Intestinal Microbiome Richness of Coral Reef Damselfishes
(*Actinopterygii*: Pomacentridae)

Christopher R J Kavazos, Francesco Ricci, William Leggat, Jordan M Casey, J Howard Choat and Tracy D Ainsworth

*Biological, Earth and Environmental Sciences, The University of New South Wales, Kensington, NSW 2052, Australia; School of Environmental and Life Sciences, The University of Newcastle, 10 Chittaway Dr, Ourimbah, NSW 2258, Australia; Australian Research Council Centre of Excellence for Coral Reef Studies, James Cook University, Townsville, QLD 4811, Australia; PSL Université Paris: EPHE-UPVD-CNRS, USR 3278 CRIOBE, Université de Perpignan, Perpignan 66100, France; Laboratoire d’Excellence “CORAIL,” Université de Perpignan, Perpignan 66100, France; College of Science and Engineering, James Cook University, Townsville QLD 4814, Australia; Centre of Marine Bio-Innovation, The University of New South Wales, Kensington, NSW 2052, Australia*

1E-mail: f.ricci@unsw.edu.au
2The first two authors contributed equally to this work.

**Synopsis**  Fish gastro-intestinal system harbors diverse microbiomes that affect the host’s digestion, nutrition, and immunity. Despite the great taxonomic diversity of fish, little is understood about fish microbiome and the factors that determine its structure and composition. Damselfish are important coral reef species that play pivotal roles in determining algae and coral population structures of reefs. Broadly, damselfish belong to either of two trophic guilds based on whether they are planktivorous or algae-farming. In this study, we used 16S rRNA gene sequencing to investigate the intestinal microbiome of 5 planktivorous and 5 algae-farming damselfish species (*Pomacentridae*) from the Great Barrier Reef. We detected *Gammaproteobacteria* ASVs belonging to the genus *Actinobacillus* in 80% of sampled individuals across the 2 trophic guilds, thus, bacteria in this genus can be considered possible core members of pomacentrid microbiomes. Algae-farming damselfish had greater bacterial alpha-diversity, a more diverse core microbiome and shared 35 ± 22 ASVs, whereas planktivorous species shared 7 ± 3 ASVs. Our data also highlight differences in microbiomes associated with both trophic guilds. For instance, algae-farming damselfish were enriched in *Pasteurellaceae*, whilst planktivorous damselfish in *Vibrionaceae*. Finally, we show shifts in bacterial community composition along the intestines. ASVs associated with the classes *Bacteroidia*, *Clostridia*, and *Mollicutes* bacteria were predominant in the anterior intestinal regions while *Gammaproteobacteria* abundance was higher in the stomach. Our results suggest that the richness of the intestinal bacterial communities of damselfish reflects host species diet and trophic guild.

**Brazilian Portuguese**  O sistema gastro-intestinal de peixes abriga microbiomas diversos que afetam a digestão, nutrição e imunidade do hospedeiro. Apesar da grande diversidade taxonômica dos peixes, entende-se pouco sobre o microbioma dos peixes e fatores que determinam sua estrutura e composição. Peixes-donzela são espécies importantes em recifes de coral que exercem papéis pivotais na determinação da estrutura de algas e corais dos recifes. De forma geral, peixes-donzela pertencem à uma de duas guildas tróficas dependendo se são plantívoros ou algívoros. Nesse estudo, usamos sequenciamento do gene 16S rRNA para investigar o microbioma intestinal de cinco espécies plantívoras e cinco espécies algívoras de peixes-donzela (*Pomacentridae*) da Grande Barreira de Corais. Detectamos ASVs de *Gammaproteobacteria* pertencendo ao gênero *Actinobacillus* em 80% dos indivíduos amostrados nas duas guildas tróficas, logo, bactérias desse gênero podem ser consideradas como possíveis membros essenciais do microbioma dos pomacentrídeos. Peixes-donzela algívoros apresentaram uma maior alpha-diversidade bacteriana, um microbioma essencial mais diverso e compartilharam 35 ± 22 ASVs, e espécies plantívoras compartilharam 7 ± 3 ASVs. Nossos dados também ilustram diferenças nos microbiomas associados com ambas guildas tróficas. Por exemplo, peixes-donzela algívoros estavam enriquecidos em *Pasteurellaceae*, enquanto peixes-donzela plantívoros, em *Vibrionaceae*. Finalmente, demonstramos mudanças na composição da comunidade bacteriana associada com as classes
Bacteroidia, Clostridia e Mollicutes foram predominantes nas regiões intestinais anteriores enquanto a abundância de Gammaproteobacteria foi maior no estômago. Nossos resultados sugerem que a riqueza das comunidades bacterianas intestinais de peixes-donze-ralentia reflete a dieta da espécie do hospedeiro, bem como a sua guilda trófica.

Chinese 鱼类肠道中种类的微生物菌群对于鱼类的消化、营养及免疫都有影响。尽管鱼类的分类多样性很高，但对于我们鱼类内的微生物群能够影响其结构和组成的原因却知之甚少。鳃囊科鱼类是一种重要的珊瑚礁鱼类，并且对珊瑚礁的藻类和珊瑚群落结构起到关键性作用。概括来说，基于食性的不同（以浮游生物为食或以藻类为食），鳃囊科鱼类分属于两种摄食类群。在本研究中，我们利用16S rRNA基因序列对于大堡礁的五种以浮游生物为食的鲷科和五种以藻类为食的鲷科分别进行了研究。在这两种类群的样本中，我们发现了属于放线菌属Actinobacillus的Gammaproteobacteria的扩增子序列变体。因此，此属细菌可能是鲷科肠道微生物群的主要组成。食藻类鲷科具有更高的细菌多样性，它们的核心微生物菌群的多样性更高，共享了35±22个扩增子序列变体，而食浮游生物类鲷科的核心微生物菌群则只共享了7±3个扩增子序列变体。我们的数据还突出了两种营养类群肠道微生物的区别。例如，食藻类鲷科有更多的Pasteurellaceae，而食浮游生物类鲷科则有更多的Vibrionaceae。最后，我们还展示了肠道中细菌群落的更替。在肠道前端，Bacteroidia, Clostridia和Mollicutes占据主导地位；而在胃中，Gammaproteobacteria则丰度更高。我们的结果意味着肠道菌群的丰富性反映了鱼类宿主的食性和摄食类群。

Hindi मछली गैस्ट्रो-आंतुर परिवार में विविधता जीवाणु होते हैं जो मेजबान के पाचन, पोषण और पुर्तांकण को पुर्वाधारित करते हैं। मछली के उच्च बर्फीलकण विविधता के बावजूद, मछली से जुड़े जीवाणु और उनकी संरचना और संरचना का निर्धारण करने वाले कारकों के बारे में बहुत कम समझ है। प्रमसेतुकण महत्त्वपूर्ण पूर्वाधारित विविधता की पूर्वजातियों हैं जो मछली और पुर्वाधारित विविधता की जनसंख्या संरचनाओं का निर्धारण करने में महत्त्वपूर्ण भूमिका निभाती हैं। कोई पौर्तरिकण पर, प्रमसेतुकण दो ट्रॉफिक कण गलियों में से कम से कम है, जो इस आधार पर है कुलवटक या मछली-बेही। इस अध्ययन में, हमने गर्दन बैचरिंग रिफ्र रन बैचरिंग रिफ्र डो पूर्वाधारित और पूर्व शौक बेही मछली प्रमसेतुकण (पोमासेटुवेड) के अंतों के जीवाणु की जांच के लिए 16S rRNA जीन अनुकूलन का उपयोग किया। हमने गामापूर्वतकेंट रिडिया एएसवी का पता लगाया, जो दो ट्रॉफिक कण गलियों में 80% सैपल नमूनों में जीनों एक्स्ट्रनियैलिमिन रूप से संबंधित थे। इस पर्यावरण, इस जीनों में बैचरिंग रिडिया को पोमासेटुवेड माइक्रोबायोम के संभावित युक्त समस्या माना जा सकता है। मछली-कूपी प्रमसेतुफल के अध्यक्ष कैटरिपियल अनुप्रयोग कण विविधता, एक अध्यक्ष विविधता कोर माइक्रोबायोम और 35±22 एएसवी साझा है, जबकि पुर्तांकण की पूर्वजातियों ने 7±3 एएसवी साझा था। हमारा डेटा दोनों ट्रॉफिक कण गलियों में जुड़े माइक्रोबायोम में आंत को भी उजागर करता है। उद्धरण के लिए, शौक-बेही करने वाले डेटमशी एल्पीएलासी अध्यक्ष होता है, जबकि पूर्वतकेंट रिडिया में वृद्धिकोशीघ्र होता है। अंत में, हम आंतों के साथ जीवाणु समुदाय संरचना में गतिविधियों को दिखाता है। बैचरिंग रिडिया, क्लोस्ट्रिडिया और मॉलस्क्यूम्बम बैचरिंग रिडिया के जुड़े एएसवी पूर्ववर्ती आंतों के क्षेत्रों में पूर्वभाग थे, जबकि गामापूर्वतकेंट रिडिया पेट में धूरुत मात्रा में थे। हमारे परिणाम संबंधित है कि प्रमसेतुफल के आंतों के जीवाणु समुदायों की समुदाय भेंट्वरान पूर्वजातियों के आधार पर ट्रॉफिक गलियों को दर्शाता है।

Italian Il sistema gastrointestinale dei pesci ospita un microbiota che influenza la digestione, nutrizione e sistema immunitario dell’ospite. Nonostante l’enorme diversità taxonomico dei pesci, la nostra comprensione del microbiota di questi animali ed i fattori che determinano la sua struttura e composizione è ancora scarsa. I pesci damigella includono specie importanti per le barriere coralline che forniscono servizi in grado che influenzano la struttura delle popolazioni di alghe e coralli. In generale, i pesci damigella appartengono a due gruppi funzionali basati sul loro tipo di dieta, e vengono divisi in consumatori di plankton o alghe. In questo studio abbiamo sequenziato il gene 16S rRNA per investigare il microbiota intestinale di cinque
Intestinal microbiome richness of coral reef damselfishes

Background

Fishes represent the greatest taxonomic diversity of vertebrates, and despite our understanding of the importance of intestinal microbiota of terrestrial vertebrates, we still lack an understanding of fish microbiome diversity and functioning (Clements et al. 2014). Largely, fish microbiome studies have centered around species with commercial value, including trout, salmon, and carp (Wang et al. 2018). For example, gastrointestinal fish microbiomes are known to be important in intestinal cell proliferation (Rawls et al. 2004; Cheesman et al. 2011), nutrition (Ray et al. 2012; Clements et al. 2014), and immunity (Bates et al. 2006; Bates et al. 2007; Galindo-Villegas et al. 2012). These studies show that the intestines of fishes harbor a large abundance and diversity of bacteria (Nayak 2010) and the regulation of this diversity is important in the maintenance of host health through a complex set of microbe-microbe and microbe-host interactions (Neish 2009; Foster et al. 2017).

There are many factors that affect the structure of fish gastrointestinal microbiomes (Clements et al. 2014; Wang et al. 2018). These include host-related factors such as genetic attributes, size, age, sex (Bolnick et al. 2014; Li et al. 2016; Stephens et al. 2016), host phylogeny (Sullam et al. 2012; Li et al. 2014; Miyake et al. 2015), environmental factors (such as water quality) (Hagi et al. 2004; Sullam et al. 2012; Neuman et al. 2016), and host diet (Miyake et al. 2015; Neuman et al. 2016). Studies that investigated intestinal microbiome changes have mostly focused on the impact of fish foods on species of aquaculture importance (Ringo et al. 2006; Martin-Antonio et al. 2007), although a few studies have investigated wild fish populations (Miyake et al. 2015; Zhang et al. 2018). For instance, bacterial symbionts diversification in wild herbivorous surgeonfish intestines is thought to be an important driver of host niche-partitioning (Miyake et al. 2016; Ngugi et al. 2017), suggesting that intestinal microbiomes can influence the trophic ecology of coral reefs and facilitate resource partitioning in these hyper-diverse ecosystems. However, the involvement of intestinal bacteria in wild fish physiology remains largely unknown.

There is increasing evidence that herbivorous fishes have distinct microbiomes as compared to omnivorous and carnivorous fishes (Givens et al. 2015). Herbivorous and carnivorous diets are known to cause shifts in intestinal fish microbiomes; fishes with plant-based diets have intestinal microbiomes dominated by Firmicutes, such as Clostridium, while fishes with fat-based diets have microbiomes dominated by protease-producing Proteobacteria (Desai et al. 2012; Ingerslev et al. 2014; Liu et al. 2016). In addition, the diversity of herbivorous fish intestinal microbiomes is higher than omnivorous and carnivorous host species under similar environmental conditions (He et al. 2013), suggesting that host feeding behavior has a significant effect on fish intestinal microbiomes.

Damselﬁshes (Pomacentridae) are a diverse and abundant group of coral reef fishes (Cooper et al. 2009; Campbell et al. 2018), and they are among the most widely studied families (Choat 1991; Emslie et al. 2019). Broadly, damselfishes are grouped into either planktivorous or alga-farming trophic guilds, although some herbivorous species may also feed on zooplankton (Eurich et al. 2019). Planktivorous damselfishes play a key role in transferring energy from the plankton to higher tiers of the food chain, while alga-farming damselfishes influence sediment and algal dynamics on coral reefs and may increase the presence of coral disease-associated pathogens within their territories (Casey et al. 2015; Casey et al. 2015; Emslie et al. 2019; Randazzo Eisemann et al. 2019, Tebbett et al. 2020; Blanchette et al. 2019). Alga-farming species can be differentiated based on the algal composition within their territories, and they are divided into several behavioral guilds, including indeterminate grazers, extensive grazers, and intensive grazers (Hata and Kato 2004; Emslie et al. 2012; Casey et al. 2015;
Indeterminate and extensive grazers feed both on macroalgae and turf, while intensive grazers maintain distinct areas of turf algae through selective grazing and weeding of unpalatable algae (Gibson et al. 2001; Emslie et al. 2012). Intensive grazing damselfish are also referred to as algae farmers. Research on intensive grazers has focused on competition (Eurich et al. 2018), patterns of co-existence (Eurich et al. 2018; Eurich et al. 2019), behavioral interactions (Kasumyan 2009; Weimann et al. 2018), and their role in structuring algae and coral communities (Klumpp et al. 1987; Ceccarelli et al. 2005; Ceccarelli 2007; Gochfeld 2010; Casey et al. 2014; Casey et al. 2015).

In this study, we investigated and described the intestinal microbial diversity of ten species of planktivorous and algae-farming damselfishes, two guilds that significantly impact coral reef trophic dynamics. We hypothesized that differences in intestinal microbial communities will reflect the differences between these two trophic guilds. Specifically, across the different host species and trophic guilds, we examined (1) differences in bacterial communities across fish species and trophic guilds, (2) core microbial members, and (3) changes in microbial community structure along the length of the intestinal tract.

Methods
Species collections and dissections
Fishes were collected from the Heron Island lagoon in the southern Great Barrier Reef, Australia (23°26′53″S, 151°56′52″E) in January and February 2015. Collections occurred at a depth of 1–8 m adjacent to the Heron Island Research Station. Three individuals of ten sympatric damselfish species (Abudejuf sexfasciatus, A. whitleyi, Acanthochromis polyacanthus, A. polycanthus, Chromis atropectoralis, Dischistodus pseudochrysopoecilus, D. perspicillatus, Pomacentrus moluccensis, P. wardi, Stegastes apicalis, and S. nigricans) of similar lengths were randomly collected across the two trophic guilds planktivorous and algae-farming. Each trophic guild was represented by 5 species and 15 individuals. Collections were conducted on SCUBA, and the planktivorous species were collected using a barrier net, while the algae-farming species were collected using a speargun. Following collections, the fishes were immediately placed on ice and transported to Heron Island Research Station. In the laboratory under sterile conditions, fishes were weighed, measured and photographed, then the gastrointestinal tract was removed, and the gut length was recorded and photographed. The entire gut was fixed in 4% DNA/RNA free paraformaldehyde and sterile phosphate-buffered saline for 12 h, then it was stored in DNA/RNA free water.

DNA extraction, amplification, and sequencing
Samples were transported to James Cook University for subsampling along each intestinal tract and DNA extraction. Under sterile conditions, standardized biopsy cores (3 × 3 mm) were taken from four locations along the intestinal tract: the stomach, the anterior intestine, the mid-intestine, and the posterior intestine. DNA was extracted from tissue biopsies using a QIAamp DNA Micro Kit (Qiagen, Hilden, Germany) following the manufacturer’s guidelines. A nanodrop was used to record the quality (260/280 ratio) and quantity (ng/μL) of DNA from each extraction.

Amplification of the 16S V1-V3 rRNA gene region was done using the primers 27F (5′-AGGAGTTTGATCMTGGCT-3′) (Ludwig 2007) and 519R (5′-GTTACNGCGGCKGCTG-3′) (Lane et al. 1985) with barcodes on the forward primer. These 16S rRNA genes were amplified using the HotStarTaq Plus Master Mix Kit (Qiagen, USA) under the following conditions: 94°C for 3 min, followed by 28 cycles of 94°C for 30 s, 53°C for 40 s and 72°C for 1 min, after which a final elongation step at 72°C for 5 min was performed. After amplification, PCR products were checked in 2% agarose gel to determine the success of amplification and the relative intensity of bands. Multiple samples were pooled together (e.g., 100 samples) in equal proportions based on their molecular weight and DNA concentrations. Pooled samples were purified using calibrated Ampure XP beads. Then the pooled and purified PCR products were used to prepare a DNA library by following Illumina TruSeq DNA library preparation protocol. Sequencing was performed at the Molecular Research LP (MR DNA; Texas, USA) on a MiSeq V2 System following the manufacturer’s guidelines.

Amplicon sequence data were sorted by the sample and demultiplexed using demux for QIIME 2 (version 2018.11; (Bolyen et al. 2018,)). Sequences were screened for quality, trimmed at 450 bp after removal of primer sequences, and assigned as amplicon sequence variants (ASVs) using DADA2 (Callahan et al. 2016). Taxonomy of the ASVs was determined using a pre-trained, naïve Bayes classifier (Pedregosa F) and the q2-feature-classifier plugin (Bokulich et al. 2018). The classifier was trained on the target 480 bp region of sequences in the Greengenes 13.8 99% database. ASV clusters were arranged in a phylogenetic tree using FastTree (Price et al. 2010) and visualized using Interactive Tree of Life 3.6.1 (Letunic and Bork 2016). The feature table, metadata, and taxonomic classifications were exported from QIIME 2 in .biom format and the rooted phylogenetic tree

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was exported in .nwk format. The closest known sequences and the origin of selected ASVs were identified through a BLASTN-based search against the GenBank nr/nt database.

**Statistical analysis**

The feature table and phylogenetic tree were imported into R version 3.5.2 and stored as a *phyloseq* object (McMurdie and Holmes 2013) for downstream analyses. All ASVs not assigned to phylum were filtered from the data, and those designated as chloroplasts or cyanobacteria were removed and stored as a separate object for further analysis. Samples were rarefied to minimum sampling depth for alpha-diversity analyses, which was estimated using the R package vegan (Oksanen et al. 2017). Non-rarefied data were used for generalized linear model (GLM) analysis (McMurdie and Holmes 2014; McMurdie 2018). Data used for principal component analysis (PCA), betadisper-test and PERMANOVA were computed using centered log-transformed Euclidean distance matrices of the non-rarefied ASV table. Differences in alpha-diversity between trophic guilds were tested via $t$-test. Multivariate GLM was used to test for significant differences in bacterial communities among host fish species, trophic guild, and location along intestines using mvabund in R (Wang et al. 2012). PCA, betadisper-test and PERMANOVA were used to test differences in the communities of *Proteobacteria*, *Bacteroidetes*, and *Firmicutes* among fish species and between the two trophic guilds. Bacterial taxa were grouped by class when examining microbiome changes along the length of the intestinal tract. Bacterial community data were fitted to negative binomial distributions and tested using log-likelihood ratios (LRT) via 999 simulations using Monte Carlo resampling. A nested analysis of variance (ANOVA) used to test the role of trophic guild and gut location when accounting for species variation. Venn diagrams were produced using the *VennDiagram* package (Chen and Boutros 2011).

**Results**

A total of 1,254,909 sequences were detected in 119 samples after denoising and removing all chloroplast, mitochondria, and uncharacterized sequences. Among these sequences, 3,776 ASVs were detected; 39.4% of which belonged to the phyla *Proteobacteria*, 26.2% to *Bacteroidetes*, 13.4% to *Firmicutes*, and 12.6% to *Planctomycetes*. The 20 most abundant ASVs accounted for 41% of the total number of detected sequences. The most common ASV belonged to the genus *Actinobacillus* and accounted for 9.9% of the total detected sequences (Table 1). Two unknown species of *Mollicutes* and *Pasteurellaceae* accounted for 6.9 and 3.8% of sequences, respectively.

Different ASV richness was detected for each fish species with observed ASVs ($t = −3.15, P = <0.01$) and Shannon index ($t = −3.68, P = <0.01$) differing significantly between the two trophic guilds. The damselfish *D. perspicillatus* had the greatest mean richness of ASVs, with a total of $322 ± 17$ ASVs per individual (Fig. 1). The species with the lowest ASV richness were *C. atripectoris* and *A. sexfasciatus* with $47 ± 21$ ASVs.
and 30 ± 8 ASVs per individual, respectively (Fig. 1). Shannon diversity was greatest for two algae-farming species *D. perspicillatus* and *S. apicalis* and lowest for the planktivorous species *A. polyacanthus* and *P. moluccensis*. PCA biplots, betadisper-test, and PERMANOVA revealed that the beta-diversity of *Proteobacteria*, *Bacteroidetes*, and *Firmicutes* communities differed among fish species and trophic guilds (Fig. 2; Table 2).

### Core microbiomes

In line with previous studies that investigated the core microbiome of other organisms (Ainsworth et al. 2015; Ricci et al. 2022), we choose a minimum threshold of 30% for this metric. Most ASVs occurred in less than 30% of sampled individuals across all fish species (Fig. 3a). A total of 13 bacterial ASVs were found in more than 30% of sampled individuals; therefore, they
may represent the 30% core microbiome of pomacentrid investigated in this study (Table 3). The most common ASV in this study belonged to the genus *Actinobacillus*, which occurred in more than 80% of sampled individuals (Table 3), albeit at a low abundance in many individuals, with the highest abundances in the planktivorous damselfishes *A. polyacanthus* and *P. moluccensis*.

The core bacterial assemblages of each fish species (defined as ASVs that were shared between all sampled individuals for each species) were composed of a variable number of ASVs (Fig. 3b). For example, there were 70 bacterial ASVs shared between the three sampled individuals of *D. perspicillatus* and only two ASVs shared between the three *A. sexfasciatus* individuals. Core microbiomes within fish species were richer in...
algae-farming species than planktivorous species (Fig. 3b), with algae-farming species sharing 35 ± 22 ASVs and planktivorous species sharing only 7 ± 3 ASVs (Wilcoxon test W = 25, p < 0.01).

Core ASVs that occurred in all three individuals of a fish species belonged to the phyla Bacteroidetes, Firmicutes, Tenericutes, Spirochaetes, Planctomycetes, Proteobacteria, and Verrucomicrobia. Core ASVs belonging to Cor alienomargarita sp. and Verruco-5 (Verru-comicrobia), Pirellulaceae (Planctomycetes), and Desulfovibrio-1onaceae (Deltaproteobacteria) occurred in all three sampled D. perspicillatus individuals (Supplementary Figure S1). We also detected high diversity of an unknown clade of Gammaproteobacteria in P. moluccensis and P. wardi damselfish. There were 61 core ASVs belonging to the Bacteroidetes, 28 of which occur in S. apicalis and 38 in D. perspicillatus (Supplementary Figure S2). An unknown clade of Flavobacteriales and a diverse consortium of Rikenellaceae were core members of S. apicalis, while D. perspicillatus had a diverse core assemblage of ASVs belonging to the family Flavobacteriaceae. One ASV belonging to Spirochaetes, Brevinem a andersonii, was a core member of S. nigricans and C. atriporalis, while a Tenericutes ASV belonging to Mollicutes was a core member of all fish species except the planktivorous damselfishes A. polyacanthus and A. sexfasciatus (Supplementary Figure S3). There was a rich consortium of core Firmicutes ASVs for S. apicales and S. nigricans, which included members of the Erysipelotrichaceae, Ruminococcaceae, and Lachnospiraceae families.

### Table 2. Results of betadisper-test and PERMANOVA testing the beta-diversity of Proteobacteria, Bacteroidetes, and Firmicutes communities across fish species and between trophic guilds.

| Fish species | betadisper | PERMANOVA | Trophic guild | betadisper | PERMANOVA |
|--------------|------------|-----------|---------------|------------|-----------|
| Proteobacteria | P = 0.084 | F = 1.86; p = 0.001*** | p = 0.039* | F = 3.52; p = 0.001*** |
| Bacteroidetes | P = 0.269 | F = 1.78; p = 0.001*** | p = 0.233 | F = 2.41; p = 0.001*** |
| Firmicutes | P = 0.001*** | F = 2.17; p = 0.001*** | p = 0.355 | F = 3.92; p = 0.001*** |

Bacterial shifts along the intestinal tract

The interaction between the trophic guild and intestinal region had a significant influence on the gut bacterial community composition (LRT = 152, P = 0.001; Supplementary Table 1). The abundance of nine classes of bacteria changed significantly across the different fish species and locations along the intestinal tract (LRT = −0.0229, P < 0.001; Fig. 4; Supplementary Table 2). Members of Gammaproteobacteria were especially common throughout the planktivorous intestinal tracts, but we also found them along all the intestines regions of the algae-farming species D. perspicillatus, D. pseudochrysopeocilus, and P. wardi (Fig. 4). In intestinal regions where Gammaproteobacteria were uncommon, members of Bacteroidia and Clostridia were generally found at higher abundances—especially for algae-farming species (Fig. 4). Members of the Mollicutes and Planctomycetes were more common throughout the intestinal tracts of algae-farming hosts than planktivorous species although their abundances were generally lowest within the stomach region (Fig. 4). The stomach had 286 unique bacterial ASVs, the anterior intestine 753, while 1,139 and 656 ASVs were only found in the mid and posterior intestines, respectively (Fig. 5). Only 19 ASVs were common in the stomach and posterior intestine while 152 ASVs were found throughout the intestine (Fig. 5).

**Effect of the trophic guild on microbiomes**

There was a significant difference in the microbiome composition between trophic guilds (LRT = −0.021, P < 0.001; Supplementary Table 2). Most bacterial ASVs were unique to either of the trophic guilds, with only 124 ASVs common to both guilds (Fig. 5). A total of 78 bacterial ASVs, belonging to 20 families, were important drivers of this relationship. There were marked differences in abundances of ASVs belonging to Vibrio-2onaceae, Lachnospiraceae, and Pasteurellaceae. Two Vibrio sp. (Vibrionaceae) were more common in planktivorous species, and five ASVs of Actinobacillus (Pasteurellaceae) were more abundant in algae-farming species.

**Discussion**

Our data show that algae-farming damselfish species have richer microbiomes than planktivorous species (Fig. 1) and this result is also reflected in their core bacterial community (Fig. 3). This result is likely attributable to the specialized feeding behavior of algae-farming species, which largely consume a narrow range of turf algae species (Hata and Kato 2004; Casey et al. 2014), unlike planktivorous species that are adapted to a more opportunistic feeding strategy. These results
suggest that the microbiome structure of fish species with specialized feeding behavior has acquired specific intestinal bacteria and further research is needed to investigate how microbiome specialization affects host digestion and metabolism. We also note that other processes that were not tested in our study such as host phylogeny and functional traits could influence the composition of damselfish intestinal bacteria and ultimately influence fish physiology.

We found that similar to what was recorded in many other species of marine fish, the damselfish intestinal microbiome was dominated by members of *Proteobacteria, Bacteroidetes, Firmicutes, and Planctomycetes* (Table 1). For example, surgeonfish, parrotfish,
Table 3  Taxonomic composition of core ASVs occurring in more than 80% of sampled individuals. Accession numbers for closest GenBank sequences (similarity given in brackets) are supplied. Occurrence and relative abundances were generated from rarefied data.

| ASV  | Phylum          | Lowest taxonomic division   | Occurrence (%) | Relative abundance | GenBank accession number |
|------|----------------|-----------------------------|----------------|-------------------|--------------------------|
| b727 | Proteobacteria | Actinobacillus sp.          | 83.3           | 0.083             | KT952745 (97.5%)         |
| 94ba | Proteobacteria | Actinobacillus sp.          | 53.3           | 0.017             | KT952745 (93.5%)         |
| 9bd9 | Proteobacteria | Photobacterium damselae     | 43.3           | 0.013             | CP035457 (100%)          |
| 5647 | Tenericutes    | Malicutes                   | 40.0           | 0.022             | HG971018 (96.3%)         |
| a382 | Proteobacteria | Photobacterium damselae     | 40.0           | 0.008             | CP018297 (100%)          |
| 73d1 | Proteobacteria | Vibrio sp.                  | 40.0           | 0.010             | KT952854 (98.7%)         |
| 9b2f | Proteobacteria | Actinobacillus porcinus     | 40.0           | 0.018             | KT952745 (91.9%)         |
| 6c33 | Proteobacteria | Spirobacillales             | 37.7           | 0.002             | KU578602 (100%)          |
| dc1c | Proteobacteria | Vibrio sp.                  | 37.7           | 0.004             | CP033144 (100%)          |
| 5a8a | Proteobacteria | Vibrio parvus               | 37.7           | 0.019             | MG524941 (100%)          |
| 762a | Bacteroidetes  | Lutimonas sp.               | 30.0           | 0.001             | MG488523 (99.6%)         |
| ca47 | Proteobacteria | Vibrio Harveyi              | 30.0           | 0.009             | CP033144 (100%)          |
| 6013 | Proteobacteria | Pasteurellaceae             | 30.0           | 0.007             | KT952745 (92.3%)         |

and rabbitfish intestinal microbiomes from the Red Sea also consist of diverse assemblages of *Firmicutes* and *Proteobacteria* (Miyake et al. 2015). Another dominant ASV in the damselfish microbiome belonging to *Mollicutes* (*Tenericutes*) resembled bacteria detected in rabbitfish intestines (Zhang et al. 2018). The number of highly similar bacterial ASVs shared among pomacentrids, acanthurids, and siganids may reflect the similar feeding behaviors of these coral reef fishes. For instance, algae-farming damselfishes may also ingest prey items other than algae, such as zooplankton (Erlich et al. 2019) or other invertebrates (Letourneau et al. 1997). The functional roles of these seemingly important microbial taxa warrant further attention in order to understand the potential consequences on host metabolism and health.

Damselfish microbiomes were largely dominated by the family *Pasteurellaceae* in the phylum *Gammaproteobacteria*, with one ASV (b727) occurring in more than 80% of sampled fishes and representing almost 10% of the total detected sequences (Tables 1 and 3). Although this ASV currently represents an unknown species in the *Actinobacillus* genus, a 98% similar sequence has been retrieved from the intestines of surgeonfishes in Saudi Arabia (Miyake et al. 2016), suggesting that *Actinobacillus* are common members of reef fish microbiomes. Bacteria in the genus *Pasteurellaceae* have also been recorded in high abundances in adult damselfishes and cardinalfishes collected around Lizard Island, Australia (Parris et al. 2016), and they are deemed as common components of tropical planktivorous fish gut microbiomes (Egerton et al. 2018). The prevalence of *Pasteurellaceae* amongst the damselfishes in this study, as well as in other reef fishes, provides additional evidence that *Pasteurellaceae* are likely important members of coral reef-associated fish microbiomes.

Algae-farming damselfishes had more observed ASVs and larger core microbiomes than planktivorous species (Figs. 1 and 3), and these core microbiomes were specific to each host species (Fig. 3). For example, *P. wardi* and *P. moluccensis* microbiomes were dominated by different taxa of *Gammaproteobacteria*, while *D. perspicillatus* and *S. apicularis* had large *Bacteroidia* core communities but were dominated by *Flavobacteriaceae* and *Rikenellaceae*, respectively. Different species of algae-farming damselfishes consume different species of algae (Casey et al. 2014), and the large differences in their specialized microbiomes may reflect these narrow dietary preferences. Conversely, the small core microbiomes of the planktivorous damselfishes may reflect the high variation in consumed plankton of each species, suggesting these fishes have opportunistic feeding behaviors. These results, however, do not support the notion that fish with greater diet variability have more diverse microbiomes (Givens et al. 2015). In fact, the damselfish with narrow, algae-farming feeding behaviors tended to have the greatest diversity of intestinal bacteria, suggesting that the host-microbiome interactions may select for specialized bacteria that enhance the digestion and absorption of nutrients from specific algal diets. The richer microbiome of algae-farming fishes could also reflect the necessity
Intestinal microbiome richness of coral reef damselfishes

Intestinal microbiome richness of coral reef damselfishes

**Fig. 4** Changes in abundance of selected bacterial Classes along the four locations along the intestine of each species of damselfish as determined by nested multivariate generalized linear models. Intestinal locations include stomach (S), anterior intestine (AI), mid-intestine (MI), and the posterior intestine (PI). The top row represents planktivorous species and bottom row represent algae-farming species.

**Fig. 5** Venn diagrams depicting the number of shared ASVs for each trophic guild (left) and for each region of the intestine (right).

of this trophic guild to be associated with a pool of symbionts that facilitate the breakdown of algal cellu-
lose. We also acknowledge that some of the bacteria we retrieved from the damselfish intestine could have been associated with the food recently ingested by the fish and, therefore, not being part of the damselfish microbiome.

Evidence suggests a high degree of resource partitioning in fish communities, which is a key mecha-
nism that facilitates the high diversity of coral reefs
(Casey et al. 2019; Leray et al. 2019). The largely distinct microbiomes of each host species presented in this study may reflect the high degree of resource partitioning found in coral reef communities, whereby different species of damselfish may be consuming different size classes of zooplankton (Leray et al. 2019), farm different algal species (Casey et al. 2014), or occupy different trophic niches (Casey et al. 2019). The similarity between closely related host species and microbiomes, such as *P. wardii* and *P. moluccensis*, also demonstrates that phylogeny may influence the intestinal microbiomes of damselfishes (Sullam et al. 2012; Miyake et al. 2015; Neuman et al. 2016; Chiarello et al. 2018).

Interestingly, *Photobacterium damselae*, *Vibrio Harveyi*, *Vibrio ponticus*, and other *Vibrio* sp. were prevalent amongst the damselfishes sampled in this study (Table 3). These bacteria represent potential pathogenic members of *Vibrio acanitae* and have been detected in many fishes of aquaculture importance, including *Chromis punctipinnis* (Love et al. 1981), *Lutjanus argentinaculatus* (Reshma et al. 2018), *Seriola dumerili* (Nishiki et al. 2018), *Scophthalmus maximus* (Montes et al. 2003), *Sparus aurata* (Vera 1991), and *Solea senegalensis* (Terceti et al. 2016). Although identified as *Vibrio Harveyi* in the GreenGenes database, Gen-Bank revealed there was a high similarity of these sequences to other members of the *Harveyi* clade, such as *Vibrio owensii* (Nishiki et al. 2018). It is thought that there are up to 11 species of *Vibrio* belonging to this clade (Urbanczky et al. 2013), most of which are pathogens of fish, shrimp, and coral (Thompson et al. 2004; Austin and Zhang 2006; Ushijima et al. 2012). Given the apparently healthy state of the sampled fishes and the high abundances of potentially pathogenic *Vibrio acanitae* in the fish guts, we provide support to the idea that these organisms are natural components of healthy fish microbiomes and are opportunistic pathogens in fishes only under specific conditions (Rivas et al. 2013; Reshma et al. 2018). Future studies should also investigate the involvement of algae-farming damselfish in the spreading of pathogens across reef organisms. For instance, it has recently been reported that the seagrass pathogen *Labyrinthula* was present in the skeleton of a common coral species (Ricci et al. 2021) and probably infected the abundant endolithic algae living in the coral skeleton (Ricci et al. 2019; Iha et al. 2020; Tandon et al. 2022; Ricci et al. 2022). Thus, it is possible that damselfishes grazing near alive corals were the medium that allowed the pathogen *Labyrinthula* to infect the corals’ endolithic algae.

The facultative anaerobic bacterial classes *Bacteroidia*, *Clostridia*, and *Mollicutes* were generally in higher abundance in the mid and posterior intestinal regions than in the stomach (Fig. 4). Differences in microbiomes along the intestinal tract have been recorded in the rabbitfish *Siganus fuscensce* (Nielsen et al. 2017), with midgut communities more representative of the environmental sources and hindguts hosting a microbiome more specialized to anaerobic conditions and fermentation (Jones et al. 2018). The increase in *Bacteroidia*, *Clostridia*, and *Mollicutes* along the intestines may be due to some members of these bacterial classes being mutualistic components of the fish gastrointestinal microbiome. Some members of *Bacteroidia* are known to breakdown polysaccharides and metabolize the derived sugars (Xu et al. 2003), while members of *Clostridium* are known to metabolize cellulose (Liu et al. 2016). Our results confirm the increased prevalence of anaerobic bacteria in the hindgut of damselfishes, which probably consists of taxa responsible for the fermentation and metabolism of complex molecules before being absorbed by the host (Clements et al. 2014). We also note that *Actinobacillus* sp. that could breakdown cellulose via fermentation (Almquist et al. 2016) were more abundant in the gut of algae-farming damselfish, suggesting that these bacteria could aid the digestion of fish in this trophic guild.

**Conclusions**

In this study, we show that damselfishes have diverse intestinal microbial communities whereby the bacterial richness of a species reflects diet and trophic guild. We show that algae-farming damselfishes have richer bacterial alpha-diversity and core microbiomes, which may reflect the more specialized diets of this trophic guild. We also provide evidence that damselfish mid and posterior intestines have higher abundances of facultative anaerobic bacteria that are known to play important roles in fermentation and cellulose breakdown. These findings add to a growing body of literature that suggests that host fish feeding behavior has a strong influence on the composition of intestinal microbiomes.

**Supplementary data**

Supplementary Data available at *IOB* online.

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The authors declare that they have no competing interests.

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Consent for publication
Not applicable.

Availability of data and material
The Illumina MiSeq datasets for each damselfish species are available at the Sequence Read Archive (NCBI) repository under BioProject accession number PRJNA638998, https://www.ncbi.nlm.nih.gov/sra. Data and R-scripts used in this study are available at https://github.com/ChrisKav/WildDamselfishMicrobiomes.

Authors’ contributions
CRJK analysed and interpreted the amplicon sequence data and was the major contributor to writing the manuscript and preparing figures and tables. FR provided feedback on the data analysis, figures design, and manuscript writing. JMC undertook the fieldwork and collected all specimens, performed gut dissections, tissue biopsies, and provided feedback on the manuscript. JHC was involved with the initial synthesis and design of this study and provided feedback on the manuscript. WL and TDA were involved with the initial synthesis and design of this study, provided the facilities to undertake laboratory work and provided feedback on the manuscript. All authors read and approved the final manuscript.

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