

Trophic specializations of damselfishes are tightly associated with reef habitats and social behaviours

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Abstract Despite the increasing need to understand factors shaping community assembly, few studies have simultaneously explored the influence of niche-based and phylogenetic processes. Here, we investigate the relationships between diet, habitat and social behaviour in damselfishes (Pomacentridae) collected in 2014 at Moorea Island (17°30'S, 149°50'W), French Polynesia. Isotope ratios of carbon and nitrogen, in association with stomach contents, delineate three trophic groups: pelagic feeders consuming zooplankton, benthic feeders mainly grazing on algae and an intermediate group feeding on prey from the whole benthic-pelagic compartment. Sulphur isotope ratios indicate segregation between species of the outer reef mostly depending on oceanic input of zooplankton and the lagoonal species relying on locally produced resources or even on terrestrial supply. We demonstrate a tight association between dietary specializations, habitat characteristics and social behaviours, and these correlations are further confirmed by integrating the phylogeny of Pomacentridae. We also provide evidence of phylogenetic conservatism for

the stomach content and the habitat-behaviour characters. However, the isotopic trait is evolutionarily more labile probably because it thoroughly depicts the ecological niche of species. To summarize, pelagic feeders (mainly from the Chrominae) usually form shoals in areas close to the open ocean at a maximum depth of 20 m. Benthic feeders (well represented in the Stegastinae) are ubiquitous, solitary and mostly territorial species found at various depths. The intermediate group includes gregarious species from three subfamilies that forage in the lagoon usually above 12 m depth. Overall, we give insight into processes that have structured the damselfish community in Moorea.

Introduction

Coral reefs are highly productive and structurally complex environments that have promoted diversification of marine teleost fishes (Alfaro et al. 2007; Price et al. 2011). Indeed, they house a wide array of fishes where niche partitioning may be viewed as one of the key factors in the course of their diversification. These resulting high levels of ecological diversity have allowed the stable co-occurrence of similar species (Schoener 1974; Colwell and Fuentes 1975). Within that context, unravelling the multiplicity of factors acting upon the fish community represents a key challenge to be tackled. Community assembly is governed by niche-based processes (including habitat filtering and competitive exclusion) and by the phylogenetic relatedness among organisms (Cavender-Bares et al. 2009; Ndiribe et al. 2013). Nowadays, the use of phylogenetic information has become more widespread in ecological studies focusing on the structure of organisms (Losos et al. 2003) because it gives valuable insight into the pattern of phylogenetic niche conservatism (PNC). This pattern is defined

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as the tendency of lineages to retain their niche-related traits through speciation events and over macroevolutionary time (Crisp and Cook 2012), with a corollary that closely related species exhibit higher ecological similarities than expected based on their phylogenetic relationships (Webb et al. 2002; Losos 2008).

Niche-based processes include the study of feeding ecology of species and populations, which is usually evaluated with two methods. (1) Stomach content analysis is widely recognized as a standard technique to determine the diet but only displays the most recent meal (Hyslop 1980). (2) Stable isotope ratios of carbon (C^{13}/C^{12} , hereafter noted as $\delta^{13}C$) and nitrogen (N^{15}/N^{14} , noted as $\delta^{15}N$) are a classical approach in trophic ecology, allowing time-integrated studies of animal diet or niche assessment (Post 2002; Newsome et al. 2007). More recently, the stable isotope ratio of sulphur (S^{34}/S^{32} or $\delta^{34}S$) has been used as an additional tracer of food sources (McCutchan et al. 2003; Connolly et al. 2004; Moreno et al. 2010). The high variation in $\delta^{34}S$ values between seawater ($\sim 22.9\%$) and sediments ($\sim 1\%$) helps to discriminate the origins of prey and locations of feeding grounds within the marine environment (Kharlamenko et al. 2001; Fry et al. 2008; Mittermayr et al. 2014). Even though $\delta^{34}S$ is extremely helpful to disentangle trophic interactions among organisms, no study has integrated this element to investigate the ecology of coral reef fishes. Other information regarding habitat preferences and social behaviours, needed to appraise all aspects of ecological niches, are commonly collected by visual observations (e.g. Lecchini and Galzin 2005).

Many studies in diverse taxa (e.g. insular terrestrial organisms and marine invertebrates) used the historic relationships among species to evaluate PNC patterns of various ecological traits (e.g. Knouft et al. 2006; Roy et al. 2009; Cooper et al. 2011). The study of phylogenetic signal constitutes one method to detect PNC and greatly varies among traits (Blomberg et al. 2003; Losos 2008). The niche-complementarity hypothesis predicts that species similar along one dimension of the niche must differ along other axes to support their coexistence (Schoener 1974). However, in some clades, there is not always tangible evidence sustaining that prediction because interactions among species may be responsible for a non-random distribution of species within the ecological space. Consequently, distantly related species might share more in common than closely related ones (Losos et al. 2003). To date, there is an overall scarcity of documented cases investigating the interactions between feeding strategy, habitat and behaviour when taking into account phylogenetic information, especially in coral reef fishes.

The Pomacentridae family (damsel-fishes), which groups 399 species (Eschmeyer et al. 2016), represents a suitable system to investigate factors shaping the community in reef

ecosystems. Interestingly, damselfishes show a great diversity of habitats (Ormond et al. 1996; Lecchini and Galzin 2005) and various social behaviours (Fishelson 1998). Furthermore, recent studies have established a firm phylogenetic framework for their radiation (Cooper et al. 2009; Cooper and Santini 2016) and revealed that damselfishes have iteratively evolved along three main feeding strategies, i.e. pelagic feeders, benthic foragers and an intermediate group (Cooper and Westneat 2009; Frédérich et al. 2013, 2016a, b). Several field observations also suggested a strong link between diet, habitat use and social behaviour in Pomacentridae (Emery 1973; Frédérich et al. 2009), but they only constitute a qualitative assumption. The analysis of quantitative data is therefore needed to formally establish these correlations.

The aim of the present study is to examine the congruence between diet, habitat and social behaviour in damselfishes (Pomacentridae) from coral reefs at Moorea Island, French Polynesia. We first explore the niche partitioning of Pomacentridae through an approach combining three stable isotope ratios ($\delta^{15}N$, $\delta^{13}C$ and $\delta^{34}S$) and the analysis of stomach contents. Then, we describe the habitat and social behaviour of each species. We investigate whether there is a correlation between dietary and habitat-behaviour characteristics at the present time. We further explore the relationships between these ecological traits across the phylogeny of Pomacentridae and test whether these traits show a phylogenetic signal. Overall, our data provide a better understanding of the processes ruling the damselfish community assembly and the causes of its structure.

Materials and methods

Sampling and collection of ecological data

The sampling campaign was carried out from April to July 2014 in Moorea, an island located in the South Pacific ($17^{\circ}30'S$, $149^{\circ}50'W$; French Polynesia). The lagoon in Moorea comprises coral patches found in between sand banks and a fringing reef shaped as a wall. The barrier reef includes the outer reef slope and the reef crest and is hereafter referred as the outer reef. Sampling occurred at Opunohu Bay located on the north side of Moorea but also in front of Haapiti in south-west part of the island (Fig. 1; Table 1). Damselfishes—belonging to four subfamilies and 17 species—were either caught with a speargun or by the use of clove oil across the whole reef at Opunohu Bay (Table 1). Four species (*Chromis iomelas*, *Chromis margaritifer*, *Dascyllus flavicaudus* and *Dascyllus trimaculatus*) were ubiquitous with individuals foraging in the two main reef zones (the lagoon and the outer reef), which were treated separately during statistical analyses. Furthermore,

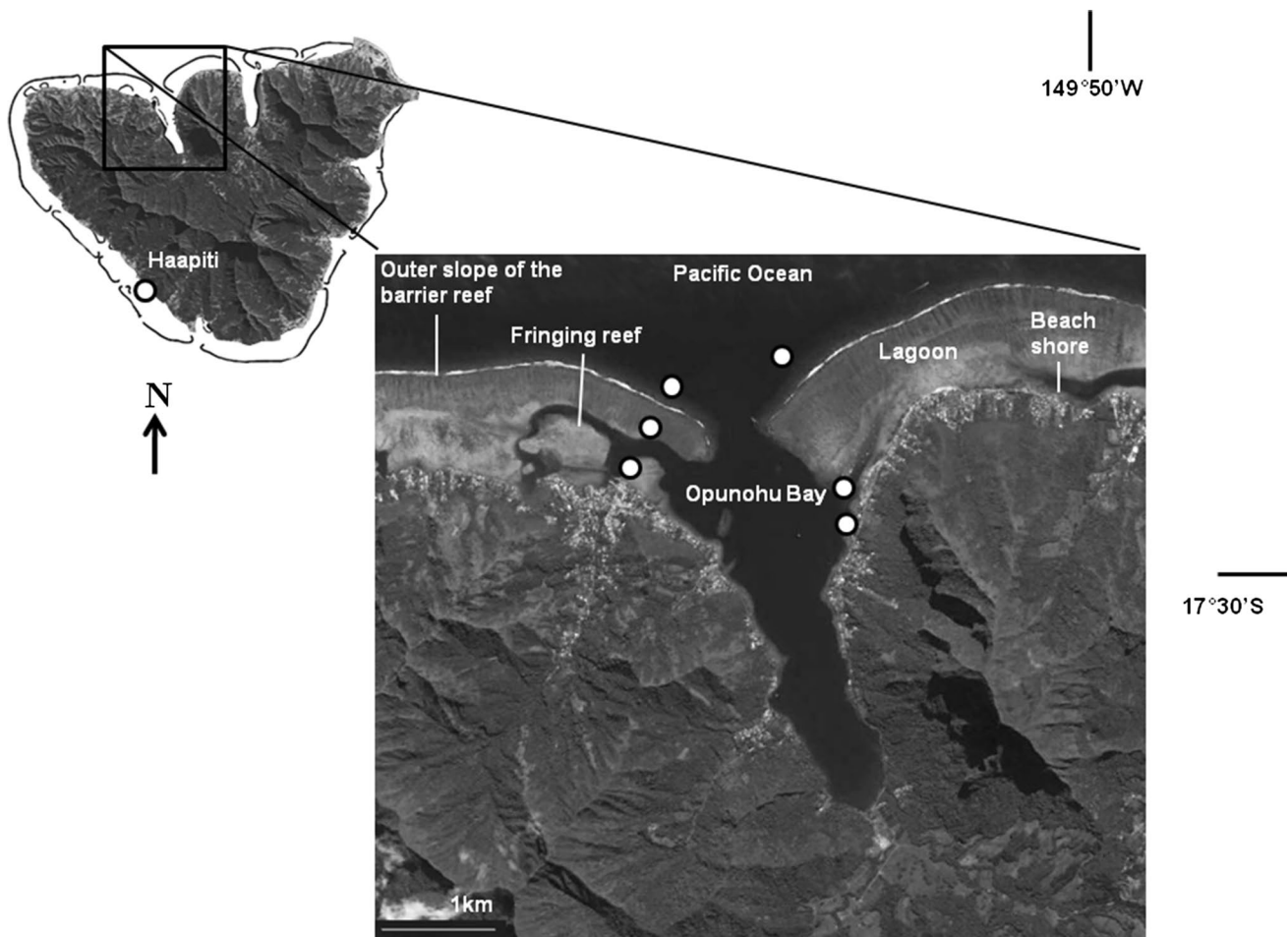


Fig. 1 Map of Moorea (French Polynesia) with sampling sites indicated by white-filled circles. At Opunohu Bay, sampling occurred in coral patches along the beach shore, in the fringing reef and also on

the outer slope of the barrier reef (the outer reef). Another site was located in front of Haapiti where one species was collected along the beach shore

individuals of *Abudefduf septemfasciatus* were collected in two distinct reefs, i.e. in the Opunohu Bay and in front of Haapiti (Fig. 1). Sampling for *A. septemfasciatus* was only conducted along the beach at both reef locations and each population was treated separately when performing statistical tests.

Fishes were euthanized by an overdose of MS-222 (tricaine-methanesulfonate) and placed on ice in cool boxes when brought back to the boat. The standard length of each fish individual was measured to the nearest millimetre using a vernier caliper (Table 1). The entire digestive tract was removed and conserved in 70% ethanol for stomach content analysis. Tissue samples of epaxial muscles were cut off and placed in a drying oven for further isotope analyses. Various food sources and primary producers (e.g. red or brown algae, benthic invertebrates and zooplankton) were taken at the same locations where fishes were collected (in the lagoon and the outer reef). Specifically, zooplankton

was sampled with a 250- μ m net towing at 2 m depth above reefs. This sampling method was replicated three times and is similar to the technique used to study the niche partitioning of damselfishes in Madagascar (Frédérich et al. 2009).

Stomach content analysis

Prey items were identified, sorted to class or order levels and counted under a binocular microscope Leica MS-5. Food sources were classified into six categories reflecting different ecological types of prey: zooplankton, benthic algae, vagile invertebrates, rubble, benthic egg and insect. Zooplankton included copepods (i.e. Poecilostomatoida, Calanoida, Harpacticoida), pelagic eggs of fishes, nauplii and zoea larvae. Benthic algae grouped filamentous and macroalgae (brown and red algae). Vagile invertebrates comprised small crabs, amphipods, isopods, gastropods, polychaetes and platyhelminthes. Rubble was also found

Table 1 Damsel fish species caught in Moorea in 2014 with indication of the number of individuals (ind.) used for each method: stomach content (SC) and stable isotope (SI) analyses

Damsel fish species	Abbreviations	Reef location	Reef zone	Standard length (mm)	Nbr. ind. SC	Nbr. empty stomachs	Nbr. ind. SI
<i>Abudefduf septemfasciatus</i>	<i>A. sept</i>	H & O	Lagoon	103.5–151.6	16	1	17
<i>Abudefduf sexfasciatus</i>	<i>A. sexf</i>	O	Lagoon	53.2–94.7	7	0	6
<i>Chromis acares</i>	<i>C. aca</i>	O	Outer reef	31.6–36.5	12	5	14
<i>Chromis iomelas</i>	<i>C. iom</i>	O	Both	35.5–47.0	11	2	12
<i>Chromis margaritifer</i>	<i>C. marg</i>	O	Both	38.2–59.9	18	0	13
<i>Chromis vanderbilti</i>	<i>C. vand</i>	O	Outer reef	32.3–40.7	19	1	15
<i>Chromis viridis</i>	<i>C. vir</i>	O	Lagoon	31.3–54.8	7	4	11
<i>Chrysiptera brownriggii</i>	<i>Ch. brown</i>	O	Lagoon	30.2–52.2	16	4	20
<i>Dascyllus aruanus</i>	<i>D. arua</i>	O	Lagoon	29.8–47.3	12	0	6
<i>Dascyllus flavicaudus</i>	<i>D. flavi</i>	O	Both	29.1–75.1	12	2	12
<i>Dascyllus trimaculatus</i>	<i>D. trima</i>	O	Both	82.6–97.3	5	2	7
<i>Plectroglyphidodon johnstonianus</i>	<i>P. johnst</i>	O	Outer reef	37.1–58.4	11	3	17
<i>Plectroglyphidodon lacrymatus</i>	<i>P. lacry</i>	O	Outer reef	33.8–66.2	17	1	18
<i>Pomacentrus pavo</i>	<i>Po. pavo</i>	O	Lagoon	33.3–72.7	20	1	20
<i>Pomachromis fuscidorsalis</i>	<i>Pom. fusc</i>	O	Outer reef	38.8–54.4	15	6	21
<i>Stegastes fasciolatus</i>	<i>S. fascio</i>	O	Outer reef	42.8–62.9	16	3	14
<i>Stegastes nigricans</i>	<i>S. nigri</i>	O	Lagoon	70.3–111.8	24	0	17

The reef location indicates where damselfish species were collected either in front of Haapiti (H) in the south-west part of Moorea or in the northern part in Opunohu Bay (O). The reef zone specifies the local sampling sites: the lagoon, the outer reef (outer slope of the barrier reef) or both

in stomachs; its origin assumed to be either rocks or corals. Benthic egg meant eggs from all types of organisms (e.g. fish or invertebrates) deposited on the substrate. Finally, the insect group included two orders of Hexapoda: Hymenoptera and Diptera. The number of items in each food category was recorded and expressed as a percentage to the total number of items for each individual stomach (Hyslop 1980).

Stable isotope analysis

Samples of fish muscle tissues and potential food sources were dried for 48 h at 60 °C and ground into a homogeneous fine powder using a mortar and a pestle. All samples of food sources containing carbonates (e.g. algae samples and zooplankton) were acidified with HCl vapours (37%) for 24 h in order to remove inorganic carbon, which is not related to diet but a source of bias in the analysis of carbon stable isotope ratios (Mateo et al. 2008). Stable isotope ratios of carbon ($^{13}\text{C}/^{12}\text{C}$), nitrogen ($^{15}\text{N}/^{14}\text{N}$) and sulphur ($^{34}\text{S}/^{32}\text{S}$) were measured at the Laboratory of Oceanology of the University of Liège, using an isotope ratio mass spectrometer (IsoPrime 100, Isoprime, UK) coupled in continuous flow to an elemental analyser (vario MICRO cube, Elementar, Germany). Isotope ratios were conventionally expressed as δ values in ‰ relative to the vPDB (Vienna Pee Dee Belemnite) for carbon (C), atmospheric N_2 for

nitrogen (N) and Canon Diablo troilite (CDT) for sulphur (S) (Coplen 2011). Certified reference materials from the International Atomic Energy Agency (IAEA, Vienna, Austria) used were sucrose (IAEA-C6, $\delta^{13}\text{C} = -10.8 \pm 0.5\text{‰}$; mean \pm standard deviation), ammonium sulphate (IAEA-N₂, $\delta^{15}\text{N} = 20.3 \pm 0.2\text{‰}$) and silver sulphide (IAEA-S1, $\delta^{34}\text{S} = -0.3 \pm 0.3\text{‰}$). Hundreds of replicate assays of internal laboratory standards (powder of sulfanilic acid) indicate measurement errors (standard deviation) of $\pm 0.2\text{‰}$ for $\delta^{13}\text{C}$, $\pm 0.3\text{‰}$ for $\delta^{15}\text{N}$ and $\pm 0.5\text{‰}$ for $\delta^{34}\text{S}$.

Description of habitat and social behaviour

The ecology of each studied damselfish species was recorded during underwater visual surveys by one observer. Observations were conducted while scuba diving or snorkelling for 50–60 min at the same location where fishes were caught, i.e. along the beach shore, in the fringing reef and the outer reef (Fig. 1). In order to capture potential variability of behaviours, duplicate observations at each sampling site were done at the same time, between 2 p.m. and 5 p.m. Our dataset was also completed using information from the literature (Fishelson 1998; Hattori and Casadevall 2016; Pratchett et al. 2016). The habitats and social behaviours for 17 species were summarized into five categorical traits (Table 2). (1) The location of fishes along

the coral reef ecosystem was distributed in three categories: “lagoon”, “outer reef” or in “both” parts. (2) Microhabitat requirements were coded as “sandy areas” for fishes found between coral patches in the lagoon, “dead corals/rocks”, “corals (O)” for damselfishes having obligated relationships with their live coral hosts and “corals (F)” for species having facultative relationships with corals which they use infrequently or opportunistically (Pratchett et al. 2016). (3) The depth range was the minimal and maximal depths expressed in metres (Pratchett et al. 2016). (4) The group type and territoriality were coded together: group type consisted of seven categories, whereas territoriality was simply defined as “yes” or “no”. The classification for the group size was: “solitary” when species were exclusively found alone, “solitary or pairing”, “small social groups” when a size-ordered dominance hierarchy existed among fishes, shoals (i.e. aggregations of individuals swimming in different directions): “small shoals” (with 3–10 individuals), “medium shoals” (with 11–20 individuals) or “large shoals” (with >21 individuals) and finally “large schools” (>21 individuals foraging around diverse shelters and swimming in a coordinate manner towards the same direction) (Fishelson 1998; Hattori and Casadevall 2016). (5) The water column position of fishes, defined as its average swimming distance from the substrate, was coded using three main categories: “OS” for species staying all the time close to the bottom, “CS” for species commonly swimming at an approximate distance of 0.5–1.0 m from the substrate and “FR” for damselfishes freely swimming throughout the water column without being restricted to one particular area.

Statistical analyses and phylogenetic comparative methods

Stomach content analysis

Stomach content data were analysed using the PRIMER software version 6.1.6 (Plymouth Routines in Multivariate Ecological Research Ltd, UK; Clarke 1993). The stomach data of each fish individual were treated as a sampling unit. These data were firstly standardized before creating a Bray–Curtis matrix to quantify the compositional dissimilarity. An analysis of similarity (ANOSIM) was then performed to test similarities among stomach contents grouped per species.

Stable isotope analysis

Food sources Differences among food sources were only investigated for zooplankton because it was the only food source with replicates. A *t* test was conducted in R version 3.2.4 (R Core Team 2016) because data were normally distributed and variances homogeneous.

Damselfishes We explored our CNS isotopic dataset of damselfishes with a principal component analysis (PCA) using the PRIMER software version 6.1.6. (Clarke 1993). PCA consists of rotating the axes such that as much variability as possible is accounted on consecutive orthogonal principal components (PCs) depicting a multidimensional space (Davis 1986). This space was thus reduced to two dimensions as the first two PCs cumulated 92.8% of the total variation (Supplementary Fig. S1). The eigenvectors of $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ were similar in magnitude and direction along both PC axes (Supplementary Fig. S1), showing that these two isotope ratios were strongly correlated among species. Since we multiplied our $\delta^{13}\text{C}$ data by -1 in the PCA, the similar $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ eigenvectors revealed that the lower the $\delta^{13}\text{C}$ values, the higher were the $\delta^{15}\text{N}$ values. Conversely, in the PC1-2 space, the $\delta^{34}\text{S}$ eigenvector appeared orthogonal to the $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ eigenvectors (Supplementary Fig. S1), indicating that $\delta^{34}\text{S}$ represented independent information regarding the segregation of species within the isotopic space. Based on these findings, we conducted a second PCA using only $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ values. This allowed us to describe most (89.9%) of the variance in $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ values with a single PC axis (hereafter referred as $\text{PC}_{\text{C-N}}$). Differences among species were thus explored using these $\text{PC}_{\text{C-N}}$ scores and the $\delta^{34}\text{S}$ values, separately. Kruskal–Wallis tests (*kruskal.test*) were conducted with the R package lattice version 0.2-33 because assumptions of normality and homogeneity of variances were not met. When significant differences ($P \leq 0.05$) were found, a post hoc test (*kruskalmc*) was conducted for multiple pairwise comparisons among species with the R package ggirmess version 1.6.4. Furthermore, a hierarchical clustering (dendrogram) was also constructed with PRIMER software version 6.1.6. (Clarke 1993) to further infer similarities among species.

Relationships between diet, habitat and social behaviour

The relationship between trophic data (defined either by stomach content (SC) or by stable isotopes (SI)) and habitat–behaviour information was first tested without phylogenetic information and then by integrating the evolutionary history of damselfishes. For both tests, the first step was to transform data by creating a dissimilarity matrix for each dataset: SC, SI and habitat–behaviour. The Bray–Curtis distance was used for both SC and SI databases given the continuous values and was performed with the R package ecodist version 1.2.9 (Goslee and Urban 2007). As the habitat–behaviour dataset contains only discrete values, we applied the Gower distance using the R package cluster version 2.0.3 (Gower 1971; Maechler et al. 2015). Then a multidimensional space with a principal coordinates analysis (Pcoa) was computed on each dissimilarity matrix

with the R package ape version 3.4 (Paradis et al. 2004). We applied the Cailliez correction, which consists of adding a constant value to the original distances in the matrix, in order to keep only positive values (Cailliez 1983; Legendre and Legendre 1998). Euclidean distances in the two-dimensional (2D) space were significantly correlated with initial Bray–Curtis distances for both SC and SI dissimilarity matrices, and with initial Gower distances for habitat–behaviour dissimilarity matrix (Mantel test with 10^5 iterations, $r = 0.92$; $P < 0.001$; $r = 0.97$, $P < 0.001$; $r = 0.99$, $P < 0.001$, respectively). This step ensures that most of the variability was robustly captured in each 2D space. For accuracy, we also tested whether adding one axis increased the quality of the multidimensional space but it did not. The coordinates in the 2D space (Pcoa 1 and Pcoa 2) for each matrix (i.e. the trophic data matrix, either with SC or with SI, and the habitat–behaviour matrix) have been used for the following tests. This data transformation into multidimensional spaces is conceptually similar to the framework developed by functional ecologists in order to quantify functional diversity (e.g. Mouillot et al. 2014; Maire et al. 2015).

The degree of association between the 2D space of trophic data and the 2D space of habitat–behaviour was assessed using a standard multivariate regression based on Procrustes distances with the function *procD.lm* (Adams and Otárola-Castillo 2013). Then, we performed another Procrustes distance-based method for evaluating regression linear models for the 2D space with phylogenetic information. This approach (D-PGLS) was used to test for independent relationships between trophic and habitat–behaviour data under a Brownian motion model of evolution (Adams 2014b; Collyer et al. 2015). This test was done with the function *procD.pgls*. Both analyses (*procD.lm* and *procD.pgls*) were performed with 10^6 iterations in the R package geomorph version 3.0.1 (Adams and Otárola-Castillo 2013).

Phylogenetic signals of ecological traits

We also explored whether trophic data and habitat–behaviour traits showed a phylogenetic signal, which is defined as the statistical dependence among species trait values due to their phylogenetic relatedness (Revell et al. 2008). This test was applied on the 2D space of SC, SI and habitat–behaviour using the function *physignal* from the R package geomorph and performed with 10^5 iterations (Adams and Otárola-Castillo 2013; Adams 2014a). The calculated K is a mathematical generalization of the Kappa statistic from Blomberg et al. (2003). A $K_{\text{mult}} > 1$ means a stronger resemblance of species than expected under a Brownian motion model of evolution, whereas $K_{\text{mult}} < 1$ indicates a greater malleability of traits.

The molecular time-calibrated and multigene phylogeny of Pomacentridae from Frédérick et al. (2013) was pruned to match the species in the dataset and provided an estimate of the evolutionary relationships among taxa. Almost all the studied species were found in that phylogeny, except for *Plectroglyphidodon johnstonianus*, *Pomachromis fuscidorsalis* and *Chromis acares*. *Plectroglyphidodon johnstonianus* and *P. fuscidorsalis* present very similar morphologies and colour patterns to *P. dickii* and *P. richardsonii*, for which molecular data are available. Thus, the phyletic position of *P. dickii* and *P. richardsonii* was used for *P. johnstonianus* and *P. fuscidorsalis*, respectively. Then, we applied the function *locate.yeti* in the R package phytools version 0.5-20 to place *C. acares* in the phylogeny of damselfishes using continuous traits (Revell 2012; Revell et al. 2015). We used meristic data from Allen (1991) which were then transformed into a multidimensional space following the same steps as for the habitat–behaviour trait (see above). We used the coordinates from the first five Pcoas into the function *locate.yeti* (Mantel test results with 10^5 iterations: $r = 0.88$; $P < 0.001$). We also tested whether adding one dimension improved the quality of the multidimensional space but it was not the case. The time-calibrated phylogeny including the 17 studied species from Moorea was used in the D-PGLS analysis and in the tests for phylogenetic signal.

Results

Trophic diversity

Stomach contents

The stomachs of 273 individuals were analysed but 35 of them were empty (Table 1). In every species, stomachs were partially filled with unidentifiable items. The origin of this amorphous material was uncertain, and thus this item was excluded from all statistical analyses. For the four ubiquitous species, we found no significant differences in stomach contents between the lagoonal individuals and the ones caught in the outer reef (Supplementary Table S1). No significant differences appeared between the population of *A. septemfasciatus* from Opunohu Bay and the one from Haapiti. Overall, we found some significant differences in the stomach contents among the 17 species (ANOSIM global test, $R = 0.501$, $P < 0.001$), but some can be clustered together (pairwise comparison tests, $R > 0.5$; Fig. 2; Supplementary Table S1). Cluster A comprises nine species (*D. trimaculatus*, *C. iomelas*, *C. margaritifera*, *Chromis vanderbilti*, *Pomacentrus pavo*, *Chromis viridis*, *Pomachromis fuscidorsalis*, *D. flavicaudus* and *Chromis acares*) for which zooplankton contributed the most to their

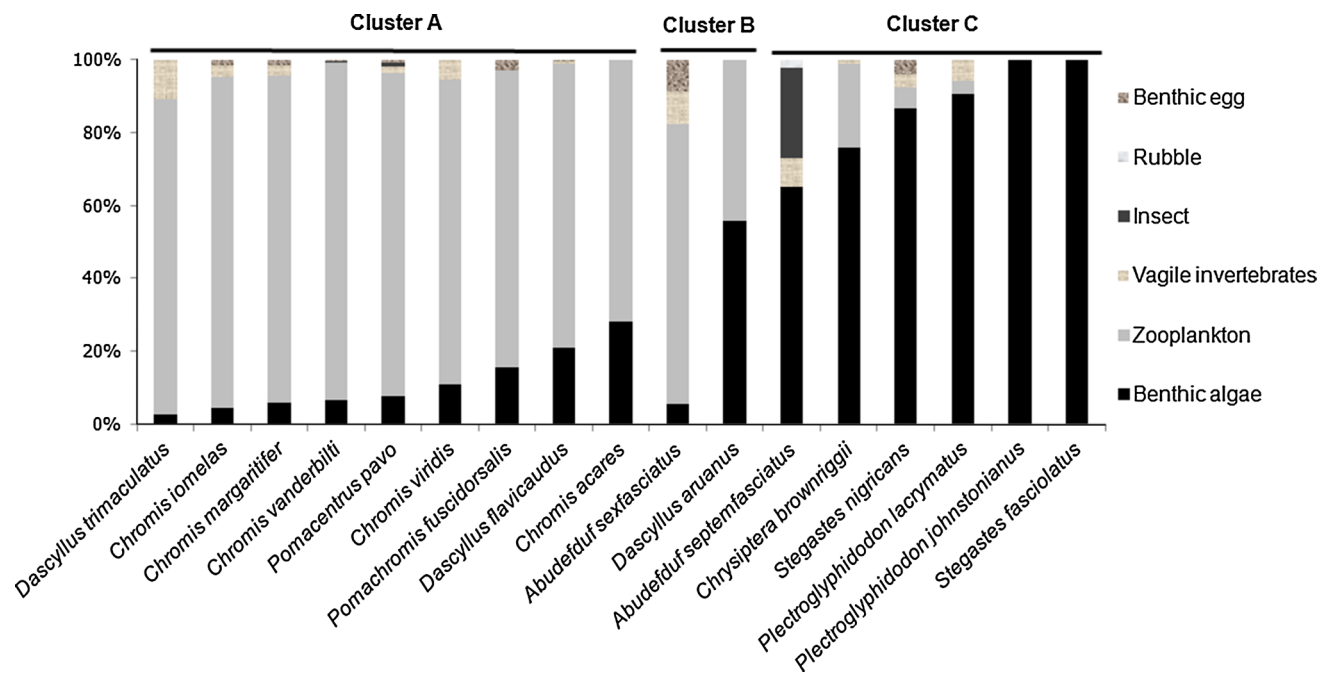


Fig. 2 Dietary compositions of the 17 damselfish species caught in Moorea in 2014. The six food categories are represented by different colour patterns. Clustering was performed with analysis of similarity (ANOSIM) on stomach content data

diet (71–91%). Individuals of *D. trimaculatus* and *C. viridis* were partially feeding on vagile benthic invertebrates, respectively, 6 and 15%. *Abudefduf sexfasciatus* and *Dascyllus aruanus* formed the second cluster (B). Zooplankton and benthic algae almost equally contributed to the diet of *D. aruanus*, whereas benthic food sources (algae, vagile invertebrates and benthic eggs) filled 23% of stomachs of *A. sexfasciatus*. Cluster C grouped *A. septemfasciatus*, *Chrysiptera brownriggii*, *Stegastes nigricans*, *Plectroglyphidodon lacrymatus*, *Plectroglyphidodon johnstonianus* and *Stegastes fasciolatus* for which stomachs were filled with 71–100% of benthic algae. Vagile invertebrates (8%) were also found in *A. septemfasciatus* but insects filled 25% of their stomachs (Fig. 2).

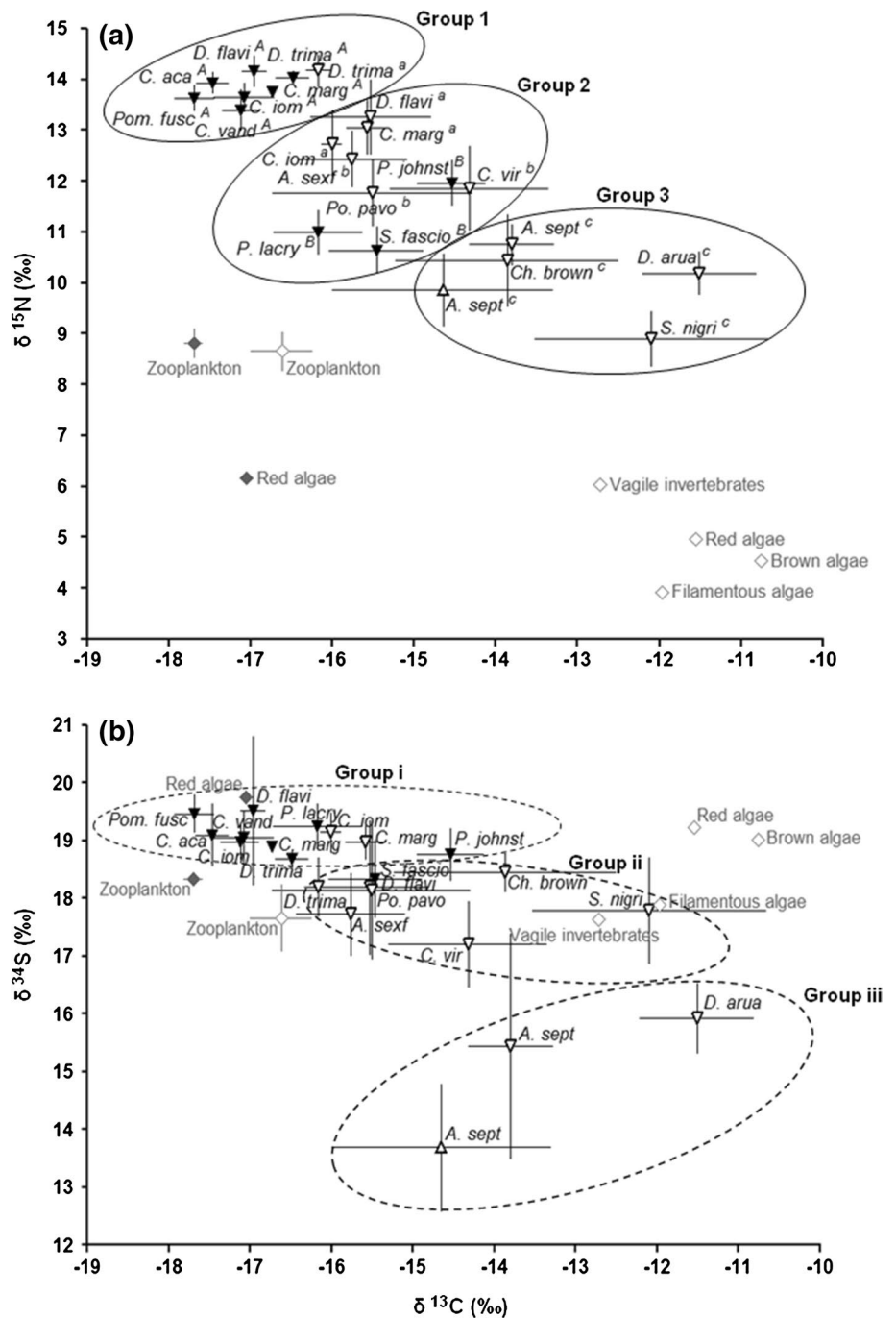
Stable isotopes

The potential food sources displayed lower $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ values than damselfishes (Fig. 3a), but their $\delta^{34}\text{S}$ values were within the same range as the ones $\delta^{34}\text{S}$ of fishes, except for red algae from the outer reef whose value was slightly higher (Fig. 3b). The algal sources from the lagoon had $\delta^{13}\text{C}$ values varying between -12‰ and -10‰ , between 4 and 5‰ for $\delta^{15}\text{N}$ values. In contrast, red algae from the outer reef displayed lower $\delta^{13}\text{C}$ values and higher $\delta^{15}\text{N}$ values. It should be noted that we sampled different types of algae in the outer reef but we only obtained isotope ratios for red algae. The values of vagile invertebrates

from the lagoon were -12.7‰ for $\delta^{13}\text{C}$ and 6.0‰ for $\delta^{15}\text{N}$. Samples of zooplankton collected in the lagoon and in the outer reef showed similar values for $\delta^{15}\text{N}$ and $\delta^{34}\text{S}$ (t tests, $t_3 = 0.46$, $P > 0.05$ and $t_3 = 1.59$, $P > 0.05$), but their $\delta^{13}\text{C}$ values were significantly different (t test, $t_3 = -3.71$, $P < 0.05$). Overall, potential food sources displayed variation in $\delta^{13}\text{C}$ values when comparing the two main reef zones. Zooplankton and algae showed less negative $\delta^{13}\text{C}$ values in the lagoon than in the outer reef (Fig. 3a). However, $\delta^{34}\text{S}$ values (17.7–19.8‰) of these food sources were found within a same range between reef zones (Fig. 3b).

Significant differences among damselfish species according to their $\text{PC}_{\text{C-N}}$ scores were found (Kruskal–Wallis test, $H_{20} = 209.62$, $P < 0.001$). Post hoc tests showed three main groups of species (referred to as groups 1, 2, and 3; Supplementary Table S2), which are illustrated on the plot of $\delta^{15}\text{N}$ versus $\delta^{13}\text{C}$ (Fig. 3a) and further supported by results of the dendrogram (Supplementary Fig. S2). Group 1 is composed of three species exclusively found in the outer reef (*C. acares*, *C. vanderbilti* and *P. fuscidorsalis*), individuals from the outer reef of the four ubiquitous species (*C. iomelas*, *C. margaritifer*, *D. flavicaudus* and *D. trimaculatus*) and also lagoonal individuals of *D. trimaculatus*. Group 2 is composed of nine species in total, including lagoonal individuals of three ubiquitous species (*C. iomelas*, *C. margaritifer* and *D. flavicaudus*), three species strictly from the outer reef (*P. johnstonianus*, *P. lacrymatus* and *S. fasciolatus*) and three species only foraging in

Fig. 3 **a** Average (\pm standard deviation, SD) $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ values of the 17 damselfish species caught in Moorea in 2014. Clustering (Groups 1, 2 and 3) was based on multiple pairwise comparisons on $\text{PC}_{\text{C-N}}$ scores of damselfishes. Grouping within the outer reef was shown with ^A and ^B; whereas it was indicated with ^a, ^b and ^c in the lagoon. **b** Average (\pm SD) $\delta^{34}\text{S}$ and $\delta^{13}\text{C}$ values of the 17 damselfish species caught in Moorea in 2014. Clustering (Groups i, ii and iii) was based on multiple pairwise comparisons on $\delta^{34}\text{S}$ values of damselfishes. Species abbreviations are provided in Table 1. Four species (*Chromis iomelas*, *Chromis margaritifer*, *Dascyllus flavicaus* and *Dascyllus trimaculatus*) were ubiquitous with individuals foraging in the two main reef zones, i.e. the lagoon and the outer reef. One species (*Abudefduf septemfasciatus*) was caught in two distinct reef locations (in front of Haapiti and in Opunohu Bay). An upward-pointing triangle represents species caught in front of Haapiti, whereas a downward-pointing triangle indicates species collected in Opunohu Bay. White-filled triangles represent species caught in the lagoon, whereas black-filled triangles mean species gathered in the outer slope of the barrier reef (the outer reef). Food sources are represented by a diamond icon and were all collected in Opunohu Bay. The white-filled diamond refers to the lagoon, while the grey-filled diamond means that food sources were gathered in the outer reef



the lagoon (*A. septemfasciatus*, *C. viridis* and *P. pavo*). Group 3 consisted of four species only present in the lagoon: *A. septemfasciatus* from Opunohu and Haapiti, *C. brownriggii*, *D. aruanus* and *S. nigricans*. Furthermore, the variation in $\delta^{13}\text{C}$ values of potential food sources might impact the trophic diversity of damselfishes along this isotopic axis. Therefore, we repeated statistical analyses on $\text{PC}_{\text{C-N}}$ scores for each damselfish assemblage: the one from the

outer reef and the one from the lagoon. Kruskal–Wallis test and multiple pairwise comparisons delineated two groups of species within the outer reef assemblage ($H_8 = 95.51$, $P < 0.001$). The first group comprised seven species including *Pomachromis* sp., four *Chromis* and two *Dascyllus* spp. (Supplementary Table S3; Fig. 3a). The second group in the outer reef contained both species of *Plectroglyphidodon* and *S. fasciolatus*. In the lagoon, significant

Table 2 Habitat and behavioural variables coded into five categories for each damselfish species caught in Moorea

Damselfish species	Reef location	Microhabitat requirements ^a	Depth (m) ^a	Group type & territoriality ^{b,c}	Position in the water column
<i>Abudefduf septemfasciatus</i>	Lagoon	Sandy areas	0–3	Solitary or pairing, No	FS
<i>Abudefduf sexfasciatus</i>	Lagoon	Corals (F)	1–15	Small shoals, No	FS
<i>Chromis viridis</i>	Lagoon	Corals (O)	1–12	Large shoals, No	CS
<i>Chrysiptera brownriggii</i>	Lagoon	Corals (F)	0–2	Solitary, No	CS
<i>Dascyllus aruanus</i>	Lagoon	Corals (O)	1–12	Small social group, No	CS
<i>Pomacentrus pavo</i>	Lagoon	Corals (F)	1–16	Small shoals, No	CS
<i>Stegastes nigricans</i>	Lagoon	Dead corals/rocks	1–12	Solitary, Yes	OS
<i>Chromis iomelas</i>	Both	Corals (F)	3–35	Large shoals, No	CS
<i>Chromis margaritifer</i>	Both	Corals (F)	2–20	Large shoals, No	CS
<i>Dascyllus trimaculatus</i>	Both	Corals (F)	1–55	Medium shoals, No	FS
<i>Dascyllus flavicaudus</i>	Both	Corals (F)	3–40	Medium shoals, No	CS
<i>Chromis acares</i>	Outer reef	Corals (F)	2–37	Large shoals, No	CS
<i>Chromis vanderbilti</i>	Outer reef	Corals (F)	2–20	Large shoals, No	CS
<i>Plectroglyphidodon johnstonianus</i>	Outer reef	Corals (O)	2–20	Solitary, Yes	OS
<i>Plectroglyphidodon lacrymatus</i>	Outer reef	Dead corals/rocks	2–12	Solitary, Yes	OS
<i>Pomachromis fuscidorsalis</i>	Outer reef	Corals (F)	1–20	Large shoals, No	OS
<i>Stegastes fasciolatus</i>	Outer reef	Dead corals/rocks	0–5	Solitary, Yes	OS

Species were classified according to their predominant (though not necessarily exclusive) ecological features. The reef location indicates where individuals were caught: lagoon, outer reef or both (lagoon and outer reef). Microhabitat requirements are “Sandy areas”, “Corals” (coral-dwelling species, further divided into those that are obligate (O) or facultative (F) coral dwellers) and “Dead corals/rocks”. The group type and territoriality were coded together. The depth range represents the minimal and the maximal depths expressed in metres. The position in the water column was coded as “FS” for freely swimmers, “CS” for species staying close to the substrate at a distance of 0.5–1.0 m and “OS” for species staying on the substrate all the time. ^a Information were completed with Pratchett et al. (2016), ^b with Hattori and Casadevall (2016) and ^c with Fishelson (1998)

differences among species revealed the presence of three groups (Kruskal–Wallis test, $H_{11} = 94.45$, $P < 0.001$; Supplementary Table S4; Fig. 3a). The first group consisted of lagoonal individuals from the four ubiquitous species (two *Chromis* spp. and two *Dascyllus* spp.) and the second group comprised *A. sexfasciatus*, *C. viridis* and *P. pavo*. Finally, the third group included the same species of Group 3.

Significant differences among damselfish species were also found according to their $\delta^{34}\text{S}$ values (Kruskal–Wallis test, $H_{20} = 153.64$, $P < 0.001$). Post hoc tests delineated three main groups (referred to as groups i, ii, and iii; Supplementary Table S5, Fig. 3b), which were further assessed by results of the dendrogram (Supplementary Fig. S3). $\delta^{34}\text{S}$ values highlighted a segregation between damselfish species from the outer reef and those from the lagoon (Fig. 3b). Group i consisted of five species exclusively caught on the outer reef (*C. acares*, *C. vanderbilti*, *P. fuscidorsalis*, *P. lacrymatus* and *P. johnstonianus*), individuals from the outer reef of four ubiquitous species (*C. iomelas*, *C. margaritifer*, *D. flavicaudus* and *D. trimaculatus*) and finally comprised lagoonal individuals belonging to *C. iomelas* and *C. margaritifer*. Group ii included five species all from the lagoon (*A. sexfasciatus*, *C. viridis*, *C. brownriggii*, *P. pavo* and *S. nigricans*) and lagoonal assemblages of *D.*

flavicaudus and *D. trimaculatus*. But it also comprised one species (*S. fasciolatus*) from the outer reef. Finally, Group iii encompassed two lagoonal species, *A. septemfasciatus* (from Haapiti and Opunohu) and *D. aruanus*.

Diversity of habitat and social behaviour

Data regarding habitat and social behaviour are summarized in Table 2. Seven species were exclusively caught in the lagoon. Five of them were gregarious or found in pairs and the other two (*C. brownriggii* and *S. nigricans*) presented a solitary behaviour. They were mostly living above 12–16 m depth; however, *A. septemfasciatus* and *C. brownriggii* were not encountered below 2–3 m. The two *Abudefduf* sp. were freely swimming in the water column, while *C. viridis*, *C. brownriggii*, *D. aruanus* and *P. pavo* were staying close to the bottom at a distance of 0.5–1.0 m. A majority of these species were found above various types of substrate even though *D. aruanus* and *C. viridis* were mainly foraging within live, branching corals. *Abudefduf septemfasciatus* was the only species found in sandy areas in between coral reef patches, and *S. nigricans* had its shelter in dead corals or rocks. Four species (*C. iomelas*, *C. margaritifer*, *D. trimaculatus*

and *D. flavicaudus*) were ubiquitous and non-territorial. They were all facultative coral-dwelling species forming medium to large shoals in maximum depths of 20 to 55 m. Only *D. trimaculatus* was observed to freely swim through the entire water column. Finally, the six remaining species exclusively foraged in the outer reef. Three of them (*C. acares*, *C. vanderbilti* and *P. fuscidorsalis*) were gregarious and facultative coral dwellers but staying close to the substrate at maximal depths of 20–37 m. The three last species (*P. johnstonianus*, *P. lacrymatus* and *S. fasciolatus*) were solitary and exhibited a territorial behaviour. *Plectroglyphidodon johnstonianus* was found deeper than the two other species and was also more restricted into the type of substrate (i.e. live corals).

Relationship between diet, habitat and social behaviour

Standard multivariate linear regressions revealed significant relationships between trophic data, defined either by stomach contents (SC) or by stable isotopes (SI), and habitat–behaviour information. These correlations were further confirmed when integrating the phylogenetic relationships among species using the D-PGLS analysis (Table 3).

Phylogenetic signals of ecological traits

Both stomach contents ($K_{\text{mult}} = 1.40$, $P = 9 \times 10^{-4}$) and habitat–behaviour information ($K_{\text{mult}} = 1.17$, $P = 1 \times 10^{-4}$) showed significant phylogenetic signals. On the other hand, trophic data from stable isotopes did not present any phylogenetic signals ($K_{\text{mult}} = 0.65$, $P = 0.22$).

Discussion

Trophic ecology of damselfishes in Moorea

At Toliara reef in Madagascar, Frédérich et al. (2009) defined three trophic groups of damselfishes, i.e. pelagic feeders, benthic feeders and an intermediate group. A similar level of trophic diversity is observed within the Pomacentridae from Moorea. The principal criterion used to differentiate the pelagic feeders from other trophic strategies is their propensity to almost exclusively feed on zooplanktonic prey (Frédérich et al. 2016b). Based on our dual dietary tracers (C, N isotope ratios and stomach contents), seven species are representative of this trophic strategy. Three of them (*C. acares*, *C. vanderbilti* and *P. fuscidorsalis*) are only foraging in the outer reef, whereas the other four species (*C. iomelas*, *C. margaritifera*, *D. flavicaudus* and *D. trimaculatus*) also have individuals distributed in lagoonal zones close to the reef crest. As previously suggested by Wyatt et al. (2012), zooplanktivory is more common in

Table 3 Results of standard multivariate linear regression and D-PGLS (multidimensional phylogenetic regression least squares) tests applied on the 17 damselfish species from Moorea (collected in 2014) to test the degree of association between the two-dimensional (2D) space of trophic data (either obtained with stomach content (SC) or with stable isotope (SI) analyses) and the 2D space depicting habitat–behaviour information

Standard multivariate linear regression analysis (without phylogeny)			D-PGLS analysis (with phylogeny)		
<i>F</i>	<i>R</i> ²	<i>P</i> value	<i>F</i>	<i>R</i> ²	<i>P</i> value
<i>2D space of SC versus 2D space of habitat–behaviour</i>					
17.19	0.71	2×10^{-4}	4.84	0.44	0.01
<i>2D space of SI versus 2D space of habitat–behaviour</i>					
17.04	0.71	1×10^{-5}	15.71	0.69	6×10^{-6}

areas in proximity with the open ocean probably because of its constant input of water column-based plankton.

Benthic feeders include algivorous and corallivorous species and also damselfish species that mainly eat small benthic invertebrates. Both trophic markers reveal that six species (*A. sexfasciatus*, *C. brownriggii*, *P. johnstonianus*, *P. lacrymatus*, *S. fasciolatus* and *S. nigricans*) belong to this trophic group. Contrary to the zooplanktivorous species, benthic feeders are found across the whole coral reef ecosystem in Moorea, i.e. from the beach shore to the outer reef. Even though benthic algae were the main food items for *P. johnstonianus* at Moorea, other studies suggest that this species might be able to bite coral polyps (Allen 1991; Kuo and Shao 1991; Ho et al. 2009).

Frédérich et al. (2009) also defined an intermediate trophic group for which the types and origins of prey are diverse. In Moorea, four species belong to this feeding strategy: *P. pavo*, *C. viridis*, *A. sexfasciatus* and *D. aruanus*. Stomach contents reveal that *P. pavo* and *C. viridis* mainly feed on zooplankton but also ingest benthic algae and invertebrates (Fig. 2), which is confirmed by their $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ values (Fig. 3a). Both species are not only chasing zooplankton but also supplement their diet with local, benthic prey. Trophic markers (C and N isotope ratios and stomach contents) provide evidence of omnivory in *A. sexfasciatus*. At Toliara reef, *A. sexfasciatus* was exclusively feeding on zooplankton (Frédérich et al. 2009), while the feeding regime of populations from Seychelles was classified as intermediate (Manica 2002). This geographic variation among populations supports trophic opportunism in *A. sexfasciatus*. Finally, *D. aruanus* also shows a dietary plasticity as this species feeds on planktonic prey and benthic algae in (almost) equal proportions, and its isotope ratios for C and N are very distinct (Fig. 3a). Frédérich et al. (2010) found a gradual shift of foraging tactics related to the size of *D. aruanus*, with smaller individuals depending more on food sources from the benthic compartment

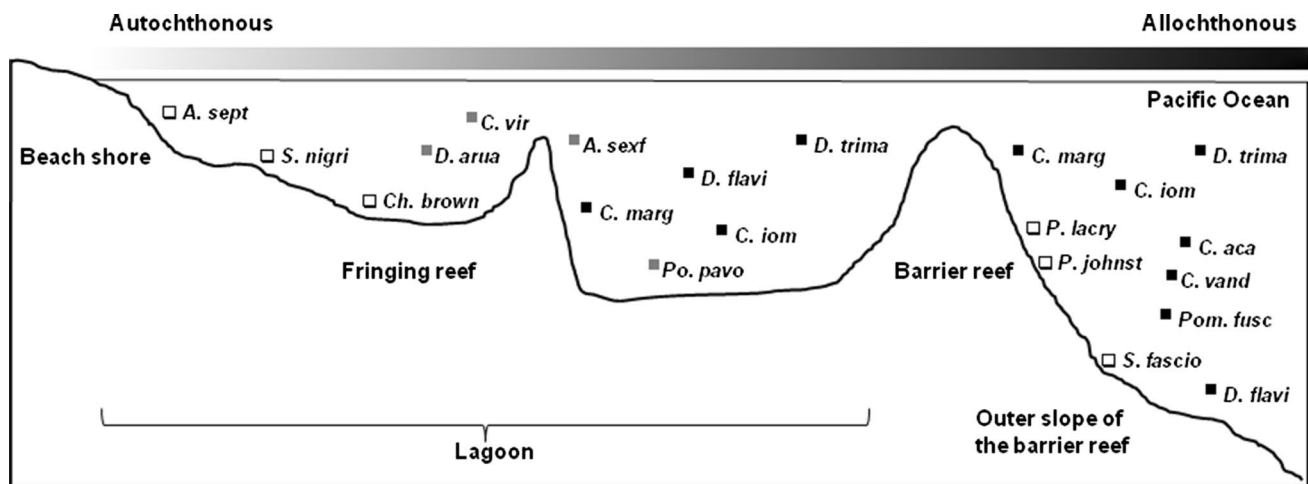


Fig. 4 Position of damselfish species along a schematic representation of the coral reef ecosystem in Moorea, French Polynesia. Species abbreviations are found in Table 1. Trophic groups are represented by *white-filled squares* for “benthic feeders”, *grey-filled squares* for “intermediate group” and *black-filled squares* for “pelagic feeders”.

and, with larger ones mainly feeding on pelagic prey. The sampled *D. aruanus* in our study differed in length (29.8–47.3 mm; Table 1), and thus the large diet breadth in Moorea could also be related to size variation.

Dietary cross-reef gradient

Research devoted to the study of variations in trophodynamics remain scarce, especially for coral reef ecosystems (see exceptions in Kolasinski et al. 2009; Greenwood et al. 2010; Wyatt et al. 2012; Davis et al. 2014; Hanson et al. 2016). The coral reef ecosystem of Moorea presents a complex topography, including a protected lagoon with many coral patches, a fringing reef shaped as a wall, and a barrier reef with a crest bounding the open ocean (Fig. 4). The influence of these morphological features on the trophic ecology of Pomacentridae is emphasized by the significant differences in S isotope ratios among species. Indeed, $\delta^{34}\text{S}$ has been used to unravel food sources for mobile consumers (Jardine et al. 2012) and to ascertain within-producer variability (Connolly et al. 2004; Hamilton 2009; Mittermayr et al. 2014). Pelagic producers using seawater sulphates ($\sim 22.9\%$) have higher $\delta^{34}\text{S}$ values, whereas consumers depending on sulphur from sediments ($\sim 1\%$) have lower $\delta^{34}\text{S}$ values (Kharlamenko et al. 2001; Connolly et al. 2004). Consequently, $\delta^{34}\text{S}$ values help to discriminate between organisms feeding on prey from the pelagic compartment and the others relying on food from the benthos. Here, the $\delta^{34}\text{S}$ results revealed that the assemblage of damselfishes from the outer reef (Group i) can be segregated from the pomacentrid assemblage (Group ii, iii) present in the protected lagoon (Fig. 3b). The diet of

The gradient from *white to black colour* indicates the degree of reliance on food sources from an autochthonous/locally produced origin towards an allochthonous/oceanic provenance. The axes are not to scale

Group i is mainly dependent on oceanic productivity and its subsequent input of planktonic prey. On the other hand, $\delta^{34}\text{S}$ values of Group ii show a higher reliance on local, autochthonous resources (filamentous algae and vagile invertebrates). The species, *A. septemfasciatus*, belonging to Group iii reinforces this dietary gradient across the coral reefs along the coastline at Moorea. This species uniquely forages above sandy banks surrounded by coral patches next to the beach shore. The low $\delta^{34}\text{S}$ value observed in *A. septemfasciatus* is probably related to insects found in their stomachs, indicating a dietary dependence on terrestrial input. The ingress of water column-based phytoplankton in the coral reef food-web is not restricted to zones in permanent contact or close to the open ocean, but is rather progressively distributed. This hypothesis is sustained by findings of two studies conducted, respectively, on the pelagic feeder *D. flavicaudus* (Hanson et al. 2016) and on the benthic feeder *S. nigricans* (McMahon et al. 2016). Results on *D. flavicaudus*, also collected within the Opunohu Bay in Moorea, revealed that this species actively selects oceanic copepods irrespective of the fish position in the reef (lagoon versus outer reef; Hanson et al. 2016). In spite of its foraging area in shallow lagoonal waters and its benthic dietary specialization, *Stegastes nigricans* still partially assimilates C from oceanic sources (McMahon et al. 2016). Overall, the trophic ecology of damselfishes in Moorea is intrinsically linked to their locations within the reef ecosystem. The trophic strategies reflect the influx of oceanic-derived plankton and its assimilation by the reef food-web. The majority of pelagic feeders is found in areas closer to the open ocean or outside the reef system, all benthic feeders are ubiquitous and the intermediate

feeding strategy is only present in the protected lagoon (Fig. 4).

The relationships between diet, habitat and social behaviour

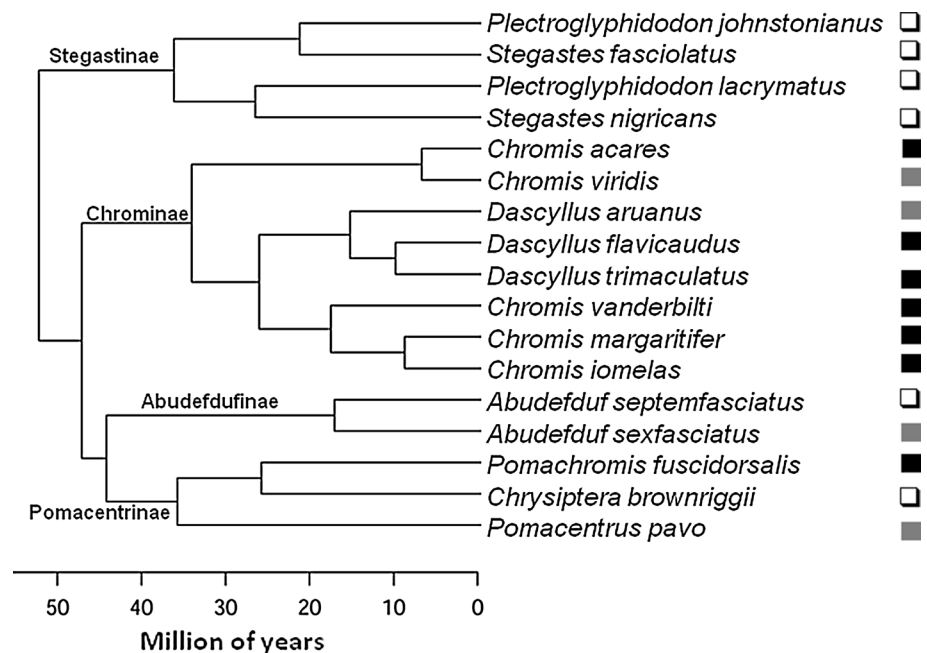
Previous studies about damselfishes carried out in Florida and Madagascar have suggested that diet, habitat and social behaviour could be correlated (Emery 1973; Frédérick et al. 2009). Here, we quantitatively demonstrate a tight association between these ecological variables in the damselfish community from Moorea. We further confirm these correlations when the phylogenetic relatedness among damselfish species is integrated. Almost all pelagic feeders (i.e. 4 *Chromis* spp. and 2 *Dascyllus* spp.) are found within the Chrominae subfamily (Fig. 5), except for *P. fuscidorsalis*. They exhibit similar social behaviours forming large aggregations (either shoals or schools) up to 20 m depth and live in areas bathed by the open ocean or just behind the barrier reef crest. They are also associated with either live or dead corals. The second most species-rich guild in Moorea is the benthic feeders, and within that group, four species belong to the Stegastinae subfamily (i.e. 2 *Stegastes* spp. and 2 *Plectroglyphidodon* spp.). In contrast to pelagic feeders, benthic feeders mostly display a territorial, solitary behaviour, and they can establish their shelter on other types of substrate than corals. Only, *P. johnstonianus* forages among branches of live corals. *Stegastes nigricans* and *P. lacrymatus* are able to harvest macroalgal lawn like farmers (Hata and Makoto 2002; Ceccarelli et al. 2005). The last trophic

strategy—the intermediate group—is represented by *A. sexfasciatus* in the Abudefdufinae, *P. pavo* in the Pomacentrinae and *C. viridis* and *D. aruanus* in the Chrominae. Species from the intermediate group are all gregarious, but only forage in the protected lagoon at usually shallower depths than pelagic feeders. Two of them (*C. viridis* and *D. aruanus*) are obligate inhabitants of live corals.

Ecological trait conservatism and community structure

Diet, defined with stomach contents, presents a phylogenetic signal suggesting some conservatism through the evolution of damselfish communities. We also provide some evidence of conservatism for the behaviour which seems, therefore, less labile than previously thought (Blomberg et al. 2003). Contrary to the study of Blomberg et al. (2003), we coded this character with ecological information and created a unique multivariate trait called “habitat–behaviour”. This combination of traits delineates a high level of habitat and behavioural diversity which appears relatively conserved through the evolution of damselfishes. In general, the causes of conservatism and PNC patterns are diverse, can interact with one another and produce complex outcomes (Revell et al. 2008; Crisp and Cook 2012). They include both direct (e.g. heritability, selection) and indirect processes (e.g. antagonistic niche interactions or dispersal opportunity) (Crisp and Cook 2012). Here, closely related pomacentrid species feed on the same types of prey and exhibit similar habitat features and social behaviours (e.g. Stegastinae and Chrominae). Thus, this

Fig. 5 Consensus time-calibrated tree illustrating the three dietary strategies for the 17 damselfish species from Moorea and with indication of the subfamilies. Trophic groups are represented by white-filled squares for “benthic feeders”, grey-filled squares for “intermediate group” and black-filled squares for “pelagic feeders”



ecological resemblance seems to be greater than expected under a Brownian motion model of evolution. In agreement with the concept of niche retention (Cooper et al. 2010), damselfish lineages could have somehow inherited these characters from their ancestors and slowly diverged. This equivalency might also arise from stabilizing selection and constraints (Cooper et al. 2010; Crisp and Cook 2012). A higher dietary specialization (primarily feeding on one particular type of prey), a narrowing of the habitat–behaviour niche (a partial exploitation of the core species niche) and a limitation of larval dispersal opportunities (induced by physical barrier) are among the possible constraints which could generate a PNC pattern for the stomach content and the habitat–behaviour traits.

Interestingly, trophic specializations defined by the CNS isotope ratios are evolutionarily more labile than diet from stomach contents. The greater malleability of isotopic trait over macroevolutionary time is probably due to its complex signature which embodies more facets of ecological niches (i.e. the dietary, habitat and behavioural components) and over a long period of time, from weeks to months (Tieszen et al. 1983). The C and N isotope ratios are primarily proxies of trophic ecology (i.e. type of prey, foraging location) but may also reflect internal physiological changes during special events such as ontogeny, metamorphosis, starvation or even pregnancy (e.g. Habran et al. 2010; Hammer-schlag-Peyer et al. 2011; Frédérick et al. 2012; Gajdzik et al. 2015). Furthermore, the S isotope ratios mostly reflect the fish position within the reef-scape (lagoon versus outer reef) and the partitioning of microhabitats within each reef zone (pelagic versus benthic compartments). Isotopic data integrate several types of biological information (including ecology and physiology), which might curb the phylogenetic signal of the isotopic trait during the evolution of lineages. The lack of evidence of PNC pattern for the isotopic trait, in contrast to the pattern of stomach content data, suggests that the true diet (a combination of these two techniques) of damselfishes could not be so well matched to the phylogeny.

Furthermore, detecting a phylogenetic signal is context-dependent and influenced by many factors (Losos 2008). Indeed, the signal of a subset of species, such as the studied community of damselfishes, may differ from the one for more inclusive groups (Losos 2008). At the family level, damselfishes observed a macroevolutionary pattern made of iterative ecological radiation and convergences (Frédérick et al. 2013, 2016a). Our results illustrate that this particular process of diversification has produced communities in which closely related species are more ecologically similar than expected by chance. This outcome differs from the case of *Anolis* lizards, where ecological similarity and phylogenetic relatedness is very weak (Losos et al. 2003). These contrasting results from two old

radiations (>15–20 million years ago) suggest that community evolution may occur in different, complex ways. Here, the phylogenetic signals detected may reflect the prevalence of geographic settings producing ecologically similar species during the evolutionary genesis of damselfish community structure.

Conclusion

The trophic ecology of damselfishes in Moorea reveals two opposing trophic strategies: pelagic feeders depending on water column-derived zooplankton and benthic feeders relying on food (algae and vagile invertebrates) from the benthos. In between, there are species displaying different degrees of food exploitation throughout the whole benthic–pelagic compartment. Sulphur isotope ratios show a gradient across the coral reef ecosystem, from a system mostly relying on oceanic support (i.e. the outer reef) towards a system more dependent on autochthonous local productivity (i.e. the lagoon). We demonstrate a tight association between dietary, habitat and behavioural traits in the community of damselfishes from Moorea. Diet defined by stomach contents and habitat–behaviour are conserved through evolution of damselfishes. On the other hand, the isotopic trait is evolutionarily more labile probably because it is affected by a complex array of factors related to diet, habitat and physiology. Further studies focusing on other clades of reef fishes and including other damselfish communities from various regions would improve our understanding of the processes shaping the assembly of fish community in the coral reef ecosystem.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

Informed consent Informed consent was obtained from all individual participants included in the study.

Human and animals rights All applicable international, national, and/or institutional guidelines for the care and use of animals were followed.

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