**Helicobacter spp. in Necropsied Southern Sea Otters (Enhydra lutris nereis) Is Associated With Gastric Ulcers and Sensitive to Multiple Antibiotics**

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Southern sea otters (Enhydra lutris nereis) are threatened marine mustelids that commonly have gastric ulcers with secondary hemorrhage (melena) as a contributing cause of death. Although Helicobacter spp. infections are known to cause gastric ulcers and gastritis in humans and ferrets, it is unknown if the sea otter bacterium, *H. enhydrae* sp. nov., causes similar gastric pathology. Determining whether *Helicobacter* detection is associated with sea otter gastric pathology is the first step toward using this information to expedite diagnosis and treatment. We investigated the proportion of *Helicobacter* infections in 46 necropsied southern sea otters via quantitative real-time polymerase chain reaction (qPCR) of the 16S rRNA gene. *Helicobacter* DNA was detected in fresh-frozen and formalin-fixed gastric body and pyloric tissues using *Helicobacter* genus-specific 16S rRNA primers. Data from gross necropsy and histopathology were analyzed for associations between *Helicobacter* detection via qPCR and presence/absence of gastric pathology. ETEST® gradient strips were utilized to investigate antimicrobial minimum inhibitory concentrations for *H. enhydrae* isolates. *Helicobacter* spp. were detected in the gastric tissue of 85% of sea otters in this study. Fresh-frozen samples were more commonly *Helicobacter* qPCR-positive than formalin-fixed tissue, indicating variable sensitivity of detection in relation to post-necropsy tissue processing methods. Diagnosis of gastric ulcers at necropsy was significantly associated with *Helicobacter* qPCR detection in gastric mucosa (*P* = 0.005), while age, sex, presence of melena, shark trauma, and protozoal infection were not associated (*P* > 0.1). *Helicobacter enhydrae* isolates were sensitive to clarithromycin and tetracycline *in vitro* at physiologically relevant concentrations. Overall, this work suggests that *Helicobacter* spp. might be commonly found in southern sea otters, particularly those with ulcers, and that *H. enhydrae* can be treated with several commonly used anti-*Helicobacter* therapies.

**Keywords:** antimicrobial sensitivity test, gastric ulcer, *Helicobacter enhydrae*, *Helicobacter* detection, quantitative PCR, sea otter
INTRODUCTION

Sea otters (*Enhydra lutris*) are one of the smallest marine mammals, but are the heaviest member of the family Mustelidae, with a unique metabolism. These animals have twice the metabolic rate of other marine mammals and 2–3 times the rate of a comparably sized terrestrial animal; as a result, they require food consumption equivalent to 25% of their body weight each day (Iversen, 1972; Morrison et al., 1974; Yeates et al., 2007). With such high caloric demands, gastrointestinal health is critical for the well-being of sea otters. Bacteria in the *Helicobacter* genus are a highly influential component of overall gastric health. In humans, gastric infection by the bacterium *Helicobacter pylori* is found in up to 50% of people worldwide and is associated with gastritis, gastric ulcers, and gastric cancers (Marshall and Warren, 1984; Cover and Blaser, 1996; Zamani et al., 2018). Mammalian infections with non-*pylori* *Helicobacter* species have been shown to co-occur with gastritis, gastric ulcer, and/or gastric adenocarcinoma development, specifically, *H. cetorum* in marine mammals and *H. mustelae* in ferrets (Fox et al., 1990, 1997; Harper et al., 2002, 2003).

A novel *Helicobacter* species, *H. enhydrae* sp. nov. (strain MIT 01-6242, GenBank No. AY203901), was isolated and characterized from the gastric mucosa of southern sea otters (*Enhydra lutris nereis*) (Shen et al., 2017). Electron microscopy and silver stains showed a slightly curved bacterial rod with lateral flagella (Shen et al., 2017). This novel bacterium was catalase- and oxidase-positive and urease-negative with 65 genes homologous to virulence factors in related genera (Shen et al., 2017). Based on targeted *Helicobacter* spp.-specific 16S rRNA polymerase chain reaction (PCR), 58% of 31 tested sea otters were positive for *Helicobacter* DNA in gastric mucosa collected at necropsy (Fox et al., 1988; Shen et al., 2017). The *H. enhydrae* sp. nov. strain demonstrated close phylogeny to *H. mustelae* from domestic ferrets (*Mustela putorius furo*), based on *Helicobacter* 16S and 23S rRNA gene sequences (Shen et al., 2017). A prior study found, but did not characterize, two southern sea otter *Helicobacter* isolates (MIT 01-5923 and MIT 01-5924) that shared phylogeny with *Helicobacter* spp. from a California sea lion (*Zalophus californianus*) (MIT 02-5519-C) and a harp seal (*Phoca groenlandica*) (MIT 01-5529-A) (Harper et al., 2003). Sea otter microbiome analyses had also identified 11 *Helicobacter* spp. amplicon sequence variants from gingival and rectal swabs (Dudek, 2018).

As part of a large retrospective study of southern sea otter mortality patterns from 1998 to 2012, gastric ulcers and melena (i.e., upper gastrointestinal bleeding) were identified as a contributing cause of death (COD) for 42.3% (237/560) of sea otters examined via necropsy and histopathology (Miller et al., 2017). While gastric ulcers have been previously reported as a cause of sea otter mortality (Kreuder et al., 2003; Williams et al., 2018; Mansour-Ghanaei et al., 2019), the high prevalence of this condition was not appreciated until this comprehensive study. In prior reports, gastric ulcers and melena in sea otters have often been attributed to captivity-associated stress or were considered a sequela to petroleum exposure (Lipscomb et al., 1993; Loughlin, 1994; Williams and Davis, 1995; Reimer and Lipscomb, 1998).

Other potential causes of gastric ulcers and melena in sea otters have not been systematically assessed, such as *Helicobacter* spp.-associated gastric infections. In a closely related mustelid, the ferret, *H. mustelae*, is strongly linked to gastric ulcers (Fox et al., 1988, 1990, 1991a; Batchelder et al., 1996).

As a federally listed threatened species with population estimates of just over 2,962 animals (Hatfield et al., 2019), it is important to determine whether gastric pathology in southern sea otters may be associated with *Helicobacter* infections. As a first step, we screened southern sea otter fresh-frozen and formalin-fixed gastric tissues for the presence of *Helicobacter* spp. DNA via quantitative real-time PCR (qPCR) using *Helicobacter* 16S rRNA primers. Demographic, gross postmortem, and histology data were evaluated to measure associations with *Helicobacter* detection via qPCR. As a secondary objective, we assessed the antibiotic efficacy of *H. enhydrae in vitro* to antibiotics tested against *H. pylori*.

MATERIALS AND METHODS

Tissue Collection and Processing Procedures

Gastric samples were from fresh, necropsied sea otters collected by staff of the California Department of Fish and Wildlife (CDFW) in the course of their duties as an official or state employee. All work was performed in accordance with Section 109(h) of U.S. Marine Mammal Protection Act (MMPA), U.S. Fish and Wildlife Service (Service) regulations implementing the MMPA at 50 CFR 18.22(a), and in accordance with Service regulations implementing the US Endangered Species Act at 50 CFR 17.21(c)(3).

Necropsy and histology sampling protocols for the 46 enrolled sea otter carcasses were performed as previously described (Kreuder et al., 2003). In brief, opportunistic sampling of minimally decomposed sea otters of various age classes and sexes was performed; the sea otters were refrigerated at 7–10°C and examined at CDFW—Marine Wildlife Veterinary Care and Research Center (CDFW-MWVCR) (Santa Cruz, CA, United States) between 2015 and 2017. During necropsy, the stomach and proximal duodenum were opened lengthwise; and the mucosa was visually inspected for erosions, ulcers, and melena. Gastric lesion locations and relative severities were documented via summary notes and, in some cases, photographs. Two sets of adjacent gastric body and pylorus samples were collected using sterile scalpel blades. One set was placed in cryovials and then frozen in a −80°C freezer until processed for DNA purification. The other set was placed in 10% buffered formalin for fixation, tissue processing, and microscopic examination. The formalin-fixed gastric mucosa was trimmed and submitted to the UC Davis Veterinary Medical Teaching Hospital for paraffin embedding, cutting of 5 μm thick sections, and hematoxylin and eosin (H&E) staining for examination by light microscopy.

Gastric erosions were defined as foci of denuded surface epithelium and underlying lamina propria where tissue loss did not extend into the muscularis mucosa; ulcers penetrated more

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deeply and extended into or through the muscularis mucosa (Tarnawski et al., 1995). Both forms of gastric lesions were combined, and the pooled dataset was referred to as "ulcers" to facilitate analyses.

**DNA Purification From Gastric Tissues and qPCR Assay**

DNeasy Blood and Tissue Kit (Qiagen) and QIAamp DNA FFPE Tissue Kit (Qiagen) were used for DNA extraction of the fresh-frozen ("frozen") gastric body and pylorus samples (25 mg/sample) and the formalin-fixed, paraffin-embedded ("fixed") samples (60–80 µm/sample), respectively, per Qiagen's extraction protocols. Primer sequences (forward primer 116F 5'-AGT AAT GCA TAG GTT ATG TGC CCT-3' and reverse primer 237R 5'-CAA GCT GAT AGG ACA TAG GCT GAT-3') were constructed by Geneious R10.1.3 (Biomatters Ltd.) and amplified a 117–122 bp product of the 16s rRNA gene from the *Helicobacter* genus. qPCR was performed on the CFX Connect™ Thermal Cycler (Bio-Rad Laboratories, Inc., United States) using a 25 µL reaction mix final volume.

The following qPCR conditions were specific for SensiMix™ SYBR® No-ROX (Bioline) and the primer sequences: 1 cycle of polymerase activation at 95°C for 10 min, then 40 cycles of denaturing at 95°C for 15 s, annealing at 57°C for 15 s, and elongation at 72°C for 15 s. The following were used as qPCR controls: no-template control (NTC) with nuclease-free water, negative control with *Escherichia coli* DH10B genomic DNA, positive control with *H. enhydrae* genomic DNA, isolated as above. All frozen and fixed gastric samples had 2–4 technical replicates (Supplementary Tables S1, S2). For the frozen samples (*n* = 92), we were able to obtain separate gastric body and pylorus for all 46 sea otters (Table 1). For the fixed samples (*n* = 59), gastric body and pylorus samples were either on separate paraffin blocks (30/59) or on a single block (29/59) (Table 1).

The quantification cycle (Cq) is the cycle (1–40, with 40 being the lowest value) in which the intensity of the fluorogenic-marked amplicons is higher than background noise. Serial dilutions of purified *H. enhydrae* genomic DNA, including 89, 8.9, 8.9 x 10^-4, 8.9 x 10^-4, and 8.9 x 10^-6 ng/µL via NanoDrop 2000 Spectrophotometer (Thermo Scientific), were used to run an amplification plot and standard curve to determine the sensitivity and specificity of the qPCR assay (Figure 1). The *H. enhydrae* genome size of 1.6 Mb was used to calculate the copy number (URI Genomics Sequencing Center, 2004).

**Statistical Analyses**

Associations between sea otter gastric *Helicobacter* detection and demographic, gross postmortem, and histologic variables were assessed using statistical software JMP Pro 15.0 (SAS Institute Inc., United States). Similar relationships were evaluated for gastric pathology. The Fisher's exact test evaluated possible associations between *Helicobacter* PCR detection and gastric ulcers, as well as demographic, gross postmortem, and histology data. The significance level was set at α = 0.05, such that *P* < 0.05 indicated a statistically significant difference.

**Histology**

Histological evaluations performed by lead author (F. Batac, MS) were reviewed by a veterinary pathologist (M. Miller, DVM, Ph.D., MS). Current human and sea otter gastric histology grading systems were reviewed and referenced in conducting sea otter histology evaluations (Supplementary Table S3). H&E stained slides were used to evaluate the fixed gastric body and pylorus for features including erosions, ulcers, melena, gastritis, inflammatory cell type (neutrophilic or lymphoplasmacytic), and detection of suspect *Helicobacter*-like organisms (Supplementary Table S4).

**H. enhydrae Culture and ETEST® Antimicrobial Sensitivity Testing**

*Helicobacter enhydrae* MIT 01-6242 from Shen et al. (2017) was used for the antimicrobial sensitivity testing. Southern sea otter *Helicobacter* isolates MIT 01-5923 and MIT 01-5924 from Harper et al. (2003) were unavailable. *Helicobacter enhydrae* MIT 01-6242 was grown on solid media consisting of Columbia agar with 5% defibrinated horse blood (CHBA; Hemostat Laboratories) containing 10 µg/mL vancomycin, 50 µg/mL cycloheximide, 5 µg/mL cefsulodin, and 2.5 units/mL polymyxin B, or liquid media containing *Brucella* broth with 10% heat-inactivated fetal bovine serum (BB10). For either growth mode, *H. enhydrae* was cultured at 37°C under microaerobic conditions (10% CO2 and

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**TABLE 1** | Number of enrolled sea otters and total samples based on gastric sampling type.

| Gastric sampling type       | Enrolled sea otters | Total gastric samples |
|-----------------------------|--------------------|-----------------------|
| Collected fresh at necropsy | 46                 | 92                    |
| frozen immediately          |                    |                       |
| Formalin-fixed paraffin     | 45                 | 59                    |
| embedded blocks             |                    |                       |
| Gross necropsy data         | 41                 | 82                    |
| Histology analyses          | 37                 | 57                    |

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**FIGURE 1** | Quantitative PCR results for serial dilutions of *Helicobacter enhydrae* genomic DNA. Amplification plots from left to right were generated from 89, 8.9, 8.9 x 10^-4, 8.9 x 10^-4, and 8.9 x 10^-6 ng/µL.
5% O₂, balance N₂). For the antimicrobial sensitivity testing, *H. enydrae* was cultured on solid CHBA media, with subculture and passage to fresh plates every 3 days for 6 days. For the BB10 overnight incubation, a small amount of the plate-grown sample was transferred to the liquid. After overnight growth with shaking at 220 rpm, the absorbance (OD₆₀₀) was determined and cultures were used when the absorbance was between 0.199 and 0.785. Then, 100 µL of this broth was spread evenly onto separate CHBA plates. After spreading, the ETEST® strips (BioMérieux, each impregnated with a separate antibiotic), were placed on the surface of individual plates, and the plates were incubated as above. The zones of bacterial growth inhibition around the surface of individual plates, and the plates were incubated for 2 days of incubation. The antibiotics tested in the study were amoxicillin, clarithromycin, chloramphenicol, kanamycin, nalidixic acid, and tetracycline.

**RESULTS**

**Helicobacter** spp. Is Frequently Detected in Frozen Sea Otter Gastric Tissue

We first examined our set of 92 frozen gastric tissue samples (body and pylorus) from 46 southern sea otters using qPCR to quantify the amount of *Helicobacter* DNA. We first ran an amplification plot and standard curve with purified *H. enydrae* DNA (Figure 1). With this approach, we could detect *H. enydrae* DNA concentrations as low as 8 × 10⁻⁶ ng/µL and a qPCR assay sensitivity of 5 copies of *Helicobacter* DNA. These samples were considered *Helicobacter* qPCR-positive, while samples below this detection were marked as *Helicobacter* qPCR-negative. Of the 46 sea otters, 39 (85%) had at least one positive tissue. By tissue, 83% of gastric pylorus (38/46) and 65% of gastric body (30/46) samples were *Helicobacter* qPCR-positive (Figure 2). Among sea otters that were qPCR-positive, nearly three-quarters had *Helicobacter* DNA detected in both the gastric body and pylorus (29/39), while a quarter had *Helicobacter* DNA detected in only one of the two anatomical locations (10/39). Overall, these data show that majority of sea otters in this study had *Helicobacter* DNA in their stomachs, with more sea otters showing positive gastric pyloric samples than gastric body ones.

**Formalin-Fixed Tissues Underrepresent Helicobacter Status**

We next asked how formalin-fixed paraffin-embedded gastric tissue samples, from the same otters as those analyzed above, would compare for *Helicobacter* spp. detection. We were curious about this question because laboratories often have preserved tissue that could be used for similar analyses. Our tissues had been preserved in 10% buffered formalin for varying lengths of time (2 days to 2–3 weeks) prior to paraffin-embedding, and the paraffin-embedded samples were used for our analysis. qPCR of DNA extracted from these fixed tissues found that only 40% (18/45, fixed samples unavailable for one otter) scored positive for *Helicobacter* DNA. The fixed samples produced a mean Cq that was 4.2–4.7 higher than frozen samples, resulting in many of them being called as *Helicobacter* qPCR-negative (Figure 3). Both the gastric body and pylorus fixed samples yielded much lower *Helicobacter* detection than did frozen tissue. These results suggested that sea otter *Helicobacter* spp. qPCR testing of frozen gastric tissue was a more sensitive method than testing fixed gastric tissues.

**Positive Association Between Helicobacter spp. Detection and Gastric Ulcers in Sea Otters**

To investigate possible associations between *Helicobacter* spp. detection and gastric pathology, we evaluated a subset of 82 frozen gastric body and pylorus samples from 41 sea otters for which gross gastric evaluations were included in the postmortem examination (Table 1). Fifty-one percent (21/41) of necropsied sea otters had gastric ulcers observed in the

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**FIGURE 2** | Helicobacter quantitative PCR results for the fresh-frozen and formalin-fixed southern sea otter gastric tissue samples. (A) Results for the fresh-frozen gastric samples (n = 92). (B) Results for the formalin-fixed paraffin embedded gastric samples (n = 59). (C) Results for the fresh-frozen gastric body samples (n = 46). (D) Results for the fresh-frozen gastric pylorus samples (n = 46).

**FIGURE 3** | Comparison of the average quantification cycle for sea otter gastric tissues as fresh-frozen (n = 92) and formalin-fixed, paraffin-embedded (n = 59) samples. Fixed both = gastric body and pylorus in same paraffin-embedded sample. Error bars = standard deviation.
gastric body and/or pylorus, with 45% (37/82) of frozen gastric samples displaying gastric ulcers. Among frozen gastric samples that had ulcers, nearly 90% (33/37) were also Helicobacter qPCR-positive (Figure 4). Helicobacter qPCR-positive gastric samples were significantly associated with gastric ulceration (2-sided Fisher’s exact test, \( P = 0.005 \)). Among the 41 sea otters with gross gastric evaluations at necropsy, the proportion of otters that were Helicobacter qPCR-positive was higher among those with gastric ulcerations (2-sided Fisher’s exact test, \( P = 0.04 \)), and in fact, 6 of the 7 Helicobacter qPCR-negative individuals had no visible gastric ulcers. Collectively, these findings suggest that Helicobacter qPCR-positive sea otters were more likely to have gastric ulcers than Helicobacter qPCR-negative animals (Figure 4).

No associations were found between Helicobacter qPCR detection and age, sex, melena, shark trauma COD, or parasitic CODs (e.g., acanthocephalan or protozoal infections) (Fisher’s exact test, \( P > 0.1 \)), although small sample sizes limited our ability to detect differences (Supplementary Table S1).

Histologic Assessments of Ulcers Did Not Reflect Gross Postmortem Assessments

We determined whether histological samples could accurately reflect grossly apparent gastric ulcer status in sea otters (Supplementary Table S3). H&E-stained histology slides determined 25% (14/57) gastric ulcer presence as compared to gross postmortem examination which reported 45% (37/82) ulcer presence (Supplementary Tables S1, S4). A histological determination of gastritis or melena were not statistically associated with qPCR Helicobacter detection (Fisher’s exact test, \( P = 0.3 \)), suggesting that these may not be useful indicators of possible Helicobacter infection. This result may be partially explained by our small sample size for histology assessment (Table 1), due in part to autolysis and varying gastric tissue sizes on the microscopic slides. Additionally, gastric histology is performed on a small sample of tissue, which may not accurately represent all pathology present in the organ itself (i.e., focal gastric ulcers may be missed).

Select Histologic Findings for Individual Helicobacter qPCR-Positive Sea Otters

While different histologic attributes were graded, the main characteristics assessed were gastric erosions/ulcers, gastritis, and melena (Supplementary Table S4). Figure 5 shows severe pyloric gastric ulcers from a Helicobacter qPCR-positive subadult female. Severe pyloric gastric mucosal ulcers with perilesional mucosal hemorrhage were confirmed at gross necropsy, based on a dark brown to black appearance due to the action of gastric acid pH on blood (Figure 5A). Histopathology of these lesions revealed mucosal coagulation necrosis associated with thrombosis of underlying blood vessels and perilesional hemorrhage (Figure 5B). This pattern of deep vascular thrombosis leading to an acute wedge-shaped mucosal infarct is more typical of stress, hypovolemia, shock, etc., and not usually Helicobacter-associated.

Chronic inflammation would be more characteristic for Helicobacter-associated lesions. Figure 6 shows pyloric gastric mucosal lesions from a Helicobacter qPCR-positive adult male. Grossly, this otter had severe pyloric gastric mucosal ulcers, severe perilesional mucosal hemorrhage, and melena (Figure 6A). Histopathology revealed focal expansion of pyloric lamina propria by nonsuppurative inflammation and mild stromal collapse (Figure 6B).

**Antimicrobial Sensitivity Tests on H. enhydrae Reveal Sensitivity to Clarithromycin and Tetracycline in vitro**

We investigated the antimicrobial sensitivity of southern sea otter H. enhydrae strain MIT 01-6242 (Shen et al., 2017). We chose ETEST® gradient strips with antibiotics commonly used to treat H. pylori or were previously tested against this H. enhydrae strain to assess minimum inhibitory concentration.
**Table 2** | The minimum inhibitory concentrations (MICs) of Helicobacter enhydrae sp. nov (MT 01-6242) against antimicrobial agents targeted toward bacteria like *H. pylori*.

| Antimicrobial Agent | MICs (µg/mL) |
|---------------------|--------------|
| Amoxicillin (AC)    | 0.5 – 2.0    |
| Clarithromycin (CH) | ≤0.016       |
| Chloramphenicol (CL)| 1.5 – 2.0    |
| Kanamycin (KM)      | 3.0 – 4.0    |
| Nalidixic acid (NA) | 16 – 24      |
| Tetracycline (TC)   | 0.047 – 0.094|

**DISCUSSION**

We found that 85% (39/46) of necropsied sea otters in our study between 2015 and 2017 were qPCR positive for *Helicobacter* DNA in fresh-frozen gastric tissues. Our reported *Helicobacter* prevalence in our sample population is higher than the 58% reported in a prior study (Shen et al., 2017) that used conventional PCR. This may be because qPCR has higher sensitivity than conventional PCR. Our study and Shen et al. (2017) described higher *Helicobacter* detection in the gastric pylorus (83 and 45%, respectively) than the gastric body (65 and 10%, respectively). The gastric body glands all contain acid-secreting parietal cells, while in the pylorus/antrum, parietal cells are either present in only half of the glands in humans or are absent in rodents (Willet and Mills, 2016). The antrum/pylorus is also the preferred *Helicobacter* colonization site for ferrets, with higher bacterial abundance and more significant gastritis in this region (Vargas et al., 1991; Yu et al., 1995; Fox et al., 1997). We tested matching formalin-fixed tissues to investigate whether we could use archived formalin-fixed gastric tissues to determine *Helicobacter* DNA presence. Fixed tissue analysis underrepresented *Helicobacter* status and produced an average overall Cq of 4.2–4.7 higher than their frozen counterparts, and thus, more fixed tissues were falsely *Helicobacter* qPCR negative. The differing Cq values may be due to the creation of abasic sites and DNA fragmentation by formalin fixatives, which can lower DNA integrity and quantity, negatively affecting its ability to be amplified by PCR as shown in other tissues (Do and Dobrovic, 2015). Our study confirms that this effect is true in sea otter tissue samples as well. Due to the differing qPCR Cq values of fixed tissues, the study did not utilize archival fixed southern sea otter gastric tissues dating back to the mid-1990s, which would have increased the sample size and provided a retrospective look into *Helicobacter* prevalence.
Southern sea otters were previously known to be infected with *H. enhydrae* sp. nov., which is related to the pathogenic species *H. mustelae* from ferrets (Shen et al., 2017). The novel species contained virulence factors, such as two copies of flagellin (flagA) and chaperone protein DnaJ (dnaj), that have aided pathogenicity in the related *H. mustelae* (Shen et al., 2017). In addition, gastric ulcers were recognized as a common pathologic lesion in sea otters, often contributing to morbidity and mortality (Miller et al., 2017). However, associations between *H. enhydrae* presence and sea otter gastric ulcers were previously undetermined. In this study, we found that *Helicobacter* qPCR detection was common among the 46 sea otters in our study and had a statistically significant association with gastric ulcers (Figure 4).

An extensively studied mustelid with *Helicobacter* are captive domestic ferrets, which are highly colonized by *H. mustelae* (Fox et al., 1990; Forester et al., 2000). Previous analysis found that 100% of 67 sampled ferrets from United States research facilities and veterinary practices in a 2-year span were *H. mustelae* positive via culture from gastric biopsies (Fox et al., 1991a). This prevalence is higher than we reported for *H. enhydrae*. *Helicobacter mustelae* is known to inhibit acid secretion by parietal cells, increasing gastric pH and creating a favorable environment for bacterial proliferation (Fox et al., 1991b; Vargas et al., 1991), possibly explaining the high levels of ferret infection. *Helicobacter mustelae* is the only non-*pylori* *Helicobacter* reported to cause gastric ulcers and cancer in its native host (O’Toole et al., 2010), and lesion development was confirmed through fulfllment of Koch’s postulates (Fox et al., 1991b).

Gastric ulcer etiology is likely multifactorial in wild and captive sea otters, including a multitude of stressors such as oil spill contamination, captivity, trauma, hypovolemia, shock, and concurrent disease (Lipscomb et al., 1993; Williams and Davis, 1995; Reimer and Lipscomb, 1998). Stress ulcers are defined as multiple superficial erosions of the gastric mucosa. Stress ulcers are considered independent of *Helicobacter* spp. infection and arise in humans from damage to the mucosal barrier that is secondary to some other systemic stress, refluxed bile, and/or high gastric acid secretion (Goldman and Rosoff, 1968; Haglund, 1990). Stress elevates glucocorticoids that can increase vasoconstriction and promote clotting, and thus ulcer development (Loughlin, 1994). Glucocorticoids also have potent anti-inflammatory and immunosuppressive effects (Munck et al., 1984; Sapolsky et al., 2000), potentially allowing pathogens like *Helicobacter* to proliferate and cause gastric lesions. Thus, stress gastric ulcers and *Helicobacter*-associated gastric ulcers may not be mutually exclusive in sea otters.

Gastric ulcers have been reported in necropsied southern sea otters since the 1960s, and captive sea otters with suspected gastric lesions have been treated medically with various approaches (Mattison and Hubbard, 1969; Williams and Davis, 1995). During the 1989 Exxon Valdez Oil Spill in Alaska, oiled northern sea otters (*Enhydra lutris kenyoni*) with suspected gastrointestinal ulcers were treated via stress reduction (i.e., reduced human interactions) and the antacid cimetidine (Williams and Davis, 1995). Cimetidine may be non-ideal, since it interacts with cytochrome p450 and may have adverse effects on hepatic petroleum hydrocarbon detoxification (Williams and Davis, 1995). Other treatment options include histamine H2-receptor antagonists (e.g., famotidine) and an antibiotic (e.g., procaine penicillin, amoxicillin, or metronidazole), which are routinely given to sick or injured southern sea otters in rehabilitation to prevent gastric ulcers (Dr. Michael Murray—Monterey Bay Aquarium, pers. comm.). Antibiotics and gastric relief medications are typically used in tandem to treat humans and ferrets with *Helicobacter* infections and gastric illnesses. Studies investigating medical treatments for ferret *H. mustelae* determined that a combination of drugs was the most effective mode of bacterial eradication, including a triple therapy consisting of amoxicillin, metronidazole, and bismuth subsalicylate (Otto et al., 1990). Bismuth subsalicylate triple therapy with amoxicillin and metronidazole cleared current *H. mustelae* infections and decreased gastritis in 100% (9/9) of ferrets tested (Czinn et al., 1996). Triple therapies are commonly used to treat human *H. pylori* infections with an increasing shift to quadruple therapy due to antibiotic resistance (Graham and Fischbach, 2011). Similar triple therapies as those used for *H. pylori* and *H. mustelae* could be explored for treating *H. enhydrae*.

A prior study reported *H. enhydrae* sensitivity toward two antibiotics, cephalothin and nalidixic acid, at 30 µg each (Shen et al., 2017). Our findings for nalidixic acid agree with the previous reports, but we did not test cephalothin due to discontinuation of the ETEST® strips by the manufacturer. We expanded the antibiotic profile to evaluate the best antibiotic choices and dosages that could be used against *H. enhydrae*. Clarithromycin and tetracycline were effective at the lowest dose. Since procaine penicillin and metronidazole had been used on aquaria sea otters (Dr. Michael Murray—Monterey Bay Aquarium, pers. comm.), it would be beneficial if additional studies determined their MICs against *H. enhydrae*.

Although our work has advanced our understanding of *Helicobacter* spp. in sea otters, some unanswered questions remain. It is unknown if *H. enhydrae* has multiple strains with varying virulence and antibiotic sensitivities, like *H. pylori*. There have been no in vivo studies on *H. enhydrae*, and murine models could be useful for experimental validation of whether or not this bacterium causes gastric pathogenesis. Fecal–oral transmission of *Helicobacter* spp. through poor sanitation and water quality is a human health concern and is a likely route of exposure for sea otters in close quarters in aquaria and/or rehabilitation. *Helicobacter* spp. have recently been isolated and sequenced from 12/22 fecal samples of marine zoo mammals in Belgium (De Witte et al., 2018). PCR analysis of sea otter fecal matter and survival of *H. enhydrae* in seawater will help elucidate whether this is a potential route of transmission for this species. An immunohistochemistry (IHC) stain for *H. enhydrae* would expedite future diagnosis. We used a *H. pylori* IHC stain on known positive and negative *H. enhydrae* cases from Shen et al. (2017) but results were inconclusive.
In summary, our results provide evidence of Helicobacter DNA in gastric samples of southern sea otters and a positive association between Helicobacter detection and gastric ulcers. Fresh frozen gastric tissue, rather than formalin-fixed tissue, is the recommended sample type for qPCR detection of Helicobacter spp. in sea otters. Furthermore, these bacteria are sensitive to antibiotics in vitro, and if future studies suggest clinical intervention is warranted, this information can help guide treatment choices. These findings provide evidence that Helicobacter spp. may be an important component of sea otter gastric health and disease and support the need for additional research to characterize the population impact on this threatened marine species.

DATA AVAILABILITY STATEMENT

All datasets presented in this study are included in the article/Supplementary Material.

ETHICS STATEMENT

All work was performed in accordance with Section 109(b) of U.S. Marine Mammal Protection Act (MMPA), U.S. Fish and Wildlife Service (Service) regulations implementing the MMPA at 50 CFR 18.22(a), and in accordance with Service regulations implementing the U.S. Endangered Species Act at 50 CFR 17.21(c)(3).

AUTHOR CONTRIBUTIONS

FB designed the research project, conducted the experiments, and wrote the manuscript. FB and MEM conducted the statistical analyses. MAM provided sea otter gastric samples, oversaw histology slide interpretation, and verified histology results. ZS and JF provided H. enhydrae isolates and unstained histology slides. KO provided E. coli isolates, access to research infrastructures, and guidance on experimental design. All authors reviewed, edited, and approved the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2020.00413/full#supplementary-material
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