

1 **Title:** Contrasting patterns of changes in abundance following a bleaching event  
2 between juvenile and adult scleractinian corals  
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## 20 **Key words**

21 coral reefs, climate change, ecology, thermal stress, juvenile corals

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## 23 **Conflict of interest statement**

24 On behalf of all authors, the corresponding author states that there is no conflict of  
25 interest.

## 26 **Abstract**

27 Coral bleaching events have caused extensive mortality on reefs around the world.  
28 Juvenile corals are generally less affected by bleaching than their conspecific adults,  
29 and therefore have the potential to buffer population declines and seed recovery. Here,  
30 we use juvenile and adult abundance data at 20 sites encircling Lizard Island, Great  
31 Barrier Reef, before and after the 2016 bleaching event to quantify: 1) correlates of  
32 changes in juvenile abundance following a bleaching event; 2) differences in  
33 susceptibility to extreme thermal stress between juveniles and adults. Declines in  
34 juvenile abundance were lower at sites closer to the 20m depth contour and higher for  
35 *Acropora* and *Pocillopora* juveniles than for other taxa. Juveniles of *Acropora* and  
36 *Goniastrea* were less susceptible to bleaching than adults, but the opposite was true for  
37 *Pocillopora* spp. and taxa in the family Merulinidae. Our results indicate that the  
38 potential of the juvenile life-stage to act as a buffer during bleaching events is taxon-  
39 dependent.

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## 41 **Introduction**

42 Coral reefs have been facing numerous anthropogenic stressors for decades, leading to a  
43 severe decline in the abundance of corals and associated organisms on reefs (Jackson et  
44 al. 2001). Among those stressors, increasing atmospheric carbon dioxide concentrations  
45 have had a strong negative effect on coral health by causing rapid and prolonged  
46 increases in sea surface temperatures that, in combination with high solar radiation,  
47 disrupt the relationship between the coral host and photosynthetic endosymbiotic algae  
48 (Lesser et al. 1990). Thermal bleaching often results in partial colony mortality or death  
49 (Harriott 1985), but the severity of bleaching depends on many factors. Importantly,  
50 juvenile corals tend to bleach less and survive better than adults (Mumby 1999; Shenkar

51 et al. 2005; Loya et al. 2001), and therefore have the potential to facilitate post-  
52 bleaching recovery.

53         A few explanations for the difference in bleaching susceptibility between  
54 juveniles and adults have been proposed. First, mass transfer to eliminate toxic by-  
55 products occurs more rapidly in small organisms than in large ones (Nakamura and van  
56 Woesik 2001) and in flat rather than branching organisms (Patterson 1992). Juvenile  
57 corals are both small and relatively flat, and mass transfer dynamics might at least partly  
58 explain their lower susceptibility to bleaching. Additionally, differences between  
59 juvenile and adult susceptibilities to bleaching might be partially driven by higher  
60 concentrations of fluorescent proteins in juveniles (Papina et al. 2002), which enhance  
61 resistance to coral bleaching by dissipating excess light energy (Salih et al. 2000).  
62 Moreover, being non-reproductive might allow juveniles to invest more energy into  
63 maintenance to survive thermal stress when compared to adults.

64         Responses to thermal stress can differ widely between corals and environments.  
65 Some coral taxa are more susceptible to bleaching than others (Marshall and Baird  
66 2000; Loya et al. 2001). Structural complexity can also be important because it  
67 increases variation in irradiance (Brakel 1979). High irradiance worsens the effects of  
68 high water temperatures on coral health (Lesser et al. 1990), and therefore colonies in  
69 shaded microhabitats typically bleach less (Hoogenboom et al. 2017; Muir et al. 2017).  
70 High water flow facilitates the removal of toxins produced at high water temperatures  
71 and high irradiance, reducing bleaching-induced mortality (Nakamura and van Woesik  
72 2001) and facilitating post-bleaching recovery (Nakamura et al. 2003). This  
73 experimental work is supported by field work indicating that sites close to deeper water  
74 were more resistant to bleaching on the Great Barrier Reef (GBR) in 2002, presumably

75 due to proximity to cooler water and a greater capacity for water mixing (Done et al.  
76 2003).

77 Warm water temperatures in the austral summer of 2016 led to the most severe  
78 bleaching event on record on the GBR and caused widespread mortality, particularly in  
79 the northern section of the GBR (Hughes et al. 2017). We use environmental data and  
80 juvenile and adult abundance before and after the 2016 bleaching event to answer the  
81 following questions: 1) What are the best environmental predictors of changes in  
82 juvenile abundance after a bleaching event? 2) Per taxon, are adult or juvenile colonies  
83 more resistant to extreme thermal disturbance?

84

## 85 **Methods**

### 86 **Study location and data collection**

87 Data were collected at 20 sites around Lizard Island in the northern GBR (14.6688° S,  
88 145.4594° E) in November in each of 2015 and 2016 (Fig. 1). At each site, five quadrats  
89 of 1m<sup>2</sup> or 0.81m<sup>2</sup> (in 2015 and 2016, respectively) were haphazardly placed and all  
90 juvenile corals within them were counted and identified to genus following Veron  
91 (2000) but updating genus and family following the World Register of Marine Species.  
92 Colonies with a maximum diameter <5cm were considered to be juveniles (*sensu* Bak  
93 and Engel 1979). Fragments were not considered juveniles. Five or six 10m line  
94 intercept transects were laid at each site to record abundances of adult corals (≥5cm  
95 diameter). Abundance was the number of individual colonies that were intercepted by  
96 the transect tape.

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100 **Predictor variables**

101 We tested the extent to which bleaching susceptibility measured by changes in  
102 abundance was predicted by five factors: thermal stress, rugosity, aspect, distance to the  
103 20m depth contour, and taxon (Table A1). We used site-level maximum water  
104 temperature recorded by temperature loggers at the sites from November 2015 to  
105 November 2016 as a proxy for thermal stress (Done et al. 2003). Rugosity was used as a  
106 proxy for structural complexity and was estimated from 3D reconstructions of 130m<sup>2</sup>  
107 areas within each site in 2015 (Pizarro et al. 2017). Rugosity was defined as the ratio of  
108 the site's 3-D surface area to the 2-D area of the plane of best fit (Friedman et al. 2012).  
109 Sites differ in water currents and wind exposure depending on their location around the  
110 island, therefore aspect (NW, NE, SW, SE) was included as a variable to capture these  
111 differences. Sites close to the 20m depth contour (m, log scale) are expected to have a  
112 greater proximity to deep and presumably cooler water and a greater capacity for water  
113 mixing than sites farther away (Done et al. 2003). Finally, taxonomic category was also  
114 included as a variable (Marshall and Baird 2000).

115

116 **Analysis**

117 To test whether juvenile abundance per m<sup>2</sup> decreased in 2016 relative to 2015, we fitted  
118 a linear mixed effects model with year as a fixed effect and site as a random effect  
119 ('lme4' R package; Bates et al. 2015). To examine the best predictors of change in  
120 juvenile abundance (number of juveniles), we calculated the yearly mean abundance (m<sup>-2</sup>  
121 <sup>2</sup>) for each taxonomic category at each site. Genera with fewer than 20 juveniles were  
122 grouped with other genera with fewer than 20 individuals belonging to the same family.  
123 Family groups with fewer than 20 juveniles were eliminated to ensure estimates for  
124 juvenile change were not strongly influenced by rare taxa. We fitted a series of linear

125 models predicting proportional change in juvenile abundance between years for  
126 taxonomic category  $i$  at site  $j$  ( $\log_{10}(\text{abundance } 2016_{ij} / \text{abundance } 2015_{ij})$ ) as a function  
127 of each combination of predictor variables. Our sample size allowed for a maximum of  
128 two predictor variables to be included in each model. We included models with site as a  
129 random effect and without random effects and we calculated the  $R^2$  value for each  
130 model ('r2glmm'; Jaeger et al. 2016).

131 To examine the similarity of changes in abundance post bleaching between  
132 juveniles and adults, we plotted mean proportional change in adults vs. mean  
133 proportional change in juveniles for each taxon, using changes in proportional  
134 abundance as a proxy for mortality. All analyses were performed in R version 3.2 (R  
135 Core Team 2016).

136

## 137 **Results and discussion**

138 Total mean juvenile abundance from did not vary significantly between 2015 and 2016  
139 at the island scale (Table A2) and changes among sites were highly variable (Fig. A1).  
140 Proportional change in juvenile abundance was best explained by distance to the 20m  
141 depth contour and taxonomic category (Table A3; site as a random effect was not  
142 significant:  $p\text{-value}=1$ ).

143 Proportional change in juvenile abundance decreased in 2016 relative to 2015  
144 with increasing distance to the 20m depth contour (Fig. 2-a; Table A4). Sites closer to  
145 the 20m depth contour are closer to deeper and potentially cooler water, as well as  
146 having a higher capacity for these waters to mix due to the greater angle of the reef  
147 slope (Done et al. 2003). On Lizard Island, sites closer to the 20m contour are on the  
148 eastern side of the island and therefore are also more exposed to the south-easterly trade  
149 winds that will stimulate water mixing and movement (Fig. A2). Moreover, high water

150 flow facilitates the mass transfer of toxic by-products of bleaching (Nakamura and van  
151 Woesik 2001), ameliorating the effects of bleaching. In contrast, Hoogenboom et al.  
152 (2017) attributed their finding that bleaching severity was higher in adults at sites that  
153 were closer to open water on Lizard Island to the fact that these sites experience less  
154 temperature variability, a factor known to promote thermal tolerance in adults (Oliver  
155 and Palumbi 2011). These contrasting results between adult and juveniles suggest that  
156 factors associated with bleaching susceptibility differ among life-history stages.

157         Some previous studies have suggested that juveniles are ‘immune’ to the effects  
158 of bleaching (e.g. Mumby 1999; Depczynski et al. 2013). In contrast, our results reveal  
159 an order of magnitude decline in the abundance of *Pocillopora* and *Acropora* (Fig. 2-b)  
160 and lesser, but still significant, declines in *Dipsastraea*, *Favites* and other Merulinidae  
161 (Fig. 2-b). Mumby (1999) conducted his research in Belize, where there are no  
162 *Pocillopora*, and *Acropora* colonies are rare: juveniles of these genera were the most  
163 susceptible to bleaching at Lizard Island. Therefore, the contrasting results are most  
164 likely driven by differences in species composition. Depczynski et al. (2013) used a  
165 different size cut-off for juveniles (less than 10 cm vs. our 5 cm). In addition, their  
166 estimates of mortality were based on the proportion of individual in different categories  
167 a few months post-bleaching (i.e. dead vs. alive). In fact, it is often difficult to detect  
168 dead coral colonies, particularly when they are small; therefore it is possible they  
169 underestimated mortality in the juvenile size class.

170         Overall, the percentage of variation in the proportional change of juvenile  
171 abundance explained by any combination of two or fewer variables was low (Table A2).  
172 More than one cohort of corals corresponds to the juvenile size class in this study.  
173 Juvenile abundance is ultimately increased by recruitment and decreased by juveniles  
174 dying or growing larger than 5cm. While bleaching can affect juvenile survival, many

175 other factors, such as the strength of yearly recruitment and the bleaching-independent  
176 mortality, will also contribute to changes in juvenile abundance.

177 Differences in susceptibility to the 2016 bleaching event between juveniles and  
178 adults differed among taxa (Fig. 3). Total mean abundance of *Cyphastrea*, *Porites*,  
179 *Montipora*, *Favites*, and *Dipsastraea* spp. remained stable in both life-stages. Overall,  
180 adults of *Acropora* and *Goniastrea* spp. were more affected than juveniles (mean  
181 abundance of *Goniastrea* juveniles actually increased; Fig. A3), a finding consistent  
182 with previous studies (Mumby 1999; Loya et al. 2001; Bena and van Woesik 2005;  
183 Depczynski et al. 2013). However, juveniles of *Pocillopora* spp. and the family  
184 Merulinidae had a stronger decline in abundance post-bleaching than adults, indicating  
185 that the decrease in susceptibility as a function of size is taxon-dependent.

186 Demographic processes of juvenile corals have important effects on population  
187 dynamics and are key to population persistence after disturbance (Connell et al. 1997).  
188 Our results show that differences in bleaching susceptibility between juveniles and  
189 adults among taxa need to be considered when predicting changes to reef communities  
190 following episodes of thermal stress.

191

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## References

- Bak RPM, Engel MS (1979) Distribution, abundance and survival of juvenile hermatypic corals (Scleractinia) and the importance of life history strategies in the parent coral community. *Mar Biol* 54:341-352
- Bates D, Mächler M, Bolker B, Walker S (2015) Fitting linear mixed-effects models using *lme4*. *J Stat Softw* 67:1-48
- Bena C, van Woesik R (2004) The impact of two bleaching events on the survival of small coral colonies (Okinawa, Japan). *Bull Mar Sci* 75:115-125
- Brakel WH (1979) Small-scale spatial variation in light available to coral reef benthos: quantum irradiance measurements from a Jamaican reef. *Bull Mar Sci* 29:406-413
- Connell JH, Hughes TP, Wallace CC (1997) A 30-year study of coral abundance, recruitment, and disturbance at several scales in space and time. *Ecol Monogr* 67:461-488
- Depczynski M, Gilmour JP, Ridgway T, Barnes H, Hyeward AJ, Holmes TH, Moore JAY, Radford BT, Thomson DP, Tinkler P, Wilson SK (2013) Bleaching, coral mortality and subsequent survivorship on a West Australian fringing reef. *Coral Reefs* 32:233-238
- Done T, Turak E, Wakeford M, De'ath G, Kininmonth S, Wooldridge S, Berkelmans R, van Oppen M, Mahoney M (2003) Testing bleaching resistance hypotheses for the 2002 Great Barrier Reef bleaching event. Australian Institute of Marine Science, 106 pp
- Friedman A, Pizarro O, Williams SB, Johnson-Roberson M (2012) Multi-scale measures of rugosity, slope and aspect from benthic stereo image reconstructions. *PloS one* 7:e50440.
- Harriott VJ (1985) Mortality rates of scleractinian corals before and during a mass bleaching event. *Mar Ecol Prog Ser* 21:81-88
- Hoogenboom MO, Frank GE, Chase TJ, Jurriaans S, Álvarez-Noriega M, Peterson K, Critchell K, Berry KLE, Nicolet KJ, Ramsby B, Paley AS (2017) Environmental drivers of variation in bleaching severity of *Acropora* species during an extreme thermal anomaly. *Front Mar Sci* 4:376
- Hughes TP, Kerry JT, Álvarez-Noriega M, Álvarez-Romero JG, Anderson KD, Baird AH, Babcock RC, Beger M, Bellwood DR, Berkelmans R, Bridge TC, Butler IR, Byrne M, Cantin NE, Comeau S, Connolly SR, Cumming GS, Dalton SJ, Diaz-Pulido G, Eakin CM, Figueira WF, Gilmour JP, Harrison HB, Heron SF, Hoey AS, Hobbs J-PA, Hoogenboom MO, Kennedy EV, Kuo C-Y, Lough JM, Lowe RJ, Liu G, McCulloch MT, Malcolm HA, McWilliam MJ, Pandolfi JM, Pears RJ, Pratchett MS, Schoepf V, Simpson T, Skirving WJ, Sommer B, Torda G, Wachenfeld DR, Willis BL, Wilson SK (2017) Global warming and recurrent mass bleaching of corals. *Nature* 543:373-377
- Jackson JBC, Kirby MX, Berger WH, Bjorndal KA, Botsford LW, Bourque BJ, Bradbury RH, Cooke R, Erlandson J, Estes JA, Hughes TP, Kidwell S, Lange CB, Lenihan HS, Pandolfi JM, Petterson CH, Steneck RS, Tegner MJ, Warner RR (2001) Historical overfishing and the recent collapse of coastal ecosystems. *Science* 293:629-638
- Jaeger BC, Edwards LJ, Das K, Sen PK (2016) An  $R^2$  statistic for fixed effects in the generalized linear mixed model. *J Appl Stat* 44:1086-1105
- Lesser MP, Stochaj WR, Tapley DW, Shick JM (1990) Bleaching in coral reef anthozoans: effects of irradiance, ultraviolet radiation, and temperature on the activities of protective enzymes against active oxygen. *Coral Reefs* 8:225-232

252 Loya Y, Sakai K, Yamazoto K, Nakano Y, Sambali H, van Woesik R (2001) Coral  
253 bleaching: the winners and the losers. *Ecol Lett* 4:122-131  
254 Marshall PA, Baird AH (2000) Bleaching of corals on the Great Barrier Reef:  
255 differential susceptibilities among taxa. *Coral Reefs* 19:155-163  
256 Muir PR, Marshall PA, Abdulla A, Aguirre JD (2017) Species identity and depth  
257 predict bleaching severity in reef-building corals: shall the deep inherit the  
258 reef? *Proc R Soc B Biol Sci* 284: 20171551  
259 Mumby PJ (1999) Bleaching and hurricane disturbances to populations of coral recruits  
260 in Belize. *Mar Ecol Prog Ser* 190:27-35  
261 Nakamura T, van Woesik R (2001) Water-flow rates and passive diffusion partially  
262 explain differential survival of corals during the 1998 bleaching event. *Mar Ecol*  
263 *Prog Ser* 212:301-304  
264 Nakamura T, Yamasaki H, van Woesik R (2003) Water flow facilitates recovery from  
265 bleaching in the coral *Stylophora pistillata*. *Mar Ecol Prog Ser* 256:287-291  
266 Oliver TA, Palumbi SR (2011) Do fluctuating temperature environments elevate coral  
267 thermal tolerance? *Coral Reefs* 30:429-440  
268 Papina M, Sakihama Y, Bena C, van Woesik R, Yamasaki H (2002) Separation of  
269 highly fluorescent proteins by SDS-PAGE in Acroporidae corals. *Comp*  
270 *Biochem Phys* 131:767-774  
271 Patterson MR (1992) A mass transfer explanation of metabolic scaling relations in some  
272 aquatic invertebrates and algae. *Science* 255:1421-1423  
273 Pizarro O, Friedman A, Bryson M, Williams SB, Madin JS (2017) A simple, fast, and  
274 repeatable survey method for underwater visual 3D benthic mapping and  
275 monitoring. *Ecol Evol* 7:1770-1782  
276 R Core Team (2016) R: A language and environment for statistical computing. R  
277 Foundation for Statistical Computing, Vienna, Austria  
278 Salih A, Larkum A, Cox G, K  hl M, Hoegh-Guldberg O (2000) Fluorescent pigments  
279 in corals are photoprotective. *Nature* 408:850-853  
280 Shenkar N, Fine M, Loya Y (2005) Size matters: bleaching dynamics of the coral  
281 *Oculina patagonica*. *Mar Ecol Prog Ser* 294:181-188  
282 Veron JEN (2000) Corals of the world. Australian Institute of Marine Science,  
283 Townsville  
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292 **Figure Legends**

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294 Figure 1. Map of Lizard Island showing the sites that were surveyed. Dotted lines show  
295 the 20m depth contour.

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297 Figure 2. Panel a- proportional change in juvenile abundance  
298 ( $\log_{10}[\text{abundance}_{2016}/\text{abundance}_{2015}]$ ) as a function of distance to the 20m depth contour  
299 (m). The black line shows the predicted proportional change in juvenile abundance, and  
300 the grey ribbon shows 95% confidence intervals. Panel b- proportional change in  
301 juvenile abundance ( $\log_{10}[\text{abundance}_{2016}/\text{abundance}_{2015}]$ ) for each taxonomic category.  
302 Error bars show standard errors. Grey dots represent observed data, with the dot size  
303 showing the number of observations.

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306 Figure 3. Proportional change in adult abundance versus proportional change in juvenile  
307 abundance (log-log scale) between 2015 and 2016 for each taxonomic category. Line-  
308 ranges indicate standard errors. The dotted vertical and horizontal lines indicate no  
309 proportional change in adult and juvenile abundance, respectively, between years. The  
310 dashed line indicates values for which the proportional change in adult abundance is  
311 equal to the proportional change in juvenile abundance.

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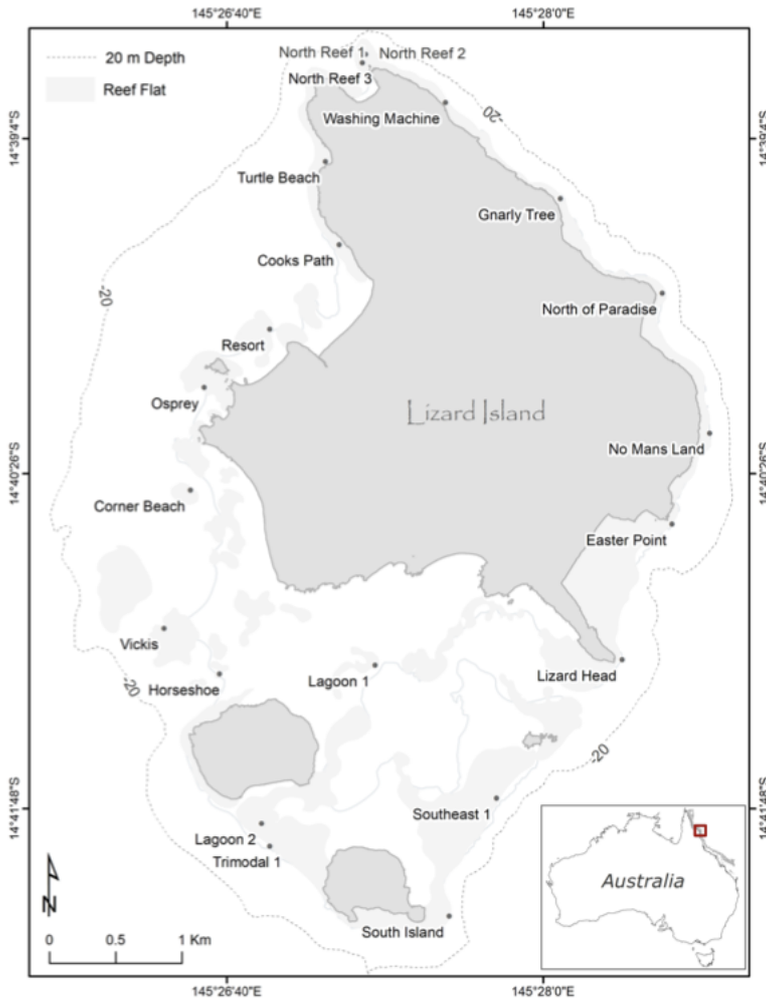
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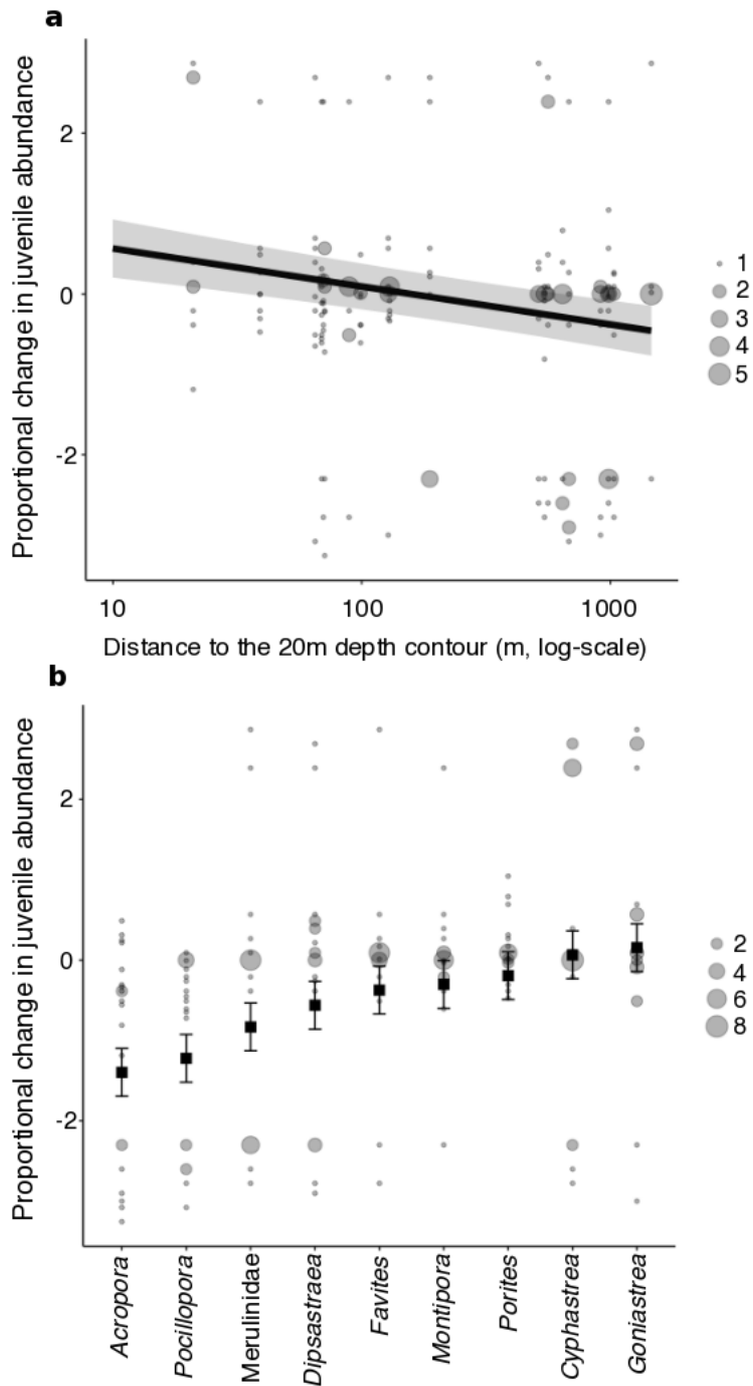
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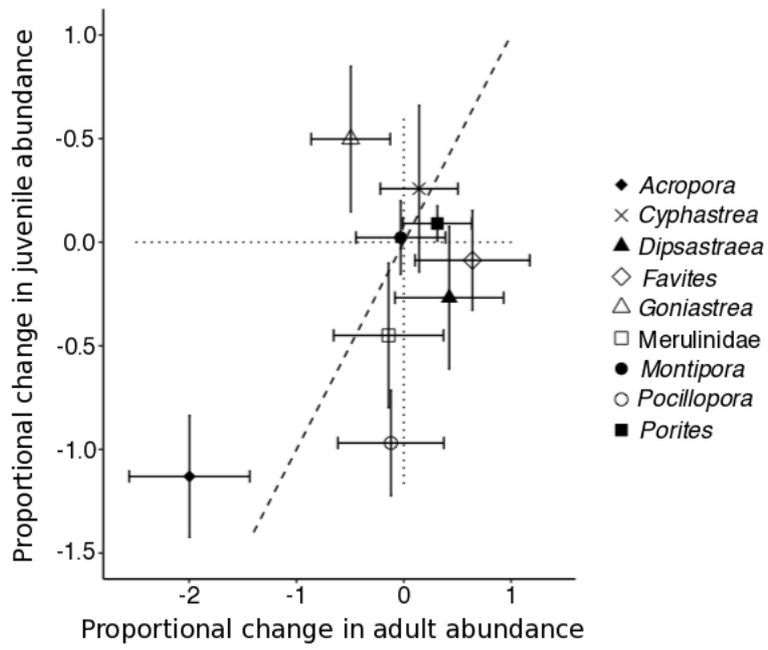
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319 Figure 1



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321 Figure 2



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323 Figure 3