The gut bacterial microbiota of sea turtles differs between geographically distinct populations

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ABSTRACT: The microbiota of metazoans can be influenced by a variety of factors including diet, environment and genetics. In this study we sampled multiple populations from 2 host species that do not overlap in distribution, in order to test whether their bacterial microbiotas are species-specific or more variable. Intestinal swabs were collected from loggerhead turtles originating from Florida, USA, and Queensland, Australia, as well as from flatback turtles from Crab Island, Queensland, and Port Hedland, Western Australia. We then manually extracted bacterial DNA and used 16S rRNA sequencing to explore bacterial microbial community composition and structure. Our investigation showed that the bacterial microbiota of sea turtles is heavily influenced by geography, with loggerhead turtles originating from the USA and Australia harbouring significantly different bacterial microbial populations in terms of composition. Similarly, we also found that flatback turtles from Crab Island had significantly less diverse microbiotas, with a predominance of the bacterial phylum Firmicutes, in comparison to their genetically similar counterparts from Port Hedland. Factors that may explain these observed differences between populations include host genetics, differences in foraging habitat quality and differences in migratory distance (and thus durations of inappetence) between foraging and breeding grounds. The mechanisms by which these factors may influence bacterial microbial composition of sea turtle gastrointestinal tracts warrants further investigation. The results of this study highlight the importance of interpreting microbiota data of wild animals in the context of geography.

KEY WORDS: Microbiota · Flatback turtle · Loggerhead turtle · Geography

1. INTRODUCTION

The composition of the gastrointestinal microbiota is determined by the same ecological principles that drive diversity and niche occupancy within macro-ecosystems. That is, colonist assemblages are established by founder populations, dispersal, diversification, local abundance, ecological drift, and habitat quality (Costello et al. 2012, Koskella et al. 2017). These are further influenced by individual bacterial traits such as dispersal efficiency and mutation rates, as well as host genetics (Costello et al. 2012, Coyte et al. 2015). Thus, the final bacterial community of an individual is determined by a complex interplay of local and external factors, with microorganisms competing to establish stable populations within the gastrointestinal environment (Costello et al. 2012, Coyte et al. 2015).

In humans, diet strongly influences the composition of gut bacterial microbiota (Albenberg & Wu 2014, David et al. 2014, Voreades et al. 2014). Transient fluctuations in community structure may
become evident in as little as 24 h after consumption of novel dietary items (David et al. 2014). Such plasticity may confer teleological advantages by permitting human dietary and lifestyle diversity (David et al. 2014), but it may also be a contributing factor in the development of disease (Albenberg & Wu 2014). Genetics are also a strong driver of microbiota composition in humans, with potential implications for host metabolism (Spor et al. 2011, Goodrich et al. 2014).

As in humans, the gut microbiota of animals may be influenced by factors such as genetics (Zhao et al. 2013) and diet (Nelson et al. 2013, Frese et al. 2015, Henderson et al. 2015). Furthermore, disturbances in habitat quality appear to have species-specific effects on microbial populations of animals. Black howler monkeys Alouatta pigra inhabiting suboptimal environments consumed less diverse diets and had a corresponding decrease in gut microbial diversity. Additionally, genetic analysis of microbes isolated from these animals revealed a reduction in genes associated with butyrate production and hydrogen metabolism, which may have implications for host health, because the animals are unable to use environmental resources as efficiently as animals from pristine habitats (Amato et al. 2013). Conversely, a study involving multiple primate species demonstrated that habitat perturbation did not influence fecal microbiotas and that differences observed were explained by differences in digestive physiology and dietary preferences (McCord et al. 2014). However, in this same study, differences were noted between populations of the same species from different countries, suggesting that geography and social group membership influence microbial communities in some primates (McCord et al. 2014).

Despite the apparent malleability of the microbiota, investigations in mice have revealed that populations harbor a core microbial community, which is subject to host genetic control (Benson et al. 2010, Zhang et al. 2010). The relative abundances of taxa within this core microbiota collectively behave as a complex polygenetic trait and they may exert their effects on other taxonomic groups and subsequently affect their ability to establish within the gastrointestinal tract (Benson et al. 2010). This multifaceted relationship between microbes and their hosts may be one of the key drivers of evolution and account for some of the enormous diversity we see in form and function of metazoans (Fraune & Bosch 2010). However, in order to understand the importance of microbes in driving host evolution, investigations of a wider variety of host taxa are needed.

In comparison to other vertebrate classes, reptiles are grossly underrepresented in bacterial microbiota studies; however, in recent times, attention has started to focus on describing the bacterial community composition of this evolutionarily significant taxon (Hong et al. 2011, 2015, Keenan et al. 2013, McLaughlin et al. 2015, Yuan et al. 2015, Ren et al. 2016, Arizza et al. 2019, Holmes et al. 2019, Lin et al. 2019, Tang et al. 2019, Willson et al. 2019, Zhang et al. 2019, Fugate et al. 2020). Like other vertebrates, the bacterial microbiota of reptiles may be influenced by intrinsic and extrinsic factors such as host lineage (Ren et al. 2016), diet (Jiang et al. 2017, Holmes et al. 2019, Bloodgood et al. 2020, Montoya-Ciriacono et al. 2020), anorexia (Lin et al. 2019), food consumption (Costello et al. 2010), habitat (Littleford-Colquhoun et al. 2019), sex (Martin et al. 2010), climate change (Beston et al. 2017), captivity (Kohl et al. 2017, García-De La Peña et al. 2019), exposure to chemicals (Indest et al. 2018), and gestation (Trevelline et al. 2019). In the largest investigation into sea turtle microbiotas conducted to date, microbial communities were strongly linked to sea turtle phylogeny, through a process of co-evolution between gut microbial inhabitants and the host (Scheelings et al. 2020a). While microbiota composition is strongly linked to health in humans (Shreiner et al. 2015), establishing these same relationships in sea turtles is difficult, with no correlation found between gut microbiota samples and a range of blood parameters in flatback turtles Natator depressus (Scheelings et al. 2020b). Exposure to glyphosphate-based herbicides has shown to reduce growth and survival of selected gut bacterial isolates from green turtles Chelonia mydas, with potential negative implications for gut physiology and health (Kittle et al. 2018). Also, in green turtles, an ontogenetic shift in microbial composition has been observed when animals transition from a pelagic lifestyle as juveniles to a neritic habitat as subadults (Price et al. 2017, Campos et al. 2018). Differences in microbiota composition of juvenile green turtles have been shown to exist between animals originating from different localities (Ahasan et al. 2017, Price et al. 2017, Campos et al. 2018); however, these investigations were conducted separately rather than as a single study aiming to determine the effects of geography on bacterial community structure. Similarly, the bacterial microbiota of loggerhead turtles Caretta caretta, seems to vary markedly between studies (Abdelrhman et al. 2016, Arizza et al. 2019, Biagi et al. 2019, Scheelings et al. 2020a), despite the fact that most of the animals from these investigations originated from the same
genetic stock in the Mediterranean Sea (Pritchard 1997, Duchene et al. 2012). Understanding the microbiota, and how it has shaped vertebrate life, is particularly important for less-studied species, such as the Reptilia, in order to gain a complete understanding of the evolutionary processes driving host–microbe relationships. To address this deficit in the reptile literature, we chose to explore the potential associations that exist between geography and bacterial gut microbiota in sea turtles.

There are 7 extant species of sea turtles, which may occupy a diverse array of ecological niches and employ a variety of foraging strategies (Bowen & Karl 2007). The global phylogeography of sea turtles is markedly influenced by habitat preference and thermal tolerance (Bowen & Karl 2007). For example, the more cold-adapted leatherback turtles *Dermochelys coriacea* show the lowest geographic structuring as their large size and pelagic lifestyle permits them to traverse through the boreal waters that act as a barrier to other warm-water adapted turtles (Bowen & Karl 2007). Conversely, tropical species, such as green turtles, hawksbill *Eretmochelys imbricata* turtles, and ridley turtles (*Lepidochelys olivacea* and *L. kempii*) have ancient separations between Atlantic and Indo-Pacific populations caused by the closure of the Isthmus of Panama around 3.5 million years ago (Bowen & Karl 2007). Within sea turtle species, there may be genetically distinct populations living within the same broad geographic regions, as well as genetically distinct populations living in very separate geographic regions. Alternatively, there may also be limited genetic variation of some species living in separate populations but within the same broad geographic area. For example, loggerhead turtles that nest on the east coast of Australia are genetically distinct from those that nest on the west coast of Australia (Limpus 2008b), and both of these populations are distinct from animals that nest in Florida, USA (Bowen & Karl 2007). In contrast, there is very little genetic diversity among flatback turtles, despite there being 4 distinct nesting populations within Australia, with no inter-population breeding (Limpus 2008a). These phylogeographic traits make sea turtles interesting subjects to study in the context of bacterial microbiotas and to explore the influence of genetic and environmental factors on microbial community structures. The aims of this investigation were to explore whether variations in microbiota can be seen between different groups of the same species of sea turtles and to identify possible explanations for any observed disparities. To do this, we investigated how microbiotas differed among populations of loggerhead turtles in Australia and the USA, as well as between different populations of flatback turtles within Australia.

2. MATERIALS AND METHODS

2.1. Study populations

Nesting adult flatback turtles of unknown age were sampled from Crab Island (n = 20), Queensland (10.972° S, 142.106° E), in September 2016, and from Port Hedland (n = 17), Western Australia (20.3107° S, 118.5878° E), in November 2016. Animals from these 2 areas are considered to be from genetically separate populations with little gene flow between them (Limpus 2008a). Loggerhead turtles were sampled from Mon Repos (n = 20), Queensland (24.8059° S, 152.4416° E), in January 2017, and from Juno Beach (n = 20), Florida, USA (26.5224° N, 80.315° E), in May 2018. These 2 populations of loggerhead turtles represent distinct genetic stocks (Duchene et al. 2012).

2.2. Sample collection

If turtles were nesting, we waited until they had finished laying and were returning to the ocean. We then flipped them into dorsal recumbency, and an equine uterine swab (Minitube) was inserted into the cloaca so that it entered the distal colon. These swabs are approximately 90 cm in length, and each swab was inserted to a depth of at least 60 cm. Based on previous research conducted by our research group in which endoscopy was used to visualise the cloacal structures of sea turtles (Rafferty et al. 2013), we are confident that inserting the swabs to this depth permitted sampling of the distal colon and not the cloaca. These swabs were approximately 90 cm in length, and each swab was inserted to a depth of at least 60 cm. Based on previous research conducted by our research group in which endoscopy was used to visualise the cloacal structures of sea turtles (Rafferty et al. 2013), we are confident that inserting the swabs to this depth permitted sampling of the distal colon and not the cloaca. These swabs were housed in a sterile sheath, the entire apparatus was inserted into the cloaca, and the swab tip was extruded when correct placement of the sheath had been achieved. The swab tip was then retracted back into the sheath prior to extraction to ensure that it had no contact with the external environment or an internal environment other than the distal colon, then the turtles were returned to ventral recumbency and permitted to return to the ocean. The tip of the swab was removed using a sterile wire cutter, placed into a sterile Eppendorf tube and sealed. Total sample collection time was approximately 10 min. Negative control swabs were not collected as we were confident in our col-
lection technique eliminating environmental contamination. The Eppendorf tube containing the swab was immediately placed into a portable cool box filled with ice, and once back at the field station they were frozen at −20°C for approximately 3–4 d. Swabs were transported back to the laboratory using dry ice (−78.5°C), where they were stored at −80°C until extraction could take place approximately 1 wk later.

2.3. DNA extraction

DNA was extracted manually using the phenol-chloroform method (Green et al. 2012). In each Eppendorf tube, 500 µl of extraction buffer consisting of 20 mM ethylenediaminetetraacetic acid (EDTA), 0.1 M Tris, 1% cetrimonium bromide (CTAB), 56 mM NaCl, pH 8 was added so that swabs were completely covered. We then added 20 µl of Proteinase K (Qiagen Proteinase K (10 ml) 10 ml (>600 mAU ml−1 solution)) to each vial, along with 60 µl of 10% sodium dodecyl sulphate (SDS). The mixture was then incubated at 55°C overnight. The next day, 50 µl of 5 M NaCl and 500 µl of phenol was added, and the Eppendorf tubes were shaken until an emulsion was formed. Eppendorf tubes were then incubated at room temperature for 10 min, with intermittent mixing. The Eppendorf tubes were then centrifuged (Microcentrifuge Fresco 21, Thermo Fisher) at 10 000 rpm (9638 × g) for 10 min and the supernatant removed and added to a new Eppendorf tube containing 250 µl phenol and 250 µl chlorophorm: isoamyl-alcohol (24:1). The tubes were again centrifuged (Microcentrifuge Fresco 21, Thermo Fisher) at 10 000 rpm (9638 × g) for 10 min and the resultant supernatant added to a new Eppendorf tube containing 500 µl of chlorophorm: isoamyl-alcohol. Once again, the Eppendorf tubes were centrifuged (Microcentrifuge Fresco 21, Thermo Fisher) at 10 000 rpm (9638 × g) for 10 min. The supernatant was then added to a new Eppendorf tube containing 3 M sodium acetate at a volume equal to 10% of the extraction solution. We then added 1 ml of ice-cold 99% ethanol to each Eppendorf tube, which was then placed into a freezer at −20°C for 1 h. The Eppendorf tubes were then centrifuged (Microcentrifuge Fresco 21, Thermo Fisher) at 4°C at 12 000 rpm (13 878 × g) for 5 min. After centrifugation the alcohol was removed and the lids left off the Eppendorf tubes to allow the DNA pellet to dry. Once dried, 25 µl of 1 × Tris-EDTA (TE) was added to each Eppendorf tube, and the extracted DNA was stored at −20°C until amplicon sequencing could take place.

2.4. 16S rRNA gene amplicon sequencing

The V3-V4 region of 16S rRNA genes were amplified with forward primer 5’ ACT CCT ACG GGA GGC AGC AG 3’ and reverse primer 5’ GGA CTG CHV GGG TWT CTA AT 3’ using Q5 high fidelity polymerase (New England Biolabs) with the barcoding strategy of Fadrosh et al. (2014). Sequencing was performed on an Illumina MiSeq system (2 × 300 bp) on all samples simultaneously, and negative controls were included in the sequence run.

2.5. Data processing

Sequence data was analysed using QIIME version 1.9.1 (Caporaso et al. 2010) using default parameters and a Phred quality threshold of >20. The UCLUST algorithm (Edgar 2010) was used to pick operational taxonomic units (OTUs) at 97% sequence identity, and a Biome table was produced. Potentially chimeric sequences were identified using Pintail (Ashelford et al. 2005). BLAST was used to assign taxonomy against the Greengenes database (DeSantis et al. 2006) and QIIME version 1.9.1 defaults. Additional assignment of taxonomy was performed using a command line version of BLASTN (Altschul et al. 1997) against the NCBI 16S Microbial database.

2.6. Statistics and data analysis

Samples were rarefied to a minimum sequencing depth of 3000 reads. Initial exploration of the Biome table data was performed using the online Calypso software (http://cgenome.net/wiki/index.php/Calypso) (Zakrzewski et al. 2017). Data was further analysed with R, utilising package ‘phyloseq’. Alpha diversity was explored using observed OTUs, Shannon index and Chao1 estimates. Alpha diversity indices were tested for normality using the Shapiro-Fisher) a final time at 4°C at 12 000 rpm (13 878 × g) for 5 min. After centrifugation the alcohol was removed and the lids left off the Eppendorf tubes to allow the DNA pellet to dry. Once dried, 25 µl of 1 × Tris-EDTA (TE) was added to each Eppendorf tube, and the extracted DNA was stored at −20°C until amplicon sequencing could take place.

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ences between groups. Shannon diversity in loggerheads \((W = 0.92, p = 0.05)\) was the only normally distributed metric, and so comparisons between populations were made using ANOVA. All other measures were non-normally distributed: observed OTUs (loggerhead: \(W = 0.85, p < 0.01\); flatback: \(W = 0.88, p < 0.01\)), Shannon diversity (flatbacks: \(W = 0.84, p < 0.01\), and Chao1 (loggerhead: \(W = 0.88, p < 0.01\); flatback: \(W = 0.87, p < 0.01\), and comparisons were made using the Kruskal-Wallis test. Linear discriminant analysis (LDA) effect size (LEfSe) was performed to determine if differential abundances of OTUs were attributable to location of capture with a size effect cut-off of 3.0 on the logarithmic LDA score used. For all statistical analyses significance was accepted if \(p < 0.05\).

3. RESULTS

3.1. Loggerhead turtles

After rarefaction, we were able to include samples from 6 of the loggerheads from Juno Beach and 18 of the loggerhead turtles from Mon Repos (Table S1 in the Supplement; www.int-res.com/articles/suppl/n042p095_supp.xlsx). In Juno Beach loggerheads we identified 201 OTUs, while from Mon Repos loggerheads we identified 272 OTUs. For both populations, the predominant phylum was Proteobacteria, with differential abundance in the 2 populations being observed for other phyla identified (Fig. 1a, Table S1). In animals from Mon Repos, the next most commonly isolated phyla were Spirochaetes, Bacteroides and Actinobacteria, while in Juno Beach animals Actinobacteria, Bacteroides and Firmicutes were the next most populous phyla (Fig. 1a). Alpha diversity differed significantly among loggerheads from the 2 different populations for all metrics: observed OTUs \((\chi^2 = 18.031, df = 1, p < 0.01)\), Shannon \((df = 1, SS_T = 36.91, MS_T = 36.91, F = 54.98, p < 0.01)\), and Chao1 \((\chi^2 = 14.86, df = 1, p < 0.01)\) (Fig. 2a). Beta diversity, as assessed by PCoA, showed that the 2 populations clustered separately and distinctly from each other (Fig. 3). These observations were significantly different \((df = 1, SS_T = 0.82, MS_T = 0.82, F = 2.6, R^2 = 0.1, p = 0.03)\). Overall, 37 (29.6%) of the OTUs were shared between the 2 populations (Table S2). Sixty-one (64.9%) OTUs were unique to animals from Mon Repos, in comparison to 27 (21.6%) unique OTUs from animals from Juno Beach (Table S2). Linear discriminant analysis revealed 27 OTUs to be responsible for explaining the variance seen between the different populations, with 20 of these being found in Mon Repos loggerheads and 7 being found in Juno Beach loggerheads (Fig. 4a, Table S3). Some of the most significant OTUs explaining variation for Mon Repos animals included Helicobacter helicobacteriustelae_327563 and Vibrio sp._452638 (Fig. 4a), while for Juno Beach turtles the major OTUs responsible for explaining variation included Stenoxysbacter sp._751919 and Pasteurellaceae_142268 (Fig. 4a).

3.2. Flatback turtles

After rarefaction, we were able to include samples from 19 of the flatbacks from Crab Island and from 10 of the flatbacks from Port Hedland (Table S4). In Crab Island flatbacks we identified 160 OTUs while in Port Hedland flatbacks we identified 241 OTUs. Firmicutes was the predominant phyla in animals from Crab Island, while Proteobacteria was the most commonly isolated phyla in Port Hedland turtles, followed by Actinobacteria, Bacteroidetes and Firmicutes (Fig. 1b, Table S4). Alpha diversity differed markedly among flatback turtle groups from the 2 different populations with significance noted for all metrics: observed OTUs \((\chi^2 = 18.031, df = 1, p < 0.01)\), Shannon \((df = 1, SS_T = 17.053, df = 1, p < 0.01)\), and Chao1 \((\chi^2 = 14.85, df = 1, p < 0.01)\) (Fig. 2b). Principal coordinate analysis of microbiota composition indicated that animals from Port Hedland and Crab Island had distinct and significant differences \((df = 1, SS_T = 2.84, MS_T = 2.84, F = 10.175, R^2 = 0.27, p = 0.001)\) in their clustering patterns (Fig. 3b). Overall, 28 (29.8%) OTUs were shared between the 2 populations (Table S5). Sixty-one (64.9%), OTUs were unique to animals from Port Hedland, in comparison to 5 (5.3%) unique OTUs from animals from Crab Island (Table S5). Linear discriminant analysis revealed 76 OTUs to be responsible for explaining the variance seen between the different populations, with 69 of these being found in Port Hedland turtles, and 7 being found in Crab Island turtles (Fig. 4b, Table S6). Some of the most significant OTUs explaining variation for Crab Island animals included Bacillus sp._161559, Bacillus sp._535651, Bacillus sp._161559, Bacillus sp._535651, and Bacillus endophyticus sp._410037 (Fig. 4b). For Port Hedland turtles the major OTUs responsible for explaining variation included Actinobacteria_108564, SUP05_285720, and Campylobacter campylobactercurvus_607889 (Fig. 4b).
4. DISCUSSION

Sea turtles are cosmopolitan species with a worldwide distribution, are unusual among vertebrates in that they occupy a wide variety of ecological niches throughout their lives, and as adults they typically undergo long and physiologically arduous journeys for reproductive purposes. The phylogenetic history of sea turtles is well described on both a species level and between populations of the same species (Pritchard 1997, Duchene et al. 2012). Understanding the role that the microbiota has played in driving sea turtle phylogeny and physiology is the next important step in unravelling the complexities of their evolutionary history. Although a number of investigations exist which explore the microbiota of sea turtles (Abdelrhman et al. 2016, Ahasan et al. 2017, 2018, Price et al. 2017, Campos et al. 2018, Kittle et al. 2018, Arizza et al. 2019, Biagi et al. 2019), making meaningful comparisons between these studies and ours is challenging. Microbiome studies are complicated by differences in methods, differences in species sampled, and the nutritional status of sampled animals, which differed between these studies and
the data we report here. Further, we sampled nesting turtles, which are physiologically distinct from sick/injured or foraging turtles. Thus, because our goal was to interpret comparisons between microbiota composition subpopulations, we must turn to similar research on other taxa.

Our analysis of the bacterial microbiota of wild nesting sea turtles indicates that microbial commu-
ties differ significantly among individuals of the same species but which originate from different populations. These differences may be apparent in both composition and diversity. This investigation highlights the importance of interpreting microbiota data in concert with environmental factors and potential genetic differences among host populations. Elucidating the intrinsic and extrinsic causes driving these differences is difficult due to the lack of basic biological data for both individual animals and the different populations of turtles in this investigation. While, broadly speaking, dietary, and migratory information exists for some species and populations of sea turtles, these generalisations should be applied with caution to individuals captured from a nesting beach. For example, nesting loggerheads show a high degree of fidelity to nesting beaches and foraging grounds, but females nesting at the one rookery will have migrated from numerous foraging areas (Limpus 2008b). Female loggerheads tagged at Queensland rookeries have been recorded from feeding areas as far away as Indonesia, Papua New Guinea, Solomon Islands, and New Caledonia, but also from local areas including the Northern Territory, Queensland and New South Wales (Limpus et al. 1992). We were unable to identify the origin of each turtle that was sampled, and therefore it is difficult to determine exactly what factors may be influencing bacterial microbiota community structure in our samples.

In addition to observed differences between sea turtle populations, we also noticed differences between individuals within populations. Such differences have been observed in separate studies looking at loggerhead turtles from the Mediterranean (Abdelrhman et al. 2016, Arizza et al. 2019, Biagi et al. 2019), but no explanations have been given to explain these differences, likely because these investigations were conducted independently. In humans, the establishment and maintenance of the bacterial microbiota are heavily influenced by a range of factors including diet, host genotype, and microbial interactions (Dethlefsen et al. 2006). These factors are further influenced by other random, unpredictable events such as colonisation history. Within an individual, microbes are unevenly distributed, and they have differences in transmissibility (Dethlefsen et al. 2006). This variation is greatly amplified under the paradigm of stochastic niche occupancy, in which microbes must be able to persist and multiply using available resources in the face of fierce competition from existing colonists (Dethlefsen et al. 2006). Such unpredictabili-
Fig. 4. Linear discriminant analysis (LDA) effect size (LEfSe) algorithm was used on operational taxonomic units (OTU) tables to determine taxa that best characterized differences between each location of capture. (a) Comparison of loggerheads from Mon Repos, Australia, with loggerheads from Juno Beach, USA. (b) Comparison of flatbacks from Crab Island, Queensland, with flatbacks from Port Hedland, Western Australia.
ity accounts for the differences observed in microbiotas between closely related individuals such as inbred mice (Hoy et al. 2015) and human identical twins (Turnbaugh et al. 2010). Therefore, individual variation within a population of animals, such as the turtles in our investigation, is an expected finding.

The role that diet may be playing in determining bacterial microbiota composition in our samples is unknown. Loggerhead turtles are primarily carnivorous (Jones & Seminoff 2013), feeding on a wide range of prey items with considerable dietary plasticity (Park et al. 2005, Seney & Musick 2007, Limpus 2008b, Frick et al. 2009). Although there are differences in the species of prey items consumed among loggerhead populations, the taxonomic classes of ingested prey are relatively similar over phylogeographic separations (Jones & Seminoff 2013). The major differences in dietary preferences come from the proportion of benthic or pelagic fauna in the diet, which is predominately affected by the life stage of the turtle (Jones & Seminoff 2013). All of the loggerheads in this investigation were adult, and therefore, broad dietary preference is likely to be similar for the 2 populations. In humans, diet markedly affects the microbiota and is principally driven by the volume of fibre consumed (Albenberg & Wu 2014). It remains to be seen whether or not the observed species-level prey differences across loggerhead populations, rather than dietary variations at genus, family or higher taxonomic levels, selects for specific microbial OTUs in sea turtles.

In addition to food items consumed, food acquisition and digestive strategy can play a role in altering microbial communities in animals. For example, the Burmese python *Python molurus* is a sit-and-wait predator that consumes enormous, infrequent meals, often punctuated by prolonged periods of starvation (Costello et al. 2010). This feast or famine tactic means that the gastrointestinal tract, including the microbiota, undergoes significant change depending on the presence or absence of food in the gut (Costello et al. 2010). These changes start to become evident between 12 h to 3 d after prey ingestion (Costello et al. 2010). Similar patterns of fluctuations in microbial community composition are seen in other vertebrates in response to caloric restriction including humans (Remely et al. 2015), fish (Xia et al. 2014), bears (Sommer et al. 2016), alligators (Keenan et al. 2013), mice (Beli et al. 2018), and penguins (Dewar et al. 2014). Like Burmese pythons, sea turtles may undergo long periods of fasting, particularly during breeding and migration (Hays et al. 2002, Jessop et al. 2004), but how this affects gastrointestinal bacterial community composition is unknown. All animals sampled during this investigation were nesting females, and it is likely that they had been inappetant for an extended period, which may be influencing the bacterial microbiota composition of our populations. It is of particular relevance to study microbiotas of female sea turtles during their egg-laying reproductive period because it is the microbiota during this time that is likely to be transferred to their offspring rather than the microbiota during foraging, non-reproductive periods (Scheelings 2019). Given the rapidity with which the bacterial microbiota fluctuates in other vertebrates, it is reasonable to assume that the bacterial microbiota of nesting females is not the same as the bacterial microbiota of foraging animals. Although we were not able to determine how fasting in nesting turtles affects microbial diversity, some authors propose that fasting samples represent the core microbial OTUs, with other OTUs fluctuating in number in response to post-prandial physiological changes (Costello et al. 2010, Keenan et al. 2013). If, as hypothesised, diet has little effect on this core bacterial microbiota, then a possible explanation for differences observed in our loggerhead samples is differences in host genetics. In humans, host genetic variation has been shown to account for microbiota composition from a number of different body sites (Goodrich et al. 2014, Blekhman et al. 2015), and this phenomenon may be a significant contributing factor in the differences observed in our loggerhead samples, given their genetic separation (Bowen & Karl 2007). In contrast it seems reasonable that genetics is less likely to be contributing to observed differences among flatback populations, given the limited genetic variability in this species compared with loggerheads (Dutton et al. 2002), and so other factors are more important in shaping bacterial microbial communities in flatback populations.

We identified that flatback turtles from Crab Island had much lower microbial diversity than their counterparts from Port Hedland. Such observations in other species have been attributed to habitat degradation and may result in decreased ability of animals to utilise environmental resources (Amato et al. 2013, Barelli et al. 2015, Rosenfeld 2017). We did not attempt to assess habitat quality in this investigation, but Crab Island is a remote, uninhabited island in the Endeavour Strait in the Gulf of Carpentaria with little signs of human impact (authors’ pers. obs.). Importantly, it is believed that flatback turtles nesting on Crab Island do not feed in the Gulf of Carpentaria, and so migrate considerable distances from foraging...
grounds for reproduction (Limpus et al. 1993). Thus, in the context of their lives, flatback turtles only spend a relatively short period of time in this area, and therefore any environmental influence is most likely coming from elsewhere. Further work is required to identify the effects that habitat disturbances have on sea turtle bacterial microbials, and whether or not they can act as sentinels for oceanic health.

The observed increase in relative abundance of Firmicutes in flatback turtles from Crab Island may possibly be associated with increased adiposity (Bäckhed et al. 2004, Cui et al. 2013). Bacteria from this phylum are able to increase fat metabolism through modulation of the host genes that regulate fat storage (Turnbaugh et al. 2006). Flatback turtles from Western Australia undergo a shorter migration to their natal beaches than Crab Island turtles, as they forage relatively close to their nesting rookeries (Pendoley et al. 2014). This might mean that turtles from Crab Island are inappetant for a longer period of time, and therefore will have a greater reliance on stored body fat to fuel their migration. This physiological phenomenon may preferentially select for Firmicutes as an adaptation to their prolonged fast.

We showed that within sea turtle species, geographic location has a major effect on bacterial microbiota composition. Factors that may explain these observed differences include host genetics, differences in foraging habitat quality and differences in migratory distance (and thus durations of inappetence) between foraging and breeding grounds. The mechanisms by which these factors may influence microbial composition of sea turtle gastrointestinal tracts warrants further investigation. The results of this study highlight the importance of interpreting bacterial microbiota data of wild animals in the context of geography. This study represents the first investigation to explore the effects of population on sea turtle bacterial microbials and fills an important knowledge gap in the vertebrate microbiota literature.

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