by gonad development was considered as an estimation of OSD (Suppl.
250 Fig. 3). Subadults were subsequently identified as individuals resembling adults with sizes < OSD.

251 Similarly, the size of effective sexual maturity (ESM) for females was estimated from the
252 proportion of ovigerous females per size class, corrected by the maximum proportion of ovigerous
253 females (King 2007). We applied the previous logistic equation, substituting Psd by the corrected
254 proportion of ovigerous females ($Povf$) on the equation. The ESM was estimated by the size at
255 which $Povf$ is 50% (Suppl. Fig. 4).

256 For each sample, juvenile ratio, subadult ratio and sex ratio (F:M) were estimated.
257 Deviation from a sex ratio of 1:1 was tested with a χ^2 test, using Yates correction in samples with
258 few specimens ($n < 30$). In order to determine the variations in sex ratio within each habitat at
259 each vent field, we performed a heterogeneity χ^2 test (Zar 2010). Variations in the proportions of
260 females with size $>$ ESM and the proportions of ovigerous females between vents were tested
261 using χ^2 test.

262 Size class structures were analyzed for each sample, estimating kurtosis and skewness
263 for size class aggregation and deviation from the mean, respectively (Zar 2010). Although
264 histograms were elaborated denoting juveniles, subadults, males and females, size structure
265 comparisons were performed including all specimens. Normality tests were performed for each
266 sample using the Shapiro-Wilk test (Zar 2010). Discrete size cohorts in the samples were
267 identified using the statistical package `mixdist` (MacDonald 2003) running also in RTM. The
268 goodness of fit of the identified size cohorts was verified using χ^2 test. The analyses were
269 performed for each sample collected in the AEH and compared to verify the consistency of
270 identified cohorts. Identification of cohorts in other habitats were not performed due to insufficient
271 sample size.

272 For AEH samples, differences in body sizes associated with sex and vent fields were
273 tested using multifactorial analysis of variance (ANOVA), after \log_{10} transformation. For this
274 analysis, samples were nested at the factor vent field, representing the variation in body size

275 between samples collected within the same vent field. Similarly, an ANOVA test was performed
276 in order to examine differences in male body sizes between habitats (AEH vs DEH) at the TAG
277 vent field. In this case, samples were nested at the factor habitat which represents the small-scale
278 variation. Data normality and homoscedasticity were tested using χ^2 for frequency distribution and
279 Levene test respectively (Underwood 1997, McGuinness 2002).

280 In crustaceans, brood size increases proportionally with female size (Oh & Hartnoll 2004).
281 In order to compare reproductive outputs at the two vent fields, we estimated the relation between
282 realized fecundity (number of embryos per brood) and female size (CL) using regression
283 analyses, after \log_e transformation. The difference in size-specific fecundity between vent fields
284 was tested using a t-test analysis. Variations in embryo size associated with parental female,
285 embryo stage and vent fields were analyzed with a multifactorial ANOVA test. For this analysis
286 the factor parental female was considered as nested to the combination of vent field and embryo
287 stage.

288

289 **3. Results**

290

291 **3.1. *Rimicaris* shrimp populations at TAG and Snake Pit in January 2014**

292 Overall, 3445 individuals from 14 samples collected at both vent fields were examined.
293 Among these, we determined developmental stage and sex for 3388 individuals and measured
294 3379 individuals (missing data are due to body damages preventing accurate measurements or
295 sex/stage determination). The global dataset includes 1925 females (56.8%), 292 males (8.6%),
296 882 subadults (26.1%) and 289 juveniles (8.5%). Global sex-ratio clearly deviates from 1:1 (χ^2 ,
297 $df= 1$, $p < 0.01$). Of the 1925 females, 136 (7.1%) are either brooding eggs (125), or have recently
298 hatched larvae (11).

299 Smaller specimens are early juveniles (CL = 4.4 mm), whereas the largest one is a female
300 with 24.4 mm CL (but being an outlier specimen as other largest females reach ~20.6 mm CL).
301 Size-range of the juveniles varies between 4.4 and 10.3 mm, and overlaps subadult size range
302 (7-9.9 mm CL) (Suppl. Table 2). Subadult maximal size was determined by our estimation of the
303 onset size of sexual differentiation (OSD = 10 mm, Suppl. Fig. 3). The OSD value is also
304 consistent with the size of the smaller adult male found in our samples (CL = 9.9 mm). Although
305 the size of some juveniles exceeds the OSD size, they are morphologically distinct from
306 subadults: their rostrum is not completely reduced, and their carapace not fully inflated.

307 Overall, size ranges of males and females are similar with respective CL ranges = 9.9-
308 19.1 mm and 10 -24.4 mm (Suppl. Table 2). However, the average size of males was higher than
309 the average size of females, with 15.1 mm CL and 12.5 mm CL respectively (t-test= 20.71, p<
310 0.001). Most ovigerous females exhibit large sizes, with CL ranging from 12 mm to 20.6 mm
311 (average size: 16.5 mm CL). We estimate the size at effective sexual maturity (ESM) at 15.1 mm
312 CL for females (Suppl. Fig. 4). Hereafter, females with size \geq 15.1 mm CL are called sexually
313 mature females.

314

315 **3.2. Variation in population structure across habitats and vent fields**

316

317 **3.2.1. Different types of habitats**

318 Visually, striking differences characterized shrimp assemblages when we collected our 14
319 samples. Most samples (9) were collected among dense aggregations of actively swimming
320 shrimps gathering around vigorous fluid emissions (Fig. 1-B, Suppl. Fig. 1). In these aggregations,
321 we recorded steep temporal variations in temperature with maximum varying from 14 to 33°C
322 (Table 1). Resting shrimps scattered on inactive sulfide substratum at the periphery of dense

323 aggregations were collected 3 times at TAG (Fig. 1-C, Suppl. Figs. 2 A-C). In this habitat, no fluid
324 exit was visible and temperature was low and stable, with a maximum of 2.8°C (Table 1), whereas
325 ambient seawater temperature was 2.6°C. At TAG, aggregations of very small individuals
326 characterized by their bright red color were sampled around diffusions of translucent fluids exiting
327 from very small chimneys or cracks (Fig. 1-D, Suppl. Figs. 2 E, F). Temperatures among those
328 young individuals varied from 2.8°C to 5.3°C (Table 1). We thus defined 3 types of aggregations
329 characterized by different temperature regimes, with visually distinct shrimp assemblages in terms
330 of density, size and behavior. These are called Active Emission Habitat (AEH), Inactive Emission
331 Habitat (IEH) and Diffuse Emission Habitat (DEH).

332 3.2.2. *Variations in population structure between habitats (TAG)*

333 At TAG, we observe striking differences in terms of population structure, size-frequency
334 distribution, and reproductive features among the three habitats. In AEH, with 71% of females
335 and 8,5% of males, sex-ratio is clearly biased for females (8.4:1). Although sex ratios are
336 significantly different among AEH samples from TAG ($\chi^2_{het}= 50.05$, $df= 2$, $p< 0.001$, Table 2), all
337 of them are significantly female biased (χ^2 , $df= 1$, $p< 0.02$ in all cases, Table 2).

338 In contrast, in IEH, 90.3% of the individuals are males, while females represent only 6.5%
339 of the shrimps collected. Sex ratio is significantly biased for males (1:14) and in each TAG IEH
340 samples (χ^2 , $df= 1$, $p\leq 0.002$ in all cases, Table 2).

341 Subadults represent 17.5% of the individuals on average in AEH (Table 2), with
342 significant variation between samples (from 3% to 23.2% of the individuals in each sample, $\chi^2_{het}=$
343 16.4, $df= 2$, $p< 0.001$), but they are rare in IEH with only two individuals collected overall.

344

345 Table 2. *R. exoculata* specimens and sex ratios at different vent fields and habitats. J: juveniles,
 346 <OSD (onset of sexual differentiation): subadults, F: females (non-brooding); OF: ovigerous
 347 females; M: males.

Active Emission Habitat (AEH)										
Vent field	Sample	J	<OSD	F	OF	M	n	F : M	χ^2	p
Snake Pit	S01	1	8	109	1	20	139	5.5:1	62.308	<0.001
Snake Pit	S02	14	175	181	2	9	381	20.3:1	157.688	<0.001
Snake Pit	S03	6	177	522	29	53	787	10.6:1	410.603	<0.001
Snake Pit	S04	16	188	174	0	6	384	29:1	156.8	<0.001
Snake Pit	S05	26	22	39	1	20	108	2:1	6.667	0.01
Snake Pit	S06	21	143	158	22	19	363	9.5:1	130.256	<0.001
Total								9.8:1	904.264	<0.001
								Heterogeneity	20.056	0.001
TAG	S07	19	138	380	35	23	595	18:1	350.831	<0.001
TAG	S08	1	6	163	25	7	202	26.7:1	168.005	<0.001
TAG	S09	6	23	58	20	51	158	1.5:1	5.651	0.017
Total								8.5:1	472.441	<0.001
								Heterogeneity	50.046	<0.001
Inactive Emission Habitat (IEH)										
TAG	S10	1	0	2	0	15	18	1:7.5	9.941	0.002
TAG	S11	0	2	2	1	33	38	1:16	25	<0.001
TAG	S12	0	0	1	0	36	37	1:35	33.108	<0.001
Total								1:16	67.6	<0.001
								Heterogeneity	0.449	0.993
Diffuse Emission Habitat (DEH)										
TAG	S13	77	0	0	0	0	77			
TAG	S14	101	0	0	0	0	101			

348

349 Ovigerous females are almost exclusively observed in AEH at TAG (only one hatched
 350 female observed in IEH and none in DEH), representing 11.7 % of the females on average, with
 351 strong variations between samples (from 8.4% to 25.6%). These variations reflect both variations
 352 in the proportion of sexually mature females among all females (22.4% on average, with variations
 353 between samples: 13.3% to 64.9%), and, to a lesser extent, variations in the proportion of

354 ovigerous females among sexually mature females (36.8 % on average, with variations between
355 samples: 29.8% to 43.6%).

356 Juveniles are not abundant in AEH (less than 3% of the total population, Table 2, no
357 significant heterogeneity between samples: $\chi^2_{het}= 3.64$, $df= 2$, $p= 0.602$), and are rare in IEH with
358 only one early stage juvenile collected. In contrast, DEH samples are exclusively composed of
359 early stage juveniles. Although they were not collected, few large adult individuals (both *R.*
360 *exoculata* and *R. chacei*) were observed crawling around these nurseries (*in situ* observations
361 and Fig. 1-D).

362 3.2.3. Variation in population structure between vent fields (AEH)

363 AEH Snake Pit samples exhibit similar population structure to those from TAG, with a
364 strong dominance of females. Females, males, subadults and juveniles represent respectively
365 57.2%, 5.9%, 33 % and 3.9% of the overall population. Sex-ratio is female-biased (9.8:1 overall)
366 and similar to the ratio observed in AEH at TAG. Like in TAG AEH samples, sex-ratio vary
367 significantly among AEH samples at Snake Pit ($\chi^2_{het}= 20.06$, $df= 5$, $p= 0.001$, Table 2), but all
368 are significantly female-biased (χ^2 , $df= 1$, $p\leq 0.01$ in all cases).

369 Overall, subadults are more abundant in Snake Pit samples than in TAG samples,
370 representing almost a third of the population. However, their proportion varies strongly between
371 samples (from 5.8% to 45.9% of the individuals, $\chi^2_{het}= 164.9$, $df= 5$, $p< 0.001$). Like in TAG AEH,
372 the proportion of juveniles in Snake Pit samples is generally low, except in one sample where
373 they reach 24.1% of the individuals, resulting in significant heterogeneity between Snake Pit
374 samples ($\chi^2_{het}= 89.32$, $df= 5$, $p< 0.001$).

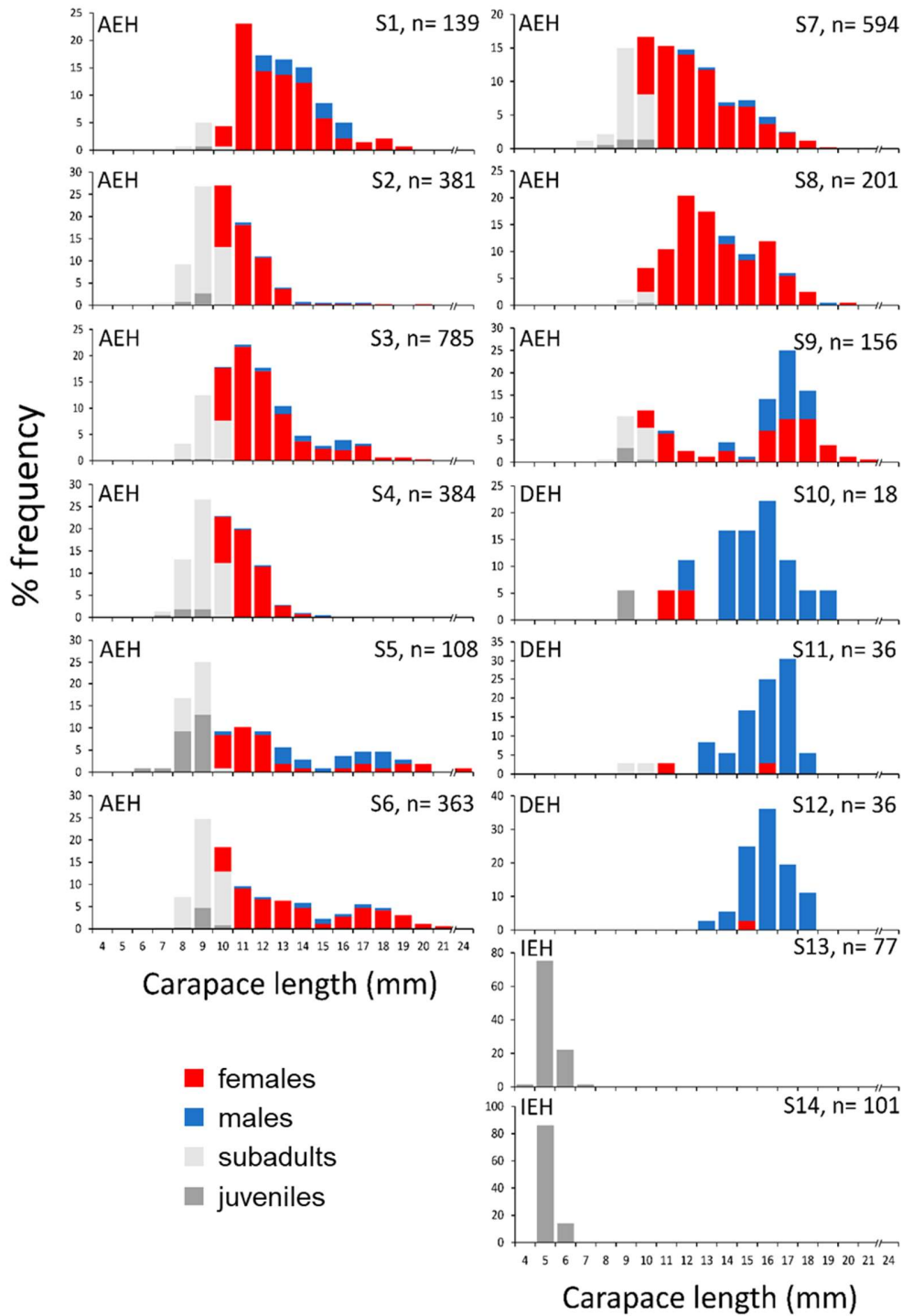
375 Although the proportion of ovigerous females among sexually mature females is similar
376 between vent fields (36.6 % on average at Snake Pit, $\chi^2= 0.003$, $p= 0.956$), the proportion of

377 sexually mature females among all females is significantly lower in Snake Pit (11.7 %) than in
378 TAG (22.4 %) AEH samples ($\chi^2= 34.24$, $p < 0.001$), resulting in lower proportion of ovigerous
379 females overall (4.4% of all females).

380 *3.2.4. Variations in size frequency distributions among habitats and vent fields*

381 Overall, reflecting the differences in sex and stage distributions between habitats, size
382 distributions also differed between habitats. While DEH host very small shrimps almost not
383 represented in other habitats, mostly large individuals inhabit IEH. In AEH, shrimp sizes vary over
384 a wide range, overlapping slightly both size ranges of shrimps in DEH and IEH (Fig. 3, Suppl. Fig.
385 5).

386 Size frequency distributions vary among samples and habitats in both vent fields, both
387 in terms of kurtosis and skewness (Fig. 3). General trends in AEH size frequency distributions
388 include bias towards small sizes (skewness= 0.945) and slightly leptokurtic distribution (kurtosis=
389 0.423). In some samples, the distribution is clearly non-unimodal. In others, deviation in skewness
390 and kurtosis suggests a mixture of cohorts. Based on a size cohort analysis, 5 different cohorts
391 are identified overall in AEH (Table 3). In individual samples, cohort number vary between 2 to 4
392 (Table 3, Suppl. Fig. 6), with some correspondence observed between samples across the two
393 vent fields. Overall in AEH, we identify one cohort of juveniles and subadults, one cohort of
394 subadults and small adults (<12 mm CL) and three cohorts of adults, two of which include females
395 \geq ESM. In both vent fields, the cohorts corresponding to juveniles, subadults and small adults
396 represent an important proportion of the population of AEH.



397

398 Figure 3. Size class structure of *R. exoculata* in different habitats of the Snake Pit (left side) and
 399 TAG (right side) vent fields. AEH active emission habitat, DEH diffuse emission habitat, IEH
 400 inactive emission habitat.

401 Table 3. Identified cohorts in AEH at the Snake Pit and TAG vent fields. Mean and standard
 402 deviations are shown for each sample, proportion of each cohort are in brackets. χ^2 denote the
 403 deviation of the sample from the cohort estimation. ns: non-significant.

Cohort:	1	2	3	4	5	χ^2
Snake Pit						
S1		11.45±1.75 (0.59)	13.24±2.03 (0.41)			12.93ns
S2	8.86±0.87 (0.61)	10.78±1.06 (0.37)		16.14±1.58 (0.02)		3.058ns
S3	9.14±0.89 (0.33)	11.25±1.09 (0.54)		15.49±1.50 (0.13)		11.16ns
S4	8.45±0.77 (0.50)	10.47±0.96 (0.48)	12.92±1.18 (0.02)			1.995ns
S5	8.30±0.87 (0.52)	11.40±1.19 (0.28)			17.28±1.80 (0.20)	15.32ns
S6	8.70±0.70 (0.47)	10.44±0.84 (0.17)	12.81±1.03 (0.17)		17.11±1.38 (0.19)	3.247ns
TAG						
S7	9.83±1.40 (0.59)		13.38±1.90 (0.41)			25.11*
S8		10.12±0.83 (0.15)	12.17±1.00 (0.51)	15.39±1.27 (0.34)		3.556ns
S9	8.98±0.53 (0.21)	10.50±0.62 (0.11)	13.37±0.79 (0.06)		16.73±0.99 (0.62)	8.230ns

404 *Despite the deviation of the size structure from the cohort model ($p=0.0485$), it was the closest
 405 model to the input data

406

407 Male and female body sizes in AEH are significantly different among samples, indicating
 408 small scale variations in sizes, within habitats at each vent field (ANOVA, $p < 0.001$, Suppl. Table
 409 3). Body sizes also vary significantly with sex (ANOVA, $p < 0.001$, Suppl. Table 3), males being
 410 larger than females at each vent field (Suppl. Table 2). Size distribution of ovigerous females
 411 clearly departed from the rest of the females, with sizes similar to the male average size in TAG
 412 and even larger than the male average size in Snake Pit. Although males and females tend to be
 413 smaller in Snake Pit than in TAG AEH, size differences are not significant between the two vent
 414 fields for each sex (ANOVA, $p=0.083$, Suppl. Table 3). In contrast, ovigerous females are slightly

415 larger in Snake Pit than in TAG AEH, but this variation between the two vent fields was not
416 statistically significant (ANOVA $F = 1.649$, $p = 0.246$, $df_2 = 6$).

417 Size frequency distributions of IEH samples are leptokurtic (kurtosis 0.12 to 1.99) and
418 biased towards larger sizes (skewness -0.45 to -1.52). Males do not exhibit different body sizes
419 between AEH and IEH, although significant differences are detected between samples of a given
420 habitat (ANOVA, $p < 0.001$, Suppl. Table 4). Females also exhibit similar size ranges between
421 both habitats, although the low number of females collected in IEH prevents statistical
422 comparisons.

423 3.2.5. Juvenile distribution in AEH and DEH: detection of *Rimicaris chacei* nurseries

424 Juvenile sizes at the AEH are similar between vent fields, only showing differences
425 between samples (ANOVA $F_{vents} = 5.2001$, $p = 0.057$, $df_2 = 7$, $F_{samples} = 2.643$, $p = 0.015$, $df_1 = 7$, $df_2 =$
426 101). However, juveniles from DEH are much smaller (ANOVA $F_{vents} = 804.91$, $p < 0.001$, $df_2 = 3$,
427 $F_{samples} = 1.580$, $p = 0.195$, $df_1 = 3$, $df_2 = 199$) and form a distinct cohort from that of AEH. Based on
428 morphological features described in Komai & Segonzac (2008), we suspected that juveniles from
429 DEH were possibly a mixture of *R. exoculata* and *R. chacei*. Molecular identification using COI
430 barcode reveals that all sequences of these juveniles are consistent with *R. chacei* affiliation,
431 whereas juveniles from AEH (with clear *R. exoculata* juvenile morphology according to Komai &
432 Segonzac 2008) are consistent with *R. exoculata* affiliation (Suppl. Fig. 7).

433

434 3.3. Reproductive features

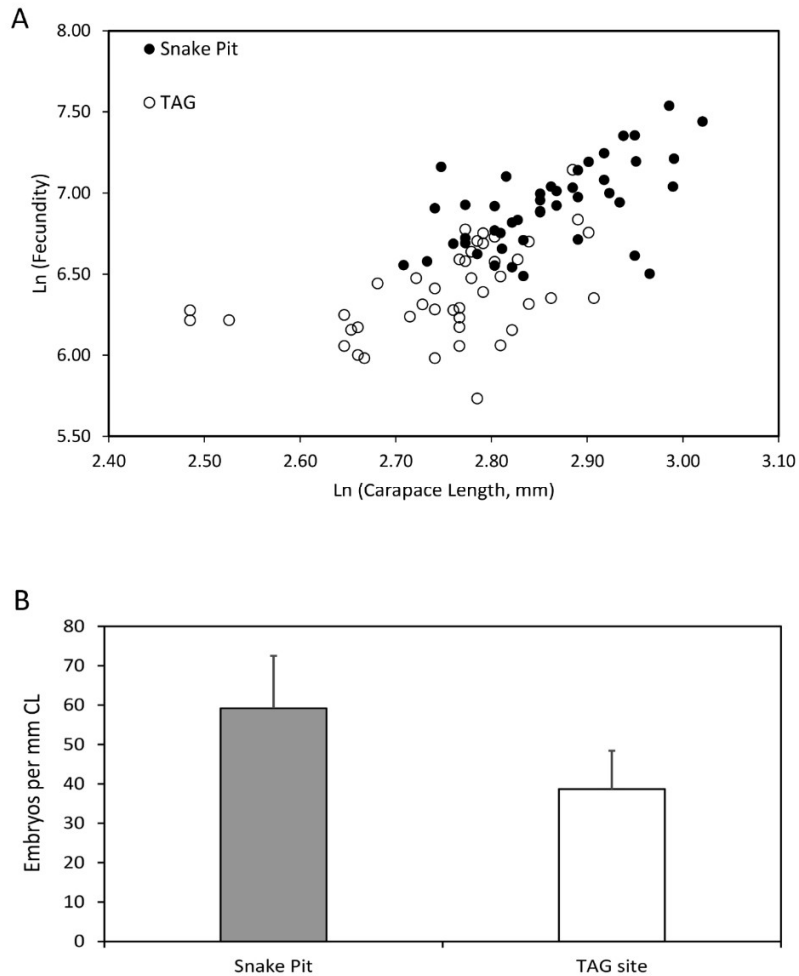
435

436 3.3.1. Fecundity

437 Of the 125 brooding females found in our samples, 36 had obviously lost part of their
438 broods, not because of hatching -since embryos were clearly not yet fully developed- but rather
439 due to either abortion or more probably lost during sampling. These are not included in our
440 fecundity analyses.

441 Fecundity varies from 304 eggs in a female from TAG with 16.2 mm CL, to 1879 eggs in
442 a female from Snake Pit with 19.8 mm CL. The largest brooding females are observed at Snake
443 Pit (with 1704 eggs and 20.5 mm CL for the largest), while the smallest brooding females are from
444 TAG (with 500 and 532 eggs for the two smallest -12 mm CL- individuals). The average fecundity
445 is overall of 833 eggs, and is higher among Snake Pit brooding females (1045 eggs, with an
446 average CL of 17.4 mm) than among TAG ones (616 eggs, with an average CL of 15.9 mm). As
447 expected, a positive correlation is observed between carapace length of the females and fecundity
448 (Pearson correlation, $R= 0.682$, $t\text{-test}= 8.71$, $p< 0.001$) (Fig. 4A). We consider that more data are
449 necessary to estimate accurate linear regression models of fecundity and compare differences
450 between populations or with other alvinocaridid shrimps.

451 Overall size-specific fecundity ranges from 19 to 95 embryos.mm⁻¹ CL. The size-specific
452 fecundity of females does not change with the developmental stage of the brood (Early vs Mid
453 stage, $t\text{-test}= 0.98$, $p= 0.164$, $df=57$; Mid vs Late stage, $t\text{-test}= 0.17$, $p= 0.432$, $df=57$), but is
454 significantly lower in females from TAG (39 ± 10 embryos.mm⁻¹ CL) than in females from Snake
455 Pit (59 ± 13 embryos.mm⁻¹ CL) ($t\text{-test}=8.16$, $p<0.001$, $df=87$) (Fig. 4B). Finally, more females with
456 damaged broods are observed at TAG, although we could not identify the cause of loss (abortion
457 or sampling).



458

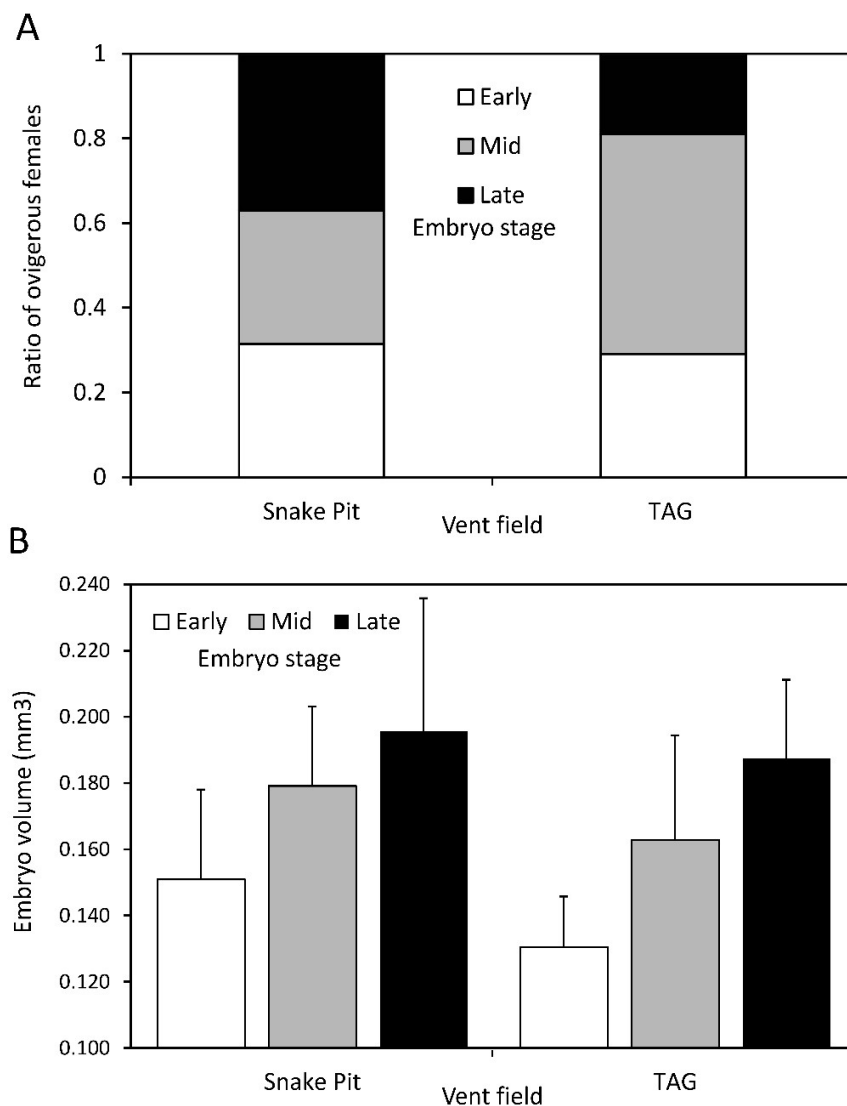
459 Figure 4. Fecundity in *R. exoculata* from TAG and Snake Pit vent fields. A. Number of embryos
460 related with female size (carapace length). B. Size-specific fecundity, number of embryos per unit
461 of size (mm of carapace length).

462

463 3.3.2. Reproductive synchrony

464 Within each individual brood examined, all eggs are at the same developmental stage
465 (early, mid or late), except for occasional dead embryos or non-fertilized eggs. However, embryos
466 at all developmental stages are observed in females from both vent fields, showing a lack of
467 synchrony between the broods of different females. Overall, the distribution of brood stages are
468 different between vent fields ($\chi^2= 7.097$ $p=0.014$), with some variability between samples of a

469 given vent field (Suppl. Fig. 8). At both vent fields, a third of the females carry early stage broods,
470 however at Snake Pit late-stage brooding and hatched females were slightly more frequent (37%)
471 than at TAG (19%). At TAG, most of the brooding females are at the mid stage (51.9%) (Fig. 5A).
472



473
474 Figure 5. Characteristics of *R. exoculata* broods at TAG and Snake Pit vent fields. A. Proportion
475 of broods with eggs at each developmental stage (including hatched females as having late
476 broods). B. Sizes of individual embryos at different developmental stages.

477

478 3.3.3. *Egg sizes and development*

479 The volume of the eggs within the brood of each female shows significant heterogeneity
480 due to inter-individual variations, developmental stage of the broods, and vent fields (Supp. Table
481 5). Despite individual variations, a clear trend of egg volume increase with developmental stages
482 is observed at both vent fields (Fig. 5B). The ratio between the minimum and maximum diameters
483 of embryos decreases along their development, with minimum diameter being on average 0.87
484 of the maximum diameter in early stage, and 0.76 of the maximum diameter in the late stage.
485 Eggs thus become more elongated at the end of embryonic development, which may reflect an
486 increase of the embryo polar axis, the distribution of the structures inside the envelope and water
487 uptake.

488 At each stage, embryos in TAG broods are smaller than in Snake Pit broods (Figure 5B).
489 The volume of early stage embryos is $0.151 \pm 0.027 \text{ mm}^3$ at Snake Pit, and $0.131 \pm 0.015 \text{ mm}^3$
490 at TAG ($t\text{-test}= 8.386$, $p < 0.001$, $df=242$). At mid-stage, the volume of embryos increases to 0.179
491 $\pm 0.024 \text{ mm}^3$ at Snake Pit and $0.163 \pm 0.032 \text{ mm}^3$ at TAG ($t\text{-test}= 3.719$, $p < 0.001$, $df= 116$). In
492 the late stage, the volume of embryos reaches $0.196 \pm 0.040 \text{ mm}^3$ at Snake Pit and 0.187 ± 0.024
493 mm^3 at TAG ($t\text{-test}= 2.458$, $p= 0.007$, $df= 238$).

494 3.3.4. *Copepod occurrence within broods*

495 The copepods collected from the brooding females were identified as *Stygiopontius*
496 *pectinatus* using molecular barcodes (Suppl. Fig. 9). These copepods appear to be truly
497 associated with eggs because they are found deep inside the broods usually attached to the setae
498 at the basis of the pleopods. In addition, no copepods were found on the abdomen of non-
499 brooding females. At Snake Pit, 33 % of the brooding females are colonized with copepods,

500 whereas only 20 % of them are infected at TAG. The proportion of infested females tends to
501 increase between early (10%) and late (40%) stage broods ($\chi^2= 6.540$, $p= 0.038$, Suppl. Figure
502 10). This trend is observed at each vent field and for each sample. In infested broods, the number
503 of copepods per brood vary between 1 and 5, and appears to be higher at Snake Pit (2 copepods
504 per brood in average, range: 1-5 copepods) than at TAG (1.1 copepods per brood in average,
505 range: 1-2 copepods). Similarly, the average number of copepod per brood tends to increase with
506 brood age from 1 copepod/brood in average in early broods to 2 copepod/brood in average in late
507 broods.

508

509 **4. Discussion**

510 In January-February 2014, samples of *Rimicaris exoculata* populations at TAG and Snake
511 Pit vent fields revealed strong female-biased sex ratios (87% females among adults overall). Sex
512 ratio biases were also striking locally, reflecting abrupt changes in population structure among
513 habitats. Dense shrimp assemblages crawling next to high temperature fluid exits (AEH)
514 consisted mainly of females and immature individuals, whereas shrimps observed scattered in
515 the cold and stable periphery of active vents (IEH) were almost only adult males. At TAG, near
516 low temperature diffusion areas (DEH), gathering of very small juveniles absent from AEH, and
517 previously hypothesized to be early juveniles of *R. exoculata*, were indeed *R. chacei* nurseries.
518 Ovigerous females were observed in larger proportion than ever reported so far for the species,
519 representing about a third of the sexually mature females in AEH. Overall, these patterns were
520 consistent across both vent fields, although a high degree of heterogeneity in population structure
521 was observed at small spatial scales, between samples from a given habitat. Ovigerous females
522 were more abundant at TAG, reflecting a higher proportion of sexually mature females. However,

523 lower fecundities, smaller eggs as well as a higher proportion of aborted broods, were observed
524 in ovigerous females from TAG, suggesting a lower individual reproductive effort.

525

526 **4.1. Spatial variation in *R. exoculata* population structure and reproduction**

527 Our study describes strongly biased sex ratios in *R. exoculata*, where sexes appear to
528 segregate between different habitat types. Similar evidence has been previously reported by
529 Shank et al. (1998) in samples collected from active chimney walls of several MAR vent fields,
530 where females were found in much larger numbers than males. Although variations in sex ratios
531 between the different vent fields were observed, variations within vent fields could not be
532 assessed because unique samples were collected in each field (Shank et al. 1998). In 2007,
533 shrimps collected in AEH at Logatchev during the Serpentine cruise also showed a female-
534 dominated population (I. Hernández-Ávila, unpublished data). Population structure of shrimps in
535 IEH has not been reported previously, probably because individuals scattered at the periphery of
536 dense aggregates have been considered as remains of the main populations observed in AEH,
537 rather than individuals preferentially occupying a specific and separated habitat. Therefore,
538 similarly to our observations, variations in sex ratios observed in previous studies could indeed
539 reflect small-scale local variations, rather than variations between vent fields.

540 Female-biased sex ratios have been reported in several vent crustaceans, often
541 associated with different spatial patterns among sexes. For instance, Nye et al. (2013) reported
542 that populations of *Rimicaris hybisae* from the Cayman Trough have a sex ratio in favor of females
543 close to the vent emissions, while more dispersed populations are dominated by males at their
544 peripheries, with some degree of local variability. In brine pools of the Gulf of Mexico, *Alvinocaris*
545 *stactophila* shows overall female-biased sex ratio with males preferentially locating on the outer
546 part of mussel beds (Copley & Young 2006). At hydrothermal vents of the East Scotia Ridge, the

547 chirostylid crab *Kiwa tyleri* exhibits similar -but inverse- patterns: areas close to vent fluids
548 emission are occupied by dense male-dominated aggregations, whereas females are more
549 numerous at the periphery (Marsh et al. 2015). *A. muricola* inhabiting cold seeps off Congo also
550 revealed a globally female-biased sex ratio, with significant local variation observed between
551 limited numbers of samples (Ramirez-Llodra & Segonzac 2006). Other vent shrimps such as the
552 Hippolytidae *Lebbeus virentova* from the Cayman Trough also exhibit female-biased sex ratio,
553 but spatial variation was not reported (Nye & Copley 2014).

554 Immature individuals including juveniles were observed in AEH and IEH. However, both
555 habitats were occupied by apparently different life stages, with very small juveniles in IEH,
556 whereas AEH harbored larger juveniles and subadults. Small juveniles of IEH had inconsistent
557 morphologies with a mixture of characters described for both early juveniles of *R. exoculata* (stage
558 *A sensus* Komai & Segonzac 2008) and juveniles of *R. chacei*. Barcoding of these IEH juveniles
559 revealed that they were indeed all *R. chacei*, whereas larger juveniles in AEH belonged to *R.*
560 *exoculata*. This observation has been repeated in 2017 and 2018, with large aggregations of small
561 juveniles of *R. chacei* observed around vent diffusion areas at the periphery of TAG and Snake
562 Pit vent fields (Methou et al. 2020). A more systematic examination of morphological characters
563 along with molecular characterization led to a redefinition of the juvenile stages of both *Rimicaris*
564 species (Methou et al. 2020). Our nurseries here are thus specific of *R. chacei*, whereas juveniles
565 of *R. exoculata* appear to be found only in AEH, where they grow to the subadult and adult stages.
566 Although this is not shown in details here, juveniles corresponding to the 2 juvenile stages of *R.*
567 *exoculata* described in Methou et al. (2020) occurred in our AEH samples. Variations in the
568 proportions of juveniles and subadults in AEH may hide further structuration at very small spatial
569 scales, perhaps corresponding to environmental gradients associated with vent fluid and
570 seawater mixing. Additional observations are necessary to elucidate such microscale population
571 structure within AEH.

572 **4.2. Reproductive efforts of *Rimicaris* and potential environmental effects**

573 The size for sexual differentiation (OSD) is 10 mm, about half of the maximal size of the
574 species (not considering our single outlier). While the OSD parameter has practical value for sex
575 separation in studies of population structure, size at first reproduction represents a key life-history
576 parameter reflecting the life-time investment in reproduction of a species (Anger & Moreira 1998).
577 In Alvinocaridid species, the size at first reproduction varies between 50% (*R. hybisae*, Nye et al.
578 2013) and 60% (*A. muricola*, Ramirez-Llodra & Segonzac 2006, *A. stactophila*, Copley & Young
579 2006, *M. fortunata*, Ramirez-Llodra et al. 2000) of the maximal size of the species. *R. exoculata*
580 thus falls within the range of its family, with the smallest brooding females measuring 12 mm CL,
581 which represents 58% of its maximal size (20.6 mm CL, not considering our outlier). However,
582 few females between 12 mm CL and 15 mm CL (i.e. < ESM) were brooding eggs in our samples
583 (3.5%) and likely represent 'premature' specimens, while much more become sexually mature
584 (36.5%) when they reach 15.1 mm CL (ESM), which represents 73% of the species maximal size
585 (excluding outliers). Shrimps of the family have been reported as iteroparous (Copley & Young
586 2006, Nye et al. 2013, Ramirez-Llodra et al. 2000), and a later onset of reproduction might limit
587 life time investment in reproduction. On the other hand, since large females have larger broods,
588 favoring reproduction to the largest individuals might be advantageous and help to maximize
589 energy investment in reproduction.

590 Fecundity in *R. exoculata* was on average of 833 eggs/female, ranging from 304 to 1879,
591 with brood sizes increasing with female sizes, which is common in carideans (Corey & Reid 1991).
592 Previous counts on the rarely collected ovigerous females fall within this range. Ramirez-Llodra
593 et al. (2000) reported a female from Snake Pit (Microsmoke cruise, November 1995) with 988
594 eggs (16.4 mm CL) and a female from TAG (Bravex cruise, September 1994) with 836 eggs (17.7
595 mm CL). *R. exoculata* appears to have fecundities similar to those reported for *R. hybisae*
596 (maximal fecundity of 1707 eggs, Nye et al. 2013) but lower than fecundities of *R. chacei* (2510

597 eggs reported in a female collected in June 1994 at Lucky Strike, Ramirez-Llodra et al. 2000).
598 Egg sizes of *R. exoculata* are also consistent with the previous report of Ramirez-Llodra et al.
599 (2000): early stage (blastula) embryos had a volume of about 0.145 mm³ (volume are calculated
600 hereafter from diameters given in reference papers, with the spheroid formula used in this paper)
601 at TAG, within the size range we observed in this study at similar developmental stage: 0.131
602 mm³ at TAG and 0.151 mm³ at Snake Pit. Eggs of *R. exoculata* are larger than those of *R. hybisae*
603 (0.08 mm³, Nye et al. 2013), or those of *R. chacei* (0.09 mm³, Ramirez-Llodra et al. 2000). They
604 are also larger than the eggs of *A. muricola* (0.1 mm³, Ramirez-Llodra & Segonzac 2006) and
605 similar in size to the eggs of *M. fortunata* (0.13 mm³, Ramirez-Llodra et al. 2000). *R. exoculata*
606 thus stands among the Alvinocarididae shrimps with the largest eggs, perhaps reflecting a specific
607 strategy of higher parental investment per egg.

608 We observed significant variations in reproductive outputs between *R. exoculata* from TAG
609 and Snake Pit. With lower realized fecundities, perhaps partly due to higher post-spawning losses,
610 and lower egg sizes, the reproductive effort of individual females at TAG indeed appears to be
611 reduced compared to that of females from Snake Pit. The difference in fecundity could be
612 explained by the larger body sizes of brooding females at Snake Pit, and the positive allometric
613 variation of fecundity with female size. Similarly, differences in fecundity between *R. hybisae*
614 females of two vent fields in the Cayman Trough were primarily attributed to the large size
615 differences observed between shrimps of the two fields (Nye et al. 2013). However additional
616 factors, such as food availability, environmental challenges (fluid toxicity, or temperature stress),
617 or variations in fertilization success may also contribute to fecundity differences between both
618 vent fields.

619 The smaller size of the embryos carried by females from TAG compared to those from
620 Snake Pit suggests that the reproductive investment per egg is lower at TAG, further contributing
621 to the lower reproductive output observed at this vent field. Both vent fields are located

622 approximately 310 km apart nearly at the same depth, and regional factors are unlikely to provide
623 environmental heterogeneity that could explain the different investment in reproductive efforts.
624 We thus hypothesize that reproduction differences between the two populations are more likely
625 related with local environmental factors associated with vent emissions. Shrimp tolerance to
626 metallic elements and dissolved gases in vent fluids depends on detoxification processes through
627 metallothioneins, antioxidants (Gonzalez-Rey et al. 2007) and metabolic activities of their
628 symbiotic bacteria (Jan et al. 2014). Higher concentrations in some metallic elements in
629 hydrothermal fluids could force the shrimps to allocate more metabolic energy to detoxification
630 processes, at the expense of reproductive functions. For instance, TAG fluids have higher iron,
631 copper and manganese concentrations than those from Snake Pit (Desbruyères et al. 2000,
632 Schmidt et al. 2008, Charlou et al. 2010), which could explain the lower reproductive output of
633 shrimps at this vent field. However, both bioenergetics and vent processes are complex and these
634 hypotheses must be explored with experimental approaches testing physiological tolerance of the
635 shrimps (e.g. August et al. 2016) in relation with more detailed exploration of local vent chemistry.

636 *Stygiopontius pectinatus* copepods are known to inhabit the cephalothoracic cavity of *R.*
637 *exoculata* (Gollner et al. 2010, Humes 1996, Ivanenko et al. 2006). They belong to the
638 Siphonostomatoida, an order of copepods comprising many cases of parasitic lifestyles.
639 *Stygiopontius* copepods of the Juan de Fuca Ridge graze bacteria in alvinellid tubes (Limen et al.
640 2008) and *S. pectinatus* are believed to feed on microbial communities developing in the
641 cephalothoracic cavity of *R. exoculata* (Gollner et al. 2010, Humes 1996). Here, they appear to
642 also inhabit broods, which could similarly provide shelter and food, since the egg envelopes are
643 colonized by bacteria (Methou et al. 2019). We observed a trend of increasing copepod infestation
644 with brood aging, which also correlates with the development of microbial communities on egg
645 envelopes, but could also simply be the result of cumulative probability of encounter with time.
646 The infestation of Alvinocaridid broods with small meiofaunal organisms has been briefly reported

647 for some brooding females of *A. muricola* in cold seeps of the Gulf of Guinea (Ramirez-Llodra &
648 Segonzac 2006). Small nematodes *Chomodorita* sp. nov. were reported among the eggs of *A.*
649 *muricola*, and were associated with the presence of decaying embryos, with one instance of
650 predatory behavior (the nematode was partly inside the embryo) reported. In *R. exoculata* broods,
651 although a more extended survey would be needed to statistically assess the impact of *S.*
652 *pectinatus* copepods on egg development, our observation did not reflect a negative effect, as
653 higher infestation rates were found at Snake Pit where broods had more numerous, larger and
654 healthier eggs.

655 **4.3. Spawning seasonality in *R. exoculata*?**

656 Despite three decades of sampling at MAR hydrothermal vents, brooding females were
657 very rarely reported in *R. exoculata* (Vereshchaka 1997, Ramirez-Llodra et al. 2000). The only
658 exception until 2014, was the observation of brooding females on structures of the Logatchev vent
659 field in March 2007, and the collection of a few of them (Komai et Segonzac 2008, Gebruck et al.
660 2010, Guri et al. 2012). In January-February 2014, large numbers of ovigerous females were
661 observed crawling among dense shrimp aggregates on sulfide structure walls. In our samples,
662 the overall abundance of ovigerous female was relatively low (7.1% of the sampled females) but
663 this probably reflects the low proportion of females reaching the size of sexual maturity. In fact,
664 while a third of the sexually mature females were ovigerous, many of the two remaining thirds
665 exhibited bright pink well developed gonads suggesting they were nearly ready to spawn
666 (observation not quantified, seen in our samples and on *in situ* video acquisition). This high
667 proportion of ovigerous females among sexually mature ones, consistent across two vent fields
668 310 km apart, and contrasting with all previous collections along the MAR (see 1985-2014
669 collection compilation, Table 4) suggests temporal variations in reproductive activity of *R.*
670 *exoculata* with a possible spawning season in winter.

671

672

673 Table 4. Occurrence of brooding females in *Rimicaris exoculata* samples from different cruises
 674 done on the MAR since 1985.

Site	Jan	Feb	Mar	Apr	May	Jun	Jul	Aug	Sep	Oct	Nov	Dec	No date
Rainbow						1998 ⁸ 2000 ⁶	1997 ⁴ 2002 ¹⁰	2005 ⁸	2005 ¹⁰	1998 ¹⁰			
Broken Spur							1997 ⁴ 2002 ¹⁰		1994 ¹⁰ 1996 ¹⁰	1994 ^{3,10}			1994, 1996 ³
TAG site	2014* 2014*, ¹¹					2002 ¹⁰	1997 ⁴	1985 ¹	1994 ³				
Snake Pit	2014* 2014*, ¹¹					2002 ¹⁰	1997 ⁴	2003 ¹⁰	1994 ³			1995 ⁸	
Logatchev			2007 ^{8,9,11}				2001 ⁸ 1997 ^{4,5}				2004 ⁷ 1998 ¹⁰		
Mephisto						2006 ⁸							

675 ¹Williams & Rona (1986), ²Copley (Copley 1998), ³Vereshchaka (1997), ⁴Shank et al. (1998), 5.
 676 Gebruk et al (2000), ⁶Ramirez-Llodra et al. (2000), ⁷Copley et al (2007), ⁸Komai & Segonzac
 677 (2008), ⁹Gebruk et al. (2010), Guri et al. (2012), ¹⁰Lunina & Vereshchaka (2014), ¹¹Methou et al.
 678 (2019), *this study. Non-direct estimations and too small samples were omitted. Color code:
 679 blue: no brooding female reported; green: 1-2 specimens reported; pink: reported as “many
 680 observed”⁸ and few specimens collected^{8,9}; red: > hundred specimens collected.

681

682 However, previous studies on *R. exoculata* reproduction suggested continuous
 683 reproduction based on asynchronous ovarian development observed among females collected in
 684 summer at the Rainbow vent field (Ramirez et al. 2000) and in autumn at TAG (Ramirez-Llodra
 685 et al. 2000, Copley et al. 2007). Nonetheless, regardless of vent origin, oocyte maximum size was
 686 significantly lower in summer than in autumn (Ramirez-Llodra et al. 2000). In addition, brooding

687 females were extremely rare in both studies (3 specimens totally reported in Ramirez-Llodra et
688 al. 2000, collected in August, September and November). This thus might suggest an extended
689 period of oocyte growth and maturation in summer and autumn, with rare occurrences of early
690 spawning, followed with a winter period during which large proportion of sexually mature females
691 spawn. Such scenario is similar to what is observed in *Alvinocaris stactophila* from cold seeps in
692 the Gulf of Mexico (Copley & Young 2006). *A. stactophila* females exhibit increasing oocyte size
693 throughout the year, with small previtellogenic oocytes in both ovigerous and non-ovigerous
694 females in February-March, larger vitellogenic oocytes in summer with no ovigerous female
695 observed and females with large vitellogenic oocytes and small previtellogenic oocytes in
696 November. In the latter, females with large vitellogenic oocytes are large non-ovigerous females
697 (but ready to spawn), while females with small previtellogenic oocytes were either ovigerous
698 females or small bodied non-ovigerous females (Copley & Young 2006). Such pattern fits a
699 seasonal spawning between November and March in *A. stactophila*, and appears very similar to
700 what is observed in *R. exoculata*. In *R. hybisae* from vents in the Cayman Trough, winter spawning
701 was also suggested (Nye 2013, Nye et al. 2013). In winter, most females had one cohort of small
702 previtellogenic oocytes, while a smaller number of them had a cohort of large vitellogenic oocytes,
703 and only a few had a cohort of intermediate size oocytes. Large and medium size vitellogenic
704 oocytes were observed only in non-ovigerous females, while small size previtellogenic oocytes
705 were observed both in ovigerous and non-ovigerous females, suggesting that while some females
706 had already spawned, others had mature oocytes and would probably spawn shortly (Nye 2013,
707 Nye et al. 2013). Inter-annual comparison of the proportion of brooding females between January
708 and February further support winter spawning in *R. hybisae*, but the effect of spatial population
709 structure and need to expand temporal coverage was recognized (Nye 2013). Although we did
710 not dissect ovaries in our study, we noticed that many non-ovigerous females in the large size
711 classes had conspicuous pink-orange gonads filled with large oocytes (visible through the
712 transparent carapace) indicative of imminent spawning, whereas ovigerous females had shrank

713 whitish gonads, which reflects observations on ovaries of *R. hybisae* during the winter spawning
714 period. As for *R. hybisae*, we need to expend the temporal breath of observations on *R. exoculata*
715 reproduction to better constrain its spawning seasonality and assess its degree of variability.

716 Although reproductive traits of vent species were suggested to be phylogenetically
717 constrained (Tyler & Young 1999), some Alvinocaridid shrimp species also exhibit continuous
718 reproduction. *Alvinocaris muricola* (Ramirez_Llodra & Segonzac 2006) from cold seeps off Congo
719 is one such example, although temporally limited observations of asynchronous ovarian
720 development could also hide seasonal patterns. In *Rimicaris chacei* from the MAR, additional
721 observations could also challenge the current view of continuous reproduction based on time
722 limited observations (Ramirez-Llodra et al. 2000). In contrast, *Mirocaris fortunata* exhibit many
723 ovigerous females throughout the year, which is truly supporting continuous reproduction:
724 Ramirez-Llodra et al. (2000) and Van Dover et al. (1996) reported many brooding females from
725 May, June, July and September at Lucky Strike. We also repeatedly observed *M. fortunata*
726 brooding females during multiple cruises at different periods of the year and different MAR vent
727 fields: August 2013 and April 2015 at Lucky Strike (Momarsat cruise series), and January 2014
728 at Snake Pit (BICOSE cruise). To conclude, Alvinocaridid reproductive strategies are diverse
729 making it difficult to draw general inferences for the family.

730 Vent species are usually believed to reproduce continuously because of the absence of
731 photoperiodic signals at depth coupled with the high and continuous productivity at vents that
732 would obscure variations in surface-derived food supply. Seasonal reproduction in Alvinocaridid
733 shrimps is however not an exception. On the MAR, *Bathymodiolus azoricus* mussels spawn in
734 January (Colaço et al. 2006, Dixon et al. 2006), and this pattern has been related to the variability
735 in the flux of particulate food material that might trigger seasonal reproductive activity (directly in
736 planktotrophic larvae and/or indirectly in adults). On the East Pacific Rise, vent crabs *Bythograea*
737 *thermydron* brood their offsprings in spring (Perovich et al. 2003), again possibly responding to

738 surface production cues. On the other hand, bythograeid crabs from Pacific-Antarctic Ridge vent
739 sites were suggested to have continuous reproduction, suggesting biogeographic effects may
740 induce variations in reproductive rhythms (Hilario et al. 2009). Similarly, the study of reproductive
741 rhythms in other *Rimicaris* shrimps from other locations may provide insights into the mechanisms
742 underlying spawning seasonality in *R. exoculata*.

743 In addition to seasonal reproduction, population size structure suggest discontinuous
744 recruitment in *R. exoculata*, with similar patterns at both vent fields. Polymodal population
745 structure was also noted by Gebruk et al. (1997) and Copley (1998). In addition, Shank et al.
746 (1998) suggested discrete recruitment of juvenile cohorts based on the observation of red
747 juveniles patches within adult swarms. Discontinuous recruitment may further suggest seasonal
748 reproduction, although growth rates and recruitment temporality remain to be assessed before
749 any supported conclusion can be drawn. *R. chacei* in nurseries appear to be recent recruits as
750 they are similar to those captured in nets above vents (Herring & Dixon 1998). It is less clear for
751 *R. exoculata* juveniles observed in active emission habitats, although the abundance of lipids and
752 isotopic signatures suggest they also have recently experienced pelagic life and were feeding
753 (Methou et al. 2020).

754

755 **4.4. Brooding eggs in vent fluids**

756 The remarkable lack of *R. exoculata* brooding females in samples collected at various
757 periods and locations, until this study, had been explained by the hypothesis of a migration of
758 brooding females towards -yet to be found- areas with moderate environmental conditions in order
759 to avoid exposure of their eggs to harmful vent fluids (Ramirez-Llodra et al. 2000). Such protective
760 behavior of brooding mothers is indeed reported in many vent crustaceans. Brooding females of
761 *B. thermydron* crabs on the East Pacific Rise were observed at the periphery of vents whereas

762 males and non-brooding females occupied areas with high fluid venting and temperatures
763 (Perovitch et al. 2003). In Alvinocaridid shrimps, brooding females of *A. stactophila* locate
764 preferentially in areas with more oxygen, and less toxic sulfur compounds than those occupied
765 by males and non-brooding females (Copley & Young 2006). In the amphipod *Ventiella sulfuris*,
766 adults migrate at the periphery of vents to mate and brood (Sheader & Van Dover 2007). In the
767 anomouran crab *Kiwa tyleri*, brooding females also locate at the periphery in less-constraining
768 conditions, whereas males remain close to vent fluid exits (March et al. 2015, Thatje et al. 2015).
769 In *R. exoculata*, however, such scenario had already been challenged by the collection of a few
770 brooding females in dense shrimp aggregates at Logatchev (Guri et al, 2012). Our study further
771 strengthen these observations, brooding females from TAG and Snake Pit being exclusively found
772 in dense shrimp aggregates where temperature records were higher, and never observed
773 (including on video footage) in cold peripheral areas where males scattered.

774 Females of *R. hybisae* also appear to brood their eggs directly within aggregates crawling
775 on chimney walls with no particular protection from vent fluids. Both *R. exoculata* and *R. hybisae*
776 are rare examples of vent species that experience vent fluid exposure during embryonic
777 development. Indeed, many species broadcast their eggs, in which case early development occur
778 in the water column at some distance from the most extreme part of the vent fluid mixing gradient,
779 and brooding species usually provide protective structures for the early development of their
780 offsprings (e.g. Reynolds et al. 2010), or relocate into milder areas.

781 As all developmental stages were observed in broods of *R. exoculata*, including hatching
782 zoea (Hernández-Ávila et al. 2015), females probably remain near vent fluid exits during the whole
783 brooding period until they release their larvae. Exposure of embryos to high temperature may
784 accelerate their development, and shorten the brooding period, while challenging normal
785 development due to heat stress. *In vitro* incubations of embryos removed from brooding females
786 of *Alvinocaris longirostris* and *Shinkaicaris leurokolos*, two Alvinocaridid species inhabiting

787 Okinawa Trough vent sites, showed increased developmental rates with increasing temperature
788 for both species, with optimal growth at temperatures within the range experienced by their adult
789 counterparts *in situ* (Watanabe et al. 2016). These authors showed that embryos of *S. leurokolos*,
790 which lives near fluid exits, developed better between 10 to 20°C, and hatched within 9-12 weeks
791 at 10°C and 3-4 weeks at 20°C. Considering the habitat of brooding females in *R. exoculata*, eggs
792 are likely to hatch rather quickly, perhaps within a month following spawning. A relatively short
793 brooding period in *R. exoculata* is also supported by the lack of macroscopic evidence of
794 exceedingly high mineral deposits in the cephalothoracic cavity of ovigerous females. The short
795 molt cycle of *R. exoculata* (10 days, Corbari et al. 2008) is believed to reset symbiotic communities
796 of the cephalothoracic cavity sufficiently often to avoid impaired symbiosis due to mineral
797 precipitation progressively overgrowing bacteria. In bearing-egg crustaceans, the molting cycle is
798 interrupted during the brooding period until hatching (Correa & Thiel 2003). As females appear to
799 spend the entire brooding period near vent fluids, a large extension of the molting cycle expose
800 them to the risk of seeing their symbiotic bacteria drowned in mineral precipitations. However, as
801 ovigerous females did not appear to suffer excessive mineral load of the cephalothoracic cavity,
802 this observation also advocates for a relatively short brooding period.

803 Brooding within vent fluids may also sustain the bacterial community including symbiotic
804 lineages observed on egg envelopes (Methou et al. 2019), and may thus participate in symbiont
805 transmission to the young shrimp (Hernandez Avila et al. 2015). Another speculative hypothesis
806 would be that mothers could imprint their offspring with vent signature by bathing their eggs within
807 vent fluids during embryonic development, which would later help them returning to vent habitats
808 after dispersal. Such homing process might involve the strongly developed higher brain centers
809 of *R. exoculata* enabling the memory and navigation skills necessary to locate suitable recruitment
810 sites (Machon et al. 2019).

811

812 **4.5. Mating system in *R. exoculata***

813 Spatial segregation of sexes in vent species has been related to reproductive processes
814 needing specific habitats, but it is not known whether such spatial structuration remains
815 throughout the year or is restricted to mating and brooding periods. Segonzac et al. (1993)
816 observed dispersed *R. exoculata* - of unknown sex - at the periphery of Snake Pit in June 1988
817 but more observations are needed to evaluate the temporal correlation between spatial
818 distribution of sexes and reproductive processes. It is however unlikely that individuals resting in
819 peripheral areas stay permanently in such habitats because of their need to supply their symbiotic
820 bacteria with reduced compounds found in vent fluids (Ponsard et al. 2013). Moreover, the highly
821 modified cephalothorax of *R. exoculata* adults hinder movement of the appendages, making any
822 foraging or grazing behavior unlikely (Segonzac et al. 1993, Zbinden & Cambon 2020). Shrimps
823 thus most likely fast when they are in IEH. Peripheral areas also harbor more predators, such as
824 *Maractis rimicarivora* anemones (Fautin & Barber 1999, Copley et al. 2007), or fishes that are
825 excluded from AEH, maybe because of the harsh vent conditions. The migration of *R. exoculata*
826 males to inactive areas is therefore not likely driven by trophic needs or interspecific interactions
827 (increased predatory risk), and given the current knowledge, the hypothesis of a reproductive
828 behavior remains the best explanation. In addition, spermatophores at the gonopores of males
829 were often observed in IEH, but never in males from other habitats (Pradillon, personal
830 observation), suggesting a greater mating readiness in IEH. Mature caridean females molt just
831 before egg extrusion (Bauer 2004), and vent shrimps may migrate briefly towards a milder habitat
832 at this stage of increased vulnerability. Males in IEH would thus have better chances to encounter
833 females just after their prespawning molt and mate successfully. After mating and egg extrusion,
834 females would return to AEH and brood their eggs, while the fate of males remains uncertain.
835 Males will however probably have to return to AEH at some point to fulfill their nutrition needs.

836 Body sizes and male weaponry in crustaceans have been associated with their mating
837 system (Correa & Thiel 2003, Baeza & Thiel 2007, Asakura 2009). In free-living caridean shrimps,
838 larger males are usually observed in mating systems that involve sexual competition for females
839 or precopulatory mate guarding (Bauer 1996, Correa & Thiel 2003, Asakura 2009). In contrast,
840 the lack of sexual competition in “pure searching” mating systems or long-term mate (monogamy
841 or semi-monogamy) is generally associated with similar or smaller-sized males (Correa & Thiel
842 2003, Bauer 1996). Although *R. exoculata* males were on average larger than females, this was
843 due to the large proportions of small females. Considering only sexually mature females, i.e. those
844 females actually involved in the courtship and mating processes, then sex size differences do no
845 longer exist, and females are even slightly larger than males. In other vent carideans such as
846 *Lebbeus virentova*, females tend to be larger than males (Nye & Copley 2014), and a similar trend
847 has been proposed for the cold-seep alvinocaridid *A. muricola*, based on maximum sizes of males
848 and females (Ramirez-Llodra & Segonzac 2006). In *R. hybisae*, males and females exhibit similar
849 sizes (Nye et al. 2013). In addition, in *R. exoculata* there is a lack of sexual dimorphism in
850 secondary characters associated with male competition (e.g. increase in cheliped size and
851 cephalotorax). We thus hypothesize that a pure searching model could better describe the mating
852 system in *R. exoculata*. In this model, males search for receptive females just after their
853 reproductive molt using mostly tactile signals or pheromones (Bauer 1996, 1976). However,
854 looking for tactile and chemical cues within a dense aggregation of congeners close to vent
855 emissions does not seem optimal. A migration to the vent periphery to courtship and mate could
856 increase the chance of male-female recognition and ensure mating success. In his description of
857 the mating behavior of the Hippolytidae *Heptocarpus pictus*, a caridean species exhibiting a pure
858 searching mating model, Bauer (1976) described a suite of events including one called “straddle”
859 when the male clutches the female with his walking legs. We observed *R. exoculata* pairs with
860 similar behavior several times *in situ* (in all habitats, but mainly at the periphery of AEH or in DEH)
861 (Supplementary video 1), which further supports the pure searching mating model in this species.

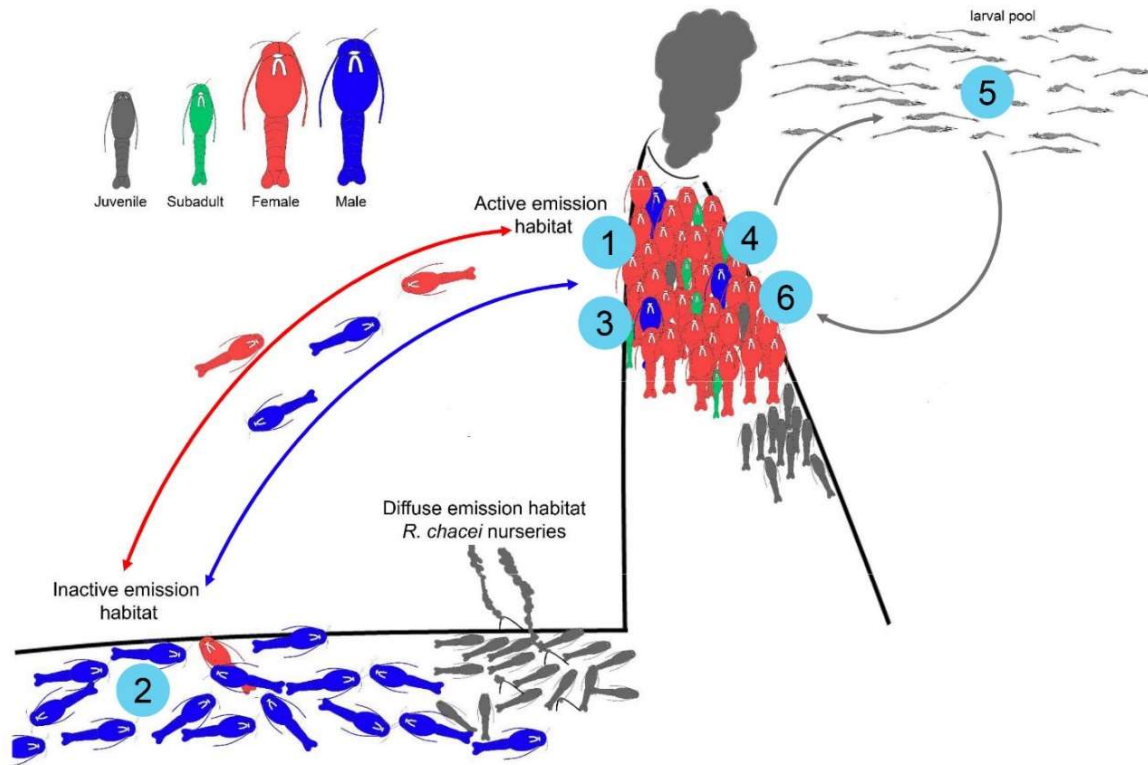
862

863 **4.6. A proposed life cycle scenario for *Rimicaris exoculata***

864 Our examination of population structure and reproductive features across different
865 habitats allows us to draw a hypothetical scenario for the benthic phase of the life cycle of *R.*
866 *exoculata* (Fig. 6). In our observations, juveniles of the species appear to recruit directly in dense
867 aggregations crawling on vent active structures. Visually, they can be distinguished from adults
868 and subadults by their bright red color. Although in this study “nurseries” near diffusion areas
869 belonged to *R. chacei*, juveniles of *R. exoculata* were observed forming gatherings within dense
870 assemblages (Pradillon, personal observation and Shank et al. 1998), perhaps related to small-
871 scale environmental conditions within AEH. Considering all observations gathered so far in this
872 study and others (Shank et al. 1998, Gebruck et al. 2000, Methou et al. 2020), juveniles found in
873 AEH are the smallest observed and identified with confidence for *R. exoculata*. We cannot
874 exclude that we are still lacking an earlier benthic stage, as smaller alvinocaridid juveniles -
875 without species assignment – were caught in bottom trawls (Herring & Dixon 1998). However,
876 isotopic signatures of the smaller benthic stages of *R. exoculata* reported by Methou et al. (2020)
877 are reflecting a pelagic lifestyle and advocate for relatively recent recruitment of those stages.

878 After recruitment, juveniles shift towards a chemosynthetic nutrition, and grow to the
879 subadult and adult stages. While small adults appear to stay within AEH as they grow, mature
880 and reach reproduction size, sexually mature reproductive adults could migrate in less active parts
881 of vent fields for mating. Sexually mature males would move there, reaching a position where they
882 may more “easily” find sexually receptive females that move out of dense aggregations to molt,
883 mate and extrude their eggs. No competitive or guarding behavior was observed in males, and
884 their size broadly similar to the size of females, as well as their lack of weaponry, do not argue
885 for competition. Instead, males appear rather inactive, and exhibit behavior resembling the “pure

886 searching” model where males contact females to find and mate with receptive females. Brooding
887 would occur entirely within vent mixing gradients and last for a few weeks before release of zoea
888 larvae. These larvae would then disperse and mix within bathypelagic waters, developing and
889 feeding for a while on pelagic food items until they reach a large post-larval stage before returning
890 to a benthic and chemosynthetic life style at vents.



Life stage	Habitat	Observation	Reference
1 Growth & maturation	AEH	All stages from juveniles to adults are present, but few males.	This study.
2 Mating & spawning	IEH	Many adult males, rare females, mating behavior observed.	This study.
3 Brooding	AEH	Brooding females with embryos at all stages.	Guri et al. 2012, this study.
4 Larvae release	AEH or closeby	Late broods and hatched females in AEH, zoe larvae collected within a few meters from adult populations.	Hernandez-Avila et al. 2015, this study.
5 Larval dispersal	bathypelagic	Alvinocaridid post-larval stages collected but not identified to the species level.	Herring & Dixon 1998
6 Recruitment	AEH	Most juveniles observed in AEH.	Methou et al. 2020, this study.

891

892 Figure 6. Proposed model of habitat use through the life cycle of the vent alvinocaridid shrimp
 893 *Rimicaris exoculata*

894

895

896 **Credit author statement:**

897 Iván Hernández-Ávila : Conceptualization, methodology, formal analysis, investigation, writing –
898 original draft.

899 Marie-Anne Cambon-Bonavita : Resources, supervision, writing – review & editing.

900 Jozée Sarrazin : Resources, writing – review & editing.

901 Florence Pradillon : Conceptualization, methodology, resource, supervision, writing – review &
902 editing.

903

904

905

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913

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