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Insights into the morphology of symbiotic shrimp eyes (Crustacea, Decapoda, Pontoniinae); the effects of habitat demands

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Morphometric differences in the optical morphology of symbiotic palaemonid shrimps can be observed among species symbiotic with different host organisms. Discriminant functional analysis revealed three distinct groups within the species examined. Of these, bivalve symbionts appear to have an eye design that is solely unique to this host-symbiont grouping, a design that spans across multiple genera of phylogenetically unrelated animals. Although some taxonomic effects may be evident, this does not explain the difference and similarities in eye morphology that are seen within these shrimps. Therefore evolutionary pressures from their host environments are having an impact on the optical morphology of eyes however, as indicated by host-hopping events there ecological adaptations occur post host invasion.

| 1 | Insights into the morphology of symbiotic shrimp eyes (Crustacea, Decapoda, Palaemonidae); the |
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8 Abstract

9 Morphometric differences in the optical morphology of symbiotic palaemonid shrimps can be observed among species symbiotic with different host organisms. Discriminant functional 10 11 analysis revealed three distinct groups within the species examined. Of these, bivalve symbionts 12 appear to have an eye design that is solely unique to this host-symbiont grouping, a design that 13 spans across multiple genera of phylogenetically unrelated animals. Although some taxonomic 14 effects may be evident, this does not explain the difference and similarities in eye morphology 15 that are seen within these shrimps. Therefore evolutionary pressures from their host environments 16 are having an impact on the optical morphology of eyes however, as indicated by host-hopping 17 events there ecological adaptations occur post host invasion.

18 **1.** Introduction

19 Symbiotic palaemonid shrimps are widespread and abundant in Indo-West Pacific reefal habitats, 20 characterised by their affinity to form associations with a wide range of taxa. Until recently these 21 shrimps were in the subfamily Pontoniinae. However in a recent phylogenetic study by De Grave 22 et al., (2015) this subfamily was synonymised with the family Palaemonidae, as were the related 23 families Gnathophyllidae and Hymenoceridae. For the purposes of this investigation and 24 throughout the remainder of this paper, we will refer to this group of shrimps as "pontoniine 25 shrimps to avoid any systematic ambiguity. Members of the previously separate families 26 Gnathophyllidae and Hymenoceridae were not included in the present analysis. Within the 27 pontoniine shrimps, an estimated 60-70% (De Grave, 2001) are known to form associations with 28 corals, sponges, ascidians, gorgonians, and so on. However this is likely to be an underestimate 29 as the host association remains unknown for several species, but is inferred to be symbiotic due to 30 their morphological similarity to other species. Pontoniine shrimps occur in a wider variety of 31 tropical and subtropical habitats, and are known from deeper water, down to about 2000 m 32 (Bruce, 2011). However, their highest species richness is on tropical coral reefs, down to about 33 100 m. The most recent catalogue (De Grave & Fransen, 2011) lists 602 species, but numerous 34 species have been described since then. 35 The traditional view of these shrimps as symbionts, has recently been challenged for a number of species dwelling in sponges, where diet studies revealed them to be parasites as their stomachs 36 37 only contained host tissue and spicules (Ďuriš et al., 2011). At present it is not known how 38 widespread parasitism is in the group, and we thus refer to them as associates, inferring no

39 trophic interaction with the host.

40 Morphological adaptation to an associated mode of life has been extensively noted in the 41 taxonomic literature for pontoniine shrimps. Such adaptations include modified pereiopods 42 (Bruce, 1977; Patton, 1994) in addition to extensive modifications in general body plan and 43 mouthparts (Bruce, 1966; Duriš et al., 2011). Additionally, a range of ecologies are recognised, 44 ranging from internally dwelling in small sized hosts like ascidians (e.g. species of the genus 45 *Periclimenaeus*) to fish cleaning species, dwelling on anemones (e.g. *Ancylomenes* spp.). Despite 46 this wealth of morphological and ecological disparity, few studies have been done linking 47 morphological disparity with ecological constraints. A recent exception to this is the study by 48 Dobson et al (2014) which examined gross eye morphology across four, broad, lifestyle

49 categories: ectosymbionts, bivalve endosymbionts, non-bivalve endosymbionts and free-living. 50 Their results clearly demonstrated considerable differences in superficial optical parameters 51 across various lifestyles. In many decapods, vision is thought to be an important feature of their 52 morphology with variations in morphology and structure reflecting ecological habitat demands 53 (Johnson, Shelton and Gaten, 2000). Differences in eye size, facet size and interommatidial angle 54 have been observed in many marine species occupying different depths (Gaten, Shelton, and 55 Herring, 1992; Johnson et al., 2000). Eve parameter (EP) has been used by a number of 56 researchers as a measure of determining the equipoise between sensitivity and resolution of 57 different organisms (Snyder, 1979; Stavenga & Hardie, 1989; Kawada et al., 2006). For 58 organisms occupying well-lit habitats EPs of between 0.45 and 1 rad-um have been recorded, 1-2 59 for crepuscular and 2-3 for nocturnal species (Kawada et al., 2006), however these values many vary in aquatic organisms due to the different refraction index of water. Pontoniine shrimps are 60 61 ideal study organisms for the relationship between eye morphology, vision and habitat demands, given their predilection for forming associations with a wide range of taxa. 62 63 The current study builds upon this previous work, by focussing on and contrasting across actual 64 host identities using a multivariate analytical framework and thus aims to further unravel

65 potential differences in gross optical morphology of pontoniine shrimps.

66 2. Methods

Optical characteristics of 96 species from 40 genera were examined from collections at the 67 68 Oxford University Museum of Natural History. A copy of the dataset used in this paper can be 69 accessed in the Supplemental Information. The work described in this paper was reviewed and approved by the Department of Biological Sciences, Faculty of Sciences ethics committee 70 71 approval number U053. To understand differences in eye morphology between host categories, 72 each species was classed into host-symbiont predefined groupings based on their most common 73 host associations (Bruce, 1994); i.e. Actiniaria, Ascidiacea, Asteroidea, Bivalvia, Crinoidea, 74 Echinoidea, Gorgonacea, Hydrozoa, Ophiuroidea, Porifera and Scleractinia or considered to be 75 free-living. For all species, eye span (ES), diameter at the base of the eyestalk (DBES), facet 76 diameter (FD) and eye diameter (ED) were measured using a dissecting microscope fitted with an 77 ocular micrometer. To reduce scaling effects ES, DBES and ED were standardised by post orbital 78 carapace length, whilst FD was standardised by eye diameter. A composite variable, ES-DBES 79 (eye span minus diameter at base of eyestalk), was also formulated to provide an indication of

- 80 eye mobility, the greater mobility of the eyes the larger the value. In addition to the variables
- 81 measured, eye parameter (EP) was calculated as an outcome of facet diameter (μ m) (FD) and
- 82 interommatidial angle ($\Delta \varphi$ in radians) using Snyder (1979) equation (Equation 1).

83 Equation 1. $EP = FD \Delta \varphi$

Interommatidial angle in radians, used in the calculation of EP, was estimated using an adaptation
of Stavenga's (2003) formula (Equation 2).

86 Equation 2.
$$\Delta \varphi = 2 \left(\frac{FD}{ED} \right)$$

The presence or absence of the nebenauge (see Dobson *et al.*, 2014) was also noted and when
present the relative size was expressed after standardisation by eye diameter (ED). Our
terminology follows Johnson *et al.*, 2015 who utilised nebenauge for the structure previously

90 referred to under several names.

91 Eye Parameter (EP) and standardised nebenauge size was compared between hosts using a

92 Kruskal Wallis test in the Statistical Software Package R 3.0.2 as this allowed for *Post Hoc*

comparisons (R Core Team, 2013), whilst Eye Diameter (ED) was analysed by the means of anANOVA.

95 Subsequently, the dataset was analysed with Discriminant Function Analysis (DFA), also known

96 as Multiple Discriminant Analysis (MDA) or Canonical Variate Analysis (CVA). DFA extracts

97 linear combinations of variables (known as roots) which maximise differences amongst a priori

98 defined groups, in this case host categories, with the percentage correctly classified providing a

- 99 goodness of fit measure, akin to more traditional P values.
- 100 As DFA requires the number of predictor variables to be fewer than the sample size of the
- 101 smallest group, a number of host-categories could not be included in the analysis, namely
- 102 Echinoidea, Hydrozoa, Ophiuroidea and Asteroidea, all of which are relatively infrequently
- 103 inhabited by pontoniine shrimp. Outliers were identified using within host category linear least-
- 104 squares regression analysis, using post-orbital carapace length as the independent variable.

- 105 Individual outliers were corrected by re-measurement (where possible), and only excluded from
- 106 the final dataset if their values still exceeded 3 standard deviation in residual plots. The final
- 107 dataset analysed with DFA thus comprised of 83 species, across 7 host categories, as well as free-
- 108 living taxa. Host categories herein analysed, comprise of Actiniaria (9 shrimp species),
- 109 Ascidiacea (7), Bivalvia (12), Crinoidea (8), Gorgonacea (7), Porifera (14) and Scleractinia (13).
- 110 Thirteen micro-predatory species, which are currently considered not to be host associated, i.e.
- 111 free-living were also included in the analysis, a combination of species living on coral reefs and
- 112 in seagrass beds.
- For consistency, statistical analysis of eye size, Eye Parameter and nebenauge was carried out onthe reduced dataset.
- 115 Prior to DFA, proportions were arcsine-transformed to meet the assumptions for statistical
- analysis of normality and homogeneity (Zuur, Ieno and Elphick, 2010). All DFA analysis was
- 117 performed in SPSS 18. In all DFA analysis, all variables were entered simultaneously, with the
- 118 contribution of each variable assessed on the basis of discriminant loadings (structure
- 119 correlations, rather than discriminant coefficients, as those are considered more valid when
- 120 interpreting the relative contributions of each variable).

121 **3. Results**

122 **3.1** Eye size, Eye Parameter and nebenauge presence

- 123 Across all species examined, mean relative ED (Fig. 1) ranged from 0.09 to 0.27, with
- significantly smaller eyes occurring in bivalve associated species (ANOVA, $F_{7,75} = 9.26$, P<0.001,
- 125 Tukey P = 0.05). Although the analysis deemed none of the remaining differences to be
- 126 statistically significant, ascidian $(\overline{x}=0.19, SD \pm 0.06)$ and sponge symbionts
- 127 $(\bar{x}=0.19, SD \pm 0.06)$ were also found to possess some of the smallest relative EDs whilst
- 128 gorgonian symbionts ($\bar{x}=0.28, SD\pm0.11$) and free-living shrimps ($\bar{x}=0.26, SD\pm0.06$) had
- 129 the largest relative EDs.

130 Eye parameter (EP) (Fig. 2) ranged from 0.44 - 8.06 rad- μ m, with a significantly larger EP found 131 in ascidian, bivalve and sponge associates (Kruskal Wallis, H (adjusted for ties) = 43.62, df = 7, P < 0.001, Post hoc pairwise comparisons P = 0.05). The smallest EP values were found in 132 133 associates of crinoid, gorgonians and in free-living shrimps. Associates of sea anemones and 134 corals were not significantly different to any other host category in terms of EP (Fig. 2), whilst 135 the widest range of values is present in sponge associates. Although not statistically considered as 136 outliers in within-host category regression analysis, three species exhibited an aberrant EP, all of 137 the genus Pontonia. Pontonia panamica an ascidian commensal has the largest EP in the dataset 138 (EP = 7.45), whilst *P. mexicana* and *P. pinnophylax* exhibited considerable larger values than

139 other species associated with bivalves.

140 A significant association was found between the presence/absence of the nebenauge and host

141 category (Chi-squared test, $\chi^2 = 24.777$, df = 7, P<0.001). High absence rates of the nebenauge

142 were observed among ascidian, bivalve and poriferan symbionts (Fig. 3), whilst it is prevalent in

143 sea anemone associates and free-living shrimps. However, the relative size is not different across

host categories (Kruskal Wallis test, H = 8.93, df = 6, P = 0.178), with ascidians excluded as only

145 one species, *Periclimenaeus hecate*, had a nebenaugen.

146 **3.2** Multivariate analysis

147 Discriminant function analysis revealed only two significant roots (Table 1), which cumulatively

148 explain 94.6% of total variance. Examination of the structure matrix (Table 2) revealed that three

149 variables were highly loaded on to the first root (EP, FD, ED), whilst a fourth variable (ES-

150 DBES) displayed greatest loading on the second function.

A classification matrix indicates that overall 50.6% of shrimp species were correctly classified in respect to their priori defined groups (host classification) (Table 3), but with significant variation

153 as to within-group classification. Bivalve associates were 100.0% correctly classified, with a high

154 number also correctly classified for sponge associates (78.6%). Over half of the free-living

155 species (61.5%) were correctly classified to their priori group, with other species classified as sea

anemone, crinoid and coral associates. Gorgonian associates correctly classified in 42.9% of

157 cases, with misclassified taxa allied to free-living, coral and crinoid associates. Coral associates

158 correctly classified in 38.5% of cases with species misclassifying as associates of sponges, sea

anemones, crinoids and free-living species. Sea anemone and crinoid associates were only 22.2
and 25.0% correctly classified. All ascidian symbionts were found to misclassify, with 71.4% of
them misclassified as sponge associates.

When comparing the relative position of the centroids for each host category (Fig. 4) it is obvious, that the eyes of ascidian and sponge associated species are very similar to each other, as are the eyes of crinoid and coral associates, both of which also group with the free-living species. Although broadly similar to the latter grouping, the eyes of gorgonian and sea anemone associates are somewhat divergent as well as divergent to each other, as evidenced by the position of their centroids. Bivalve associates clearly occupy an isolated position, relative to the other host categories.

169 When plotting only the ascidian associates in the DFA analysis (Fig. 5), a divergent position of *P*.

170 *panamica* is evident, whilst the other taxa form a loose grouping. The positions of sponge

171 associates (Fig. 6) reveal two distinct, but loose groupings, as well as a divergent species,

172 Thaumastocaris streptopus. Membership of either of the two groups does not appear influenced

by phylogeny, as either group contains species belonging to the genera *Typton* and

174 Periclimenaeus. The positions of the individual bivalve associates (Fig. 7) reveals a relatively

175 tight grouping, but with an isolated position occupied by Conchodytes nipponensis. The

176 positions of individual crinoid associates (Fig. 8) are rather scattered, but with a very isolated

177 position for Laomenes nudirostris. A similar scattered pattern is observed for the coral associates

178 (Fig. 9) and the free-living species (Fig. 10). Gorgonian associates also demonstrate this pattern

179 (Fig. 11), but with a significant, isolated position for *Pontonides loloata*. A similar pattern is

180 observed for sea anemone associates (Fig. 12), with an isolated position for Periclimenes

181 scriptus.

182 **4. Discussion**

183 Multivariate analysis clearly reveals that three distinct eye types are present in pontoniine

184 shrimps, with bivalve associates comprising a type on their own. Sponge and ascidian associates

185 have remarkably similar eyes, to the point that the majority of ascidian associates were

186 misclassified as sponge associates in the analysis. A third eye type is present in a range of

- 187 ectosymbiotic taxa, associated with sea anemones, gorgonians, corals, crinoids, as well as free-188 living species.
- 189 An examination of the structure loadings reveals that along the first root, both facet diameter
- 190 (FD) and Eye Parameter (EP) increases, but with a concomitant decrease in eye diameter (ED),
- 191 whilst along the second root eye mobility (as measured by ES-DBES) decreases. Broadly
- 192 speaking, the ectosymbiotic and free-living taxa thus have smaller facet diameters, a lower EP
- and bigger eyes, than their endosymbiotic counterparts in bivalves, sponges and ascidians.
- 194 Equally, bivalve associates display more mobile eyes than ascidian and sponge associates, but
- 195 with roughly similar facet diameter and EP. It should be noted that the relative eye size of
- 196 bivalve associates is significantly smaller than all other host groupings, this may be as a result of
- 197 their comparably larger body sizes (e.g. mean average 6.9 mm CL versus 3.0 mm CL for
- 198 Actiniaria, 2.5 mm CL for Porifera and 1.34 mm CL for Gorgonacea symbionts).

199 Within deep sea caridean species the nebenauge has been suggested to have an important role in

200 diurnal migrations (Johnson *et al.*, 2015). The concept that orientation to light is aided by the

201 presence of the nebenauge is further supported by these results with it being highly abundant

202 within sea anemone, crinoid, free-living and coral associates. However for bivalve, ascidian and

203 sponge associates both diurnal migrations and orientation to light would be of little significance

204 for species with an endosymbiotic mode of life.

205 This result is not surprising, given the clear relationship between gross eye morphology of

206 pontoniine shrimps and life style already demonstrated in Dobson et al. (2014). Therein, based on

207 a range of optical parameters, the eyes of free-living and ectosymbiotic species were found to be

208 very similar, and clearly different from both types of endosymbiotic species considered, bivalves

209 and non-bivalve associates. Further, bivalve endosymbionts exhibited an intermediary group

210 between free-living/ectosymbionts and non-bivalve endosymbionts, potentially linked to their

211 presumed more active lifestyle, with bivalve associated documented to move hosts in search of a

212 mate (Baeza et al., 2011).

213 Whilst the relationships between optical parameters and lifestyle in Dobson et al. (2014) appears

- 214 clear-cut and supported by the present analysis, by including actual host identity, rather than
- 215 lifestyle in the current analysis, a number of surprising findings emerge.

216 The eyes of ascidian associated species emerges as being remarkable similar to the eyes of 217 sponge associated species, to the point that the majority of a priori classified species in this group 218 were misclassified as sponge eves by the multivariate analysis. This is herein interpreted being 219 likely a significant signal of phylogenetic constraint, as four out of the seven species in this host 220 category belong to a primarily sponge dwelling genus, Periclimenaeus (see below) with generally 221 conservative eye morphology, potentially indicative of recent host switching event(s). Two 222 further species in this host category, phylogenetically unrelated to Periclimenaeus, Dactylonia 223 okai and Odontonia katoi are thought to be closely related species (Fransen, 2002), but with 224 significantly different gross eye morphology. Dactylonia okai possesses stout triangular shaped 225 eyes, whereas the eyes of *Odontonia* species are small and hemispherical (Fransen, 2002). 226 Whilst D. okai and O. katoi are found living within large solitary ascidians, species of 227 Periclimenaeus are found living within both ascidians and sponges. Species such as 228 Periclimenaeus orbitocarinatus and Periclimenaeus ascidiarum live in association with 229 compound ascidians that are structurally similar in morphology to the canals of sponges occupied 230 by, for example, Periclimenaeus maxillulidens. The structural similarity in hosts between the 231 symbionts of compound ascidians and sponges could be a plausibly explanation for the high 232 misclassification of ascidian symbionts to sponges. Two species were misclassified as either a sea 233 anemone or bivalve associate. Although DFA does not provide information on individual 234 classified species, it is evident from Fig. 5 that P. panamica is the species misclassified as a 235 bivalve associate. The genus Pontonia comprises of 11 species (De Grave & Fransen, 2011) and 236 is morphologically very conservative. Although the host for one species, P. longispina, is not 237 known, the majority of species associate with bivalves in the families Pinnidae and Pteriidae, 238 whilst one poorly known species *P. chimaera*, is thought to be an associate of large gastropods of 239 the genus Strombus. Pontonia panamica is the only species to associate with ascidians, the 240 solitary species Ascidia interrupta in the eastern Pacific. Although Marin and Anker (2008) 241 speculate that a host switch to ascidians occurred early on in the evolutionary history of this genus, the retention of essentially a "bivalve" eye is perhaps indicative of a more recent host 242 243 switching event. However, on balance the differences in eve morphology between the 244 phylogenetically not related genera herein analysed as ascidian associates suggests that despite 245 occurring in a similar host environment, their enclosure inside ascidians has not provided 246 pressure on their eyes to become optically similar. As to whether this lack of overall evolutionary 247 pressure is imparted by distinctive host morphologies (compound, solitary) or habitats (intertidal,

subtidal) or indeed is determined by differential behavioural attributes (social biology) of theassociates themselves remains unclear.

250 Notwithstanding their close similarity to ascidian associate eyes, the eyes of sponge associated 251 species appear to be quite uniform, with the majority being correctly classified in their a priori 252 defined host group, but seemingly forming two distinct subgroups in the analysis, in addition to 253 the outlying *T. streptopus*. We infer here that the classification into two subgroups is putatively 254 related to host morphologies, as sponge species exhibit a discrete and distinct range of canal 255 sizes. Space partitioning, as well as individual host selection is indeed known to play a 256 significant role in the sponge-dwelling gambarelloides group of Synalpheus (Duffy, 1992; 257 Hultgren and Duffy, 2010; 2012). The speculation that canal sizes of the host may play a 258 significant role in optical acuity of pontoniine species, can however not be substantiated, as the 259 host range of most species remains unknown, with even the identity of many hosts simply not 260 being known. For instance, for many species of Periclimenaeus, a primarily sponge associated 261 genus, the hosts are not known (Bruce, 2006). Of particular interest are the three ectosymbiotic 262 species included in this primarily endosymbiotic group in the present analysis, T. streptopus, 263 Periclimenes harringtoni and Periclimenes incertus. Thaumastocaris streptopus is an Indo-264 Pacific species, which dwells in the central atrium of vase-shaped sponges like Siphonochalina 265 and Callyspongia (see Bruce, 1994). Based on the present suite of optic parameters, this species 266 does not cluster with the rest of the sponge associates. Although Duriš et al. (2011) consider the 267 species to be parasitic, in common with several other sponge associates, the isolated position of 268 the species in the present analysis, combined with their asymmetrical first pereiopods and a 269 segmented carpus (both unique within the family) is indicative perhaps of a different behavioural 270 niche. The Indo-Pacific, *P. incertus* dwells on the outside of a variety of sponges, and clusters 271 reasonably close to the other sponge associates in the present analysis, potentially indicative of 272 similar relationship to the host, if external. The Caribbean P. harringtoni dwells in the atria of 273 Neofibularia nolitangere and based on the optical parameters studied herein, appears to have an 274 eye structure very similar to that of endosymbiotic species, potentially an example of habitat 275 driven adaptation, despite the significant difference in position on the host.

The sea anemone associates included in the present analysis, fall into four ecological/systematic groups, *Ancylomenes* and three different species groups of *Periclimenes*. *Ancylomenes* species are on the whole considered to be fish cleaners, who only utilise the sea anemone as an advertisement

279 for their services to client fish (Huebner & Chadwick, 2012). It should be noted that this is 280 potentially a generalisation, as direct observation of fish cleaning behaviour is not available for 281 all species, with this information lacking for one species herein included A. tosaensis. 282 Periclimenes yucatanicus and Periclimenes rathbunae are active large bodied species, associated 283 with a variety of sea anemones in the Caribbean. Fish cleaning has not been observed for either 284 species, with Limbaugh et al., (1961) considering P. yucatanicus a fish-cleaning mimic. 285 Periclimenes ornatus and P. inornatus belong to the same species complex, and are smaller 286 bodied species which hide in between the tentacles of a variety of Indo-Pacific sea anemones. 287 Finally, P. scriptus, a Mediterranean and subtropical Northeast Atlantic species which is not 288 phylogenetically closely related to the other two groups, is an active species, associated with long 289 tentacle sea anemones, with no known fish cleaning behaviour. With the exception of P. scriptus 290 (see below) these species exhibit a scattered grouping in the DFA analysis, and as a group have a 291 low percentage correctly classified, at 22%. It thus appears that despite their broad ecological 292 niche similarity as sea anemone associates, insufficient convergent pressure on their optical

293 parameters is noted, indicative of differential usage of their eyes.

294 In contrast to sea anemone associates, coral associates exhibit a reasonable level of correctly 295 classified in the DFA analysis, at 38.5%, despite the large variety of host morphotypes involved 296 in this association. Several species Coralliocaris spp., Harpilius spp. and, Harpiliopsis spp. are 297 associated with branching corals of the families Pocilloporidae and Acroporidae. Other species 298 in this group are associated with corals which extend their polyps during the day, either short 299 polyps (e.g. Hamopontonia corallicola on Goniopora) or long polyp forms, such as Cuapetes 300 kororensis on Heliofungia actiniformis. Morphologically heavily modified taxa are also present 301 in this group, such as the laterally flattened *Ischnopontonia lophos* which moves between the 302 corallites of Galaxea. It thus appears that the habitat and/or behaviour in the case of coral 303 associates is a significant driver in optical parameters, akin to the free-living species, which had 304 an approximately similar level of correctly classified species (53.8%). However, in contrast to 305 free-living taxa, which are considered to be micro-predators, several of the coral associates are 306 potentially parasites (Stella et al., 2011). The common functionality of their optic parameters (to 307 a degree) remains unclear, although it is known that several species, e.g. Coralliocaris defend 308 their coral host against predators (Marin, 2009a; Stella et al., 2011), perhaps necessitating the 309 need for similar optical acuity to free-living micro predators.

310 Bivalve associates exhibited a 100% correct classification in the DFA analysis, although with 311 reasonable scatter in the scatter plot, and a significant outlier (*C. nipponensis*). Yet the group 312 consists of several genera, including *Conchodytes* and *Anchistus*, which are phylogenetically 313 distant (Kou et al., 2014). Furthermore these species can be differentiated by general bauplan 314 morphologies, ranging from relatively unspecialized (Anchistus and Paranchistus for example) to 315 dorso-laterally compressed (e.g. Conchodytes) (Bruce 1981; Fransen & Reijnen, 2012). Their 316 phylogenetic distance is evidence of multiple host invasions (Kou et al., 2014), but the present 317 analysis reveals considerable convergence in optical parameters, indicative of profound habitat 318 induced restraints.

319 A number of species occupy isolated positions within their respective groups, notably *P. loloata*, 320 P. scriptus, C. nipponensis and L. nudirostris. Although we cannot discount variation in optical 321 parameters of individual eyes, which may have lowered the percentage correctly classified and 322 induced a higher degree of scatter, two species are worthy of further discussion. The corneal part 323 of the eye of *Laomenes* species is characterised by an apical papilla (see illustrations for several 324 species in Marin, 2009b) which contains functional facets, but which are somewhat different in 325 shape to facets elsewhere on the cornea. The relative size as well as the exact position of the 326 papilla has been used as a minor taxonomic character to differentiate between species (Marin, 327 2009b). However, it is known that a large degree of infra-specific variation is present, which 328 unquestionably would influence some of the herein included optical parameters. Periclimenes 329 scriptus appeared isolated within the sea anemone grouping however due to the small size of the 330 specimen (CL 1.25 mm) it is possible that this animal was not fully mature as ovigerous females 331 have a reported CL of 5.0 mm (Ďuriš et al., 2013).

332 **5.** Conclusion

Overall, our analysis demonstrates that there is a significant evolutionary pressure of the host environment on the optic parameters of associate shrimp species, with in many cases congruence being evident between phylogenetically unrelated taxa. This is especially evident in bivalve and sponge associates, and to a lesser extent in other host taxa. This result is in sharp contrast to the disparate morphology of many other body parts of pontoniine shrimps, with significant variation in mouthparts, pereiopods and even general body shape between genera, inhabiting the same host. At the same time, evidence emerges from the optical analysis of recent host switching

- 340 events in certain lineages, where the optical parameters have not evolved to a communality yet,
- 341 especially in the genera *Periclimenaeus* and *Pontonia*, where taxa living in different hosts appear
- 342 to retain a close optical similarity to those living in other taxa.

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Figure 1. Mean relative eye diameter (standardised by post-orbital carapace length) for 83 species of Pontoniinae associated 8 host-symbiont groupings.

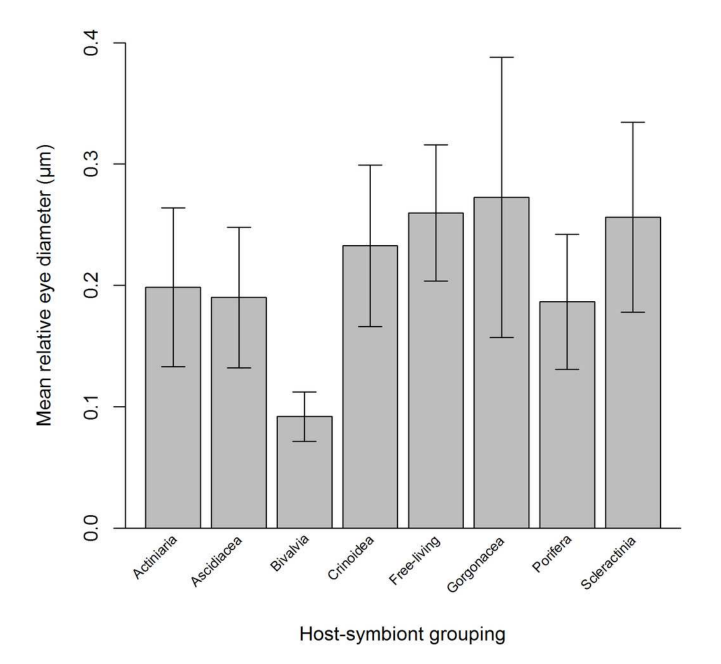


Figure 2. Median eye parameter for 83 species of Pontoniinae from 8 host-symbiont groupings. Significant differences are represented by hosts possessing the same letter A-I (Tukey HSD P<0.05).

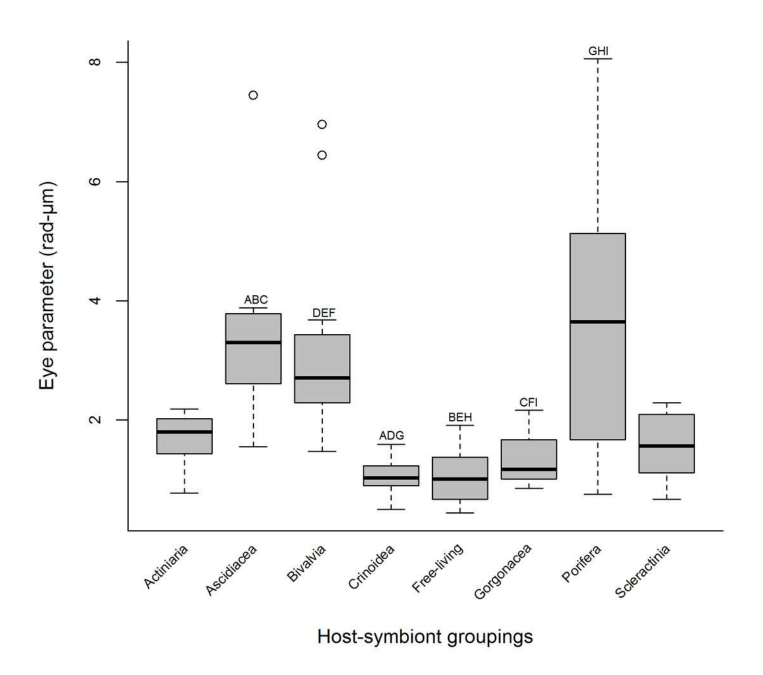


Figure 3. Percentage occurrence of the nebenauge for 83 species of Pontoniinae from 8 host-symbiont groupings.

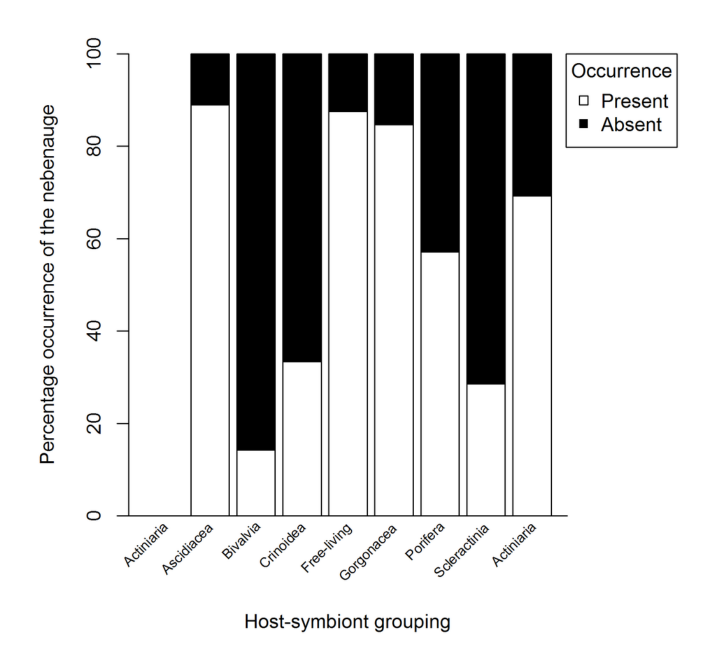


Figure 4. Morphological variation demonstrated by the DFA scores (first and second root only) of all 83 species of pontoniine shrimps (grey circles) displaying the positioning of the centroids for each of the 8 hosts-symbionts groups.

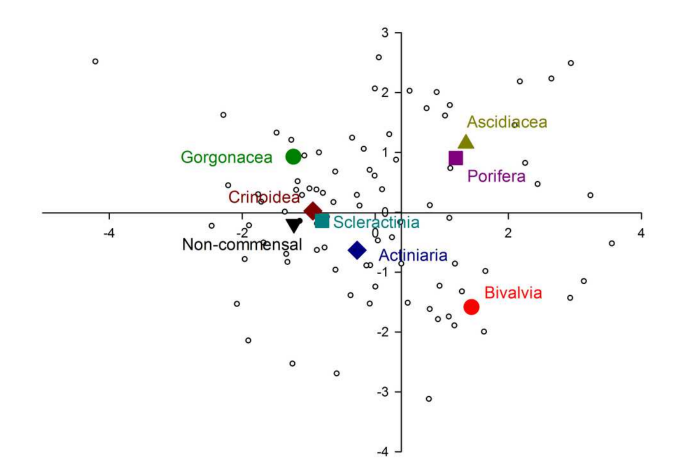


Figure 5. Morphological variation demonstrated by the DFA scores (first and second root only) of Ascidiacea associates.

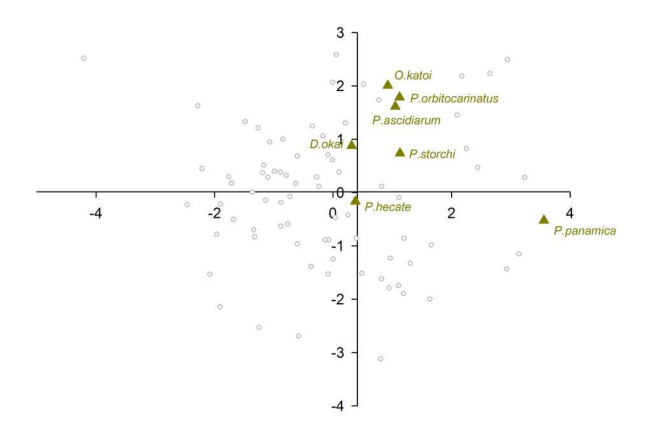


Figure 6. Morphological variation demonstrated by the DFA scores (first and second root only) of Porifera associates.

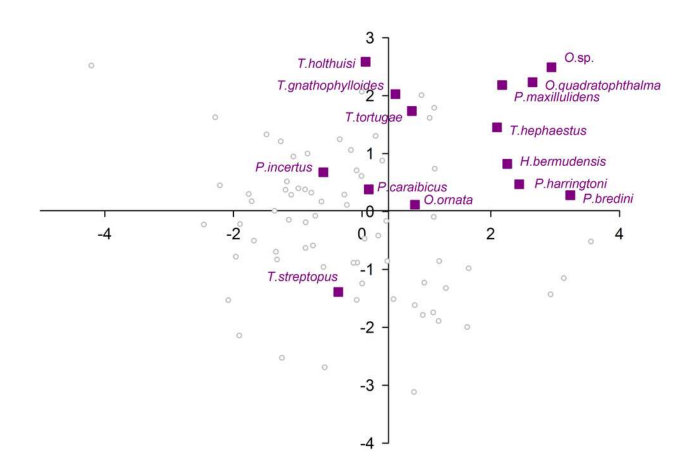


Figure 7. Morphological variation demonstrated by the DFA scores (first and second root only) of Bivalvia associates.

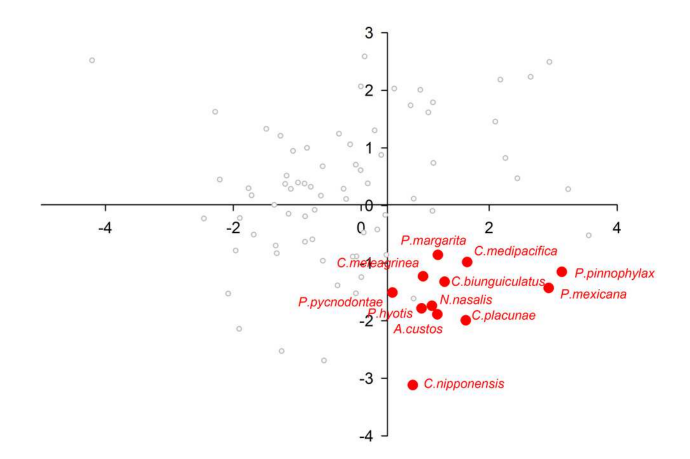


Figure 8. Morphological variation demonstrated by the DFA scores (first and second root only) of Crinoidea associates.

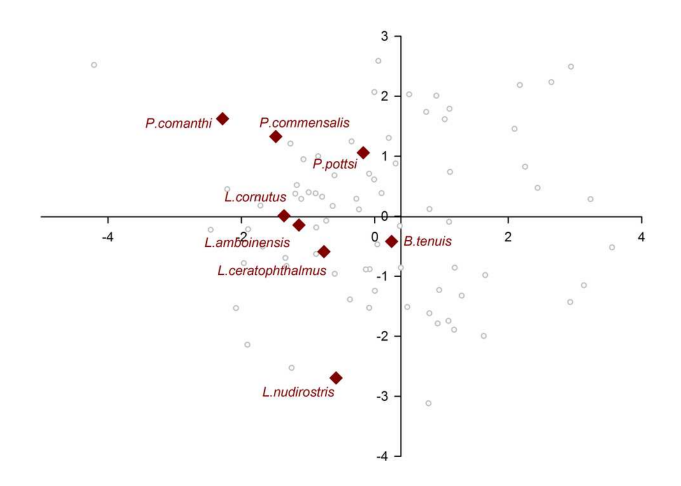


Figure 9. Morphological variation demonstrated by the DFA scores (first and second root only) of Scleractinia associates.

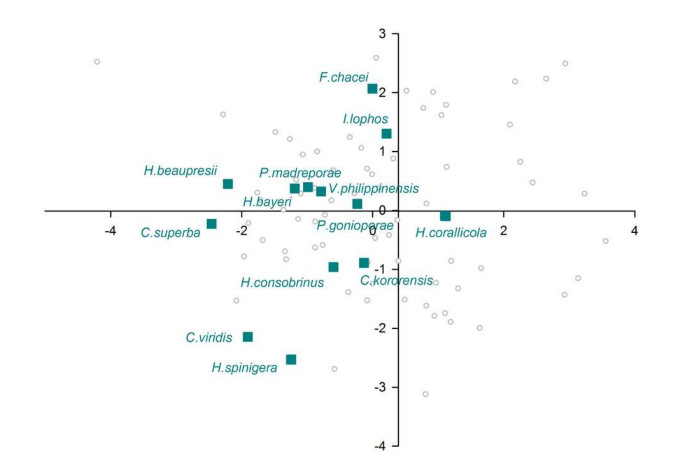


Figure 10. Morphological variation demonstrated by the DFA scores (first and second root only) of non-commensal species.

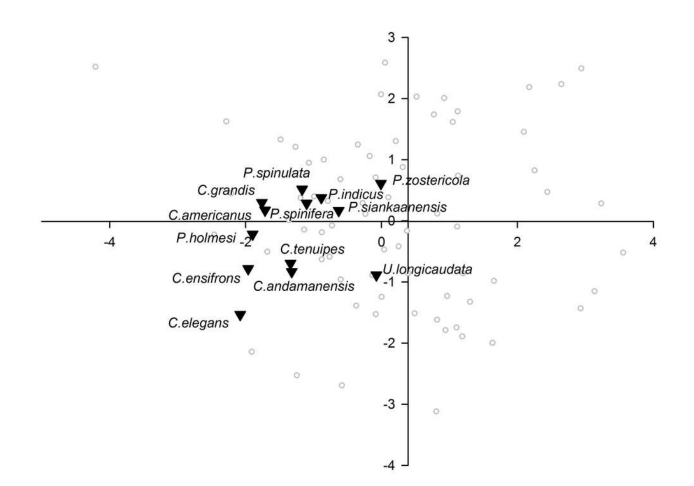


Figure 11. Morphological variation demonstrated by the DFA scores (first and second root only) of Gorgonacea associates.

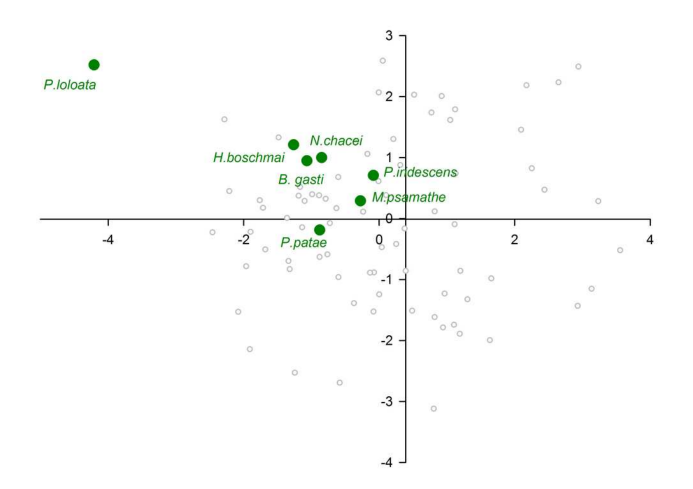


Figure 12. Morphological variation demonstrated by the DFA scores (first and second root only) of Actiniaria associates.

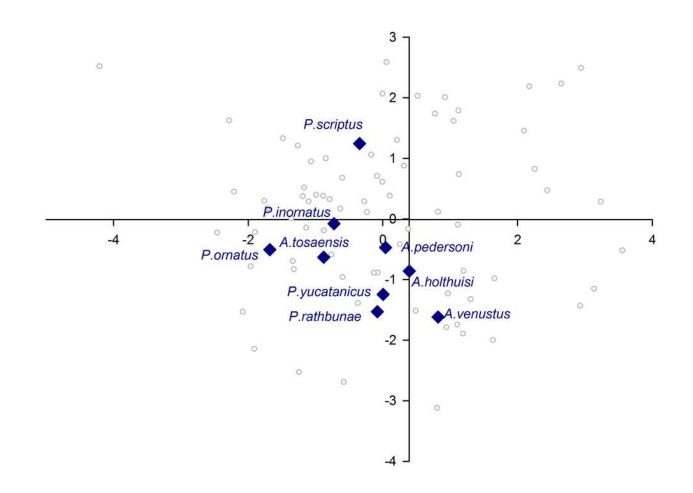


Table 1(on next page)

Table 1. Summary statistics for DFA analysis.

1 Table 1. Summary statistics for DFA analysis

2

| | Eigenvalue | % of | Cumulative | Cumulative Canonical | | Р |
|--------|------------|----------|------------|----------------------|-------|---------|
| | | variance | % | correlation | λ | value |
| Root 1 | 1.436 | 59.1 | 59.1 | 0.768 | 0.194 | < 0.005 |
| Root 2 | 0.864 | 35.5 | 94.6 | 0.681 | 0.473 | < 0.005 |
| Root 3 | 0.090 | 3.7 | 98.4 | 0.288 | 0.882 | 0.482 |
| Root 4 | 0.040 | 1.6 | 100 | 0.196 | 0.962 | 0.561 |

3

Table 2(on next page)

Table 2. Structure matrix of discriminant loadings, with the largest absolute correlation between each variable and any discriminant function indicated by *. All variables were entered simultaneously.

- 1 Table 2. Structure matrix of discriminant loadings, with the largest absolute correlation between
- 2 each variable and any discriminant function indicated by *. All variables were entered
- 3 simultaneously.

4

| Function 1 | Function 2 | | |
|------------|-----------------------------|--|--|
| 0.808* | 0.482 | | |
| 0.718* | 0.166 | | |
| -0.657* | 0.481 | | |
| -0.158 | -0.695* | | |
| | 0.808* 0.718* -0.657* | | |

5

Table 3(on next page)

Table 3. DFA Classification matrix, showing number of species correctly and incorrectly classified into a priori defined groups, expressed as a percentage of within group species numbers.

Table 3. DFA Classification matrix, showing number of species correctly and incorrectly classified into a priori defined groups, expressed as a percentage of within group species numbers.

| | | | |] | DFA classifica | tion | | | |
|-----------------|---------------|------------|------------|----------|----------------|---------------|------------|----------|--------------|
| | | Actiniaria | Ascidiacea | Bivalvia | Crinoidea | Non-commensal | Gorgonacea | Porifera | Scleractinia |
| A nriori grouns | Actiniaria | 22.2 | - | 11.1 | 11.1 | 22.2 | - | - | 33.3 |
| | Ascidiacea | 14.3 | - | 14.3 | - | - | - | 71.4 | - |
| | Bivalvia | - | - | 100.0 | - | - | - | - | - |
| | Crinoidea | 25.0 | - | - | 12.5 | 12.5 | 37.5 | - | 12.5 |
| | Non-commensal | 7.7 | - | - | 15.4 | 61.5 | - | - | 15.4 |
| | Gorgonacea | - | - | - | 28.6 | 14.3 | 42.9 | - | 14.3 |
| | Porifera | 7.1 | - | - | 14.3 | - | - | 78.6 | - |
| | Scleractinia | 15.4 | - | - | 7.7 | 15.4 | - | 23.1 | 38.5 |