Naval vessels regularly mix fuel and seawater as ballast, a practice that might exacerbate fuel biodegradation and metal biocorrosion. To investigate, a metagenomic characterization and metabolite profiling of ballast from U.S. Navy vessels with residence times of 1-, ~20-, and 31 weeks was conducted and compared with the seawater used to fill the tanks. Aerobic *Gammaproteobacteria* differentially proliferated in the youngest ballast tank and aerobic-specific hydrocarbon degradation genes were quantitatively more important compared to seawater or the other ballast tanks. In contrast, the anaerobic *Deltaproteobacteria* dominated in the oldest ballast fluid with anaerobic-specific hydrocarbon activation genes being far more prominent. Gene activity was corroborated by detection of diagnostic metabolites and corrosion was evident by elevated levels of Fe, Mn, Ni and Cu in all ballast samples relative to seawater. The findings argue that marine microbial communities rapidly shift from aerobic to anaerobic hydrocarbonoclastic-dominated assemblages that accelerate fuel and infrastructure deterioration.

**RESULTS**

**Ship ballast and seawater**

Prior to sampling, interviews with navy personnel revealed that crews retained ballast fluids on board for varying lengths of time. Since the ships were of the same class (*Arleigh Burke* destroyers), possessed seawater-compensated carbon steel ballast tanks, were stationed in the same harbor, and used the same seawater and fuel as ballast, comparison between ships as a function of ballast residence time seemed pragmatic. Nevertheless, variations in vessel operating practices, distances traveled, fuel use patterns, and other between-vessel differences were also evident. Thus, the harbor water and a research vessel that possessed carbon steel ballast tanks but did not mix fuel and seawater (Ship NF—no fuel) for ballast served as comparative controls.

---

1Department of Microbiology & Plant Biology and OU Biocorrosion Center, University of Oklahoma, Norman, OK 73019, USA. 2School of Civil Engineering and Environmental Science and OU Biocorrosion Center, University of Oklahoma, Norman, OK 73019, USA. 3Department of Physics, Barnard Hall 264, Montana State University, Bozeman, MT 59717, USA. 4Biology Department, University of Texas of the Permian Basin, Odessa, TX 79762, USA. 5email: jsulfita@ou.edu

Published in partnership with CSCP and USTB.
Characterization of fluid samples
Harbor water and ballast tank fluids of varying shipboard residence time were obtained from three naval vessels as well as from ship NF. The samples were analyzed for potential electron acceptors and selected dissolved metals (Table 1). The shortest residence time ballast water (Ship #1) contained ~50% less dissolved oxygen than the San Diego harbor, but its sulfate concentration was comparable to both the seawater and Ship #2. In contrast, the ship with the longest ballast residence time (Ship #3) exhibited 4 mM sulfate depletion in 8 mo. Dissolved Mn, Cu, Fe, and Ni concentrations in the ballast waters were substantially higher than in the harbor water. After 1 week of residence (Ship #1), these metals increased ~4.5–15-fold (Table 1). Dissolved Cu and Ni levels were much higher than Mn in Ships #1 and #2, but Ship #3 had the opposite trend with Mn being 10-fold higher than the other elements. In all ships, particles were mostly copper sulfides, with a smaller percentage of nickel sulfides, and an even smaller fraction of iron sulfides. In a few cases, both copper and nickel sulfides were present in the same particle but manganese sulfides were not detected (Table 1). Comparison of the metals in the ballast tanks relative to the harbor water suggests that the intake fluids accounted for a negligible fraction of the shipboard dissolved metal status and implies metal dissolution through corrosion as the more likely explanation. The ballast of a comparator ship that did not mix fuel and seawater (Ship NF) contained higher levels of Cu and Ni than the harbor water, but less dissolved and particulate metal than the other vessels.

Marine populations and community composition
Microbial populations in the harbor and samples from the Navy ships were quantified by direct microscopy, domain-specific qPCR, and single-copy marker gene quantification based on the metagenomic analysis (Table 2). Insufficient DNA was recovered from Ship NF for metagenomic sequencing without amplification which would have biased the findings and negated comparability. Direct counting revealed that the microbial population size was comparable in all Navy ships (10^6 cells mL^-1), independent of ballast residence time and between 2 and 10-fold less than the harbor community. Quantitative PCR enumeration of bacterial and archaeal 16S rRNA genes ranged from 10^5–10^7 and 10^2–10^3 gene copies mL^-1, respectively. By this measure, the Bacteria bloomed after 1 wk aboard Ship #1 relative to the harbor community but returned to near comparable levels with increased ballast residence time as in Ships #2 and #3 (Table 2). Archaea were several orders of magnitude less abundant than Bacteria in all samples. The largest picoeukaryote populations were found in the harbor water, whereas sharply decreased numbers were found in all ballast samples.

Microbial diversity indices, including the Shannon index, observed OTUs, and Simpson’s evenness metric (Table 3) indicate that all communities showed a bias towards some taxa (e.g. more uneven distribution of species) and a trend of increasing dominance of particular microbial types with increasing ballast tank residence time. There is also a limited change in the range of species richness (e.g. observed species) of only 8% between Ship #1 and Ship #2.

The composition of microbial populations within each sample was determined by comparison of each 16S rRNA gene library extracted from the metagenome samples. The archaeal 16S rRNA gene libraries ranged from 10^5–10^6 gene copies mL^-1, independent of residence time. The bacterial 16S rRNA genes ranged from 10^5–10^6 gene copies mL^-1.

| Table 1. Summary of water sample residence times (reported or estimated from shipboard personnel) and selected chemical properties. |
|---------------------------------------------------------------|
| **Residence time (weeks)** | **Dissolved O_2 (mg per liter)** | **Sulfate (mM)** | **Manganese (µg per liter)** | **Iron (µg per liter)** | **Nickel (µg per liter)** | **Copper (µg per liter)** |
| Harbor | Ship NF | Ship #1 | Ship #2 | Ship #3 | Ship #1 | Ship #2 | Ship #3 |
|---|---|---|---|---|---|---|---|
| 0 | >24 | 1 | ~20 | 32 | 7 | NR | ND |
| 7 | NR | 3.7 | ND | ND | 29 | 29 | 25 |
| 16.2 (NR) | 10.7 (ND) | 125.1 (ND) | 1119.3 (ND) | 119.3 (ND) | 95.8 (26%) | 1564.3 (11%) | 23.9 (39%) |
| 10.1 (NR) | 10.7 (ND) | 125.1 (ND) | 1119.3 (ND) | 1119.3 (ND) | 106.3 (47%) | 864.5 (42%) | 101.2 (85%) |

The dissolved metal content is given in concentration units and the relative frequency of the EDX-analyzed metal sulfides are shown in parentheses. Dissolved oxygen in the harbor was estimated from data presented in ref.73, ND not detected, NR Not reported.

| Table 2. Marine microbial population estimates and functional gene qPCR enumeration*. |
|-----------------------------------------------|
| **Cell Counts** | **Bacteria** | **Archaea** | **Picoeukaryotes** | **apsA** | **dsrA** |
| **Harbor** | **Ship #1** | **Ship #2** | **Ship #3** | **Ship #1** | **Ship #2** |
|---|---|---|---|---|---|
| 2 x 10^6 | 1 x 10^5 | 7 x 10^5 | 9 x 10^5 | BDL | BDL |
| 1.05 x 10^6 (1.95 x 10^5) | 2.28 x 10^6 (1.56 x 10^5) | 3.67 x 10^5 (5.45) | 2.34 x 10^5 (9.69) | 9.69 x 10^5 (3.10) x 10^5 |
| 6.44 x 10^7 (4.64 x 10^7) | 1.93 x 10^7 (7.79 x 10^7) | 9.78 x 10^6 (3.02 x 10^6) | 5.84 x 10^7 (3.72 x 10^7) | 3.38 x 10^8 (5.88 x 10^6) |
| 6.44 x 10^7 (4.64 x 10^7) | 1.93 x 10^7 (7.79 x 10^7) | 9.78 x 10^6 (3.02 x 10^6) | 5.84 x 10^7 (3.72 x 10^7) | 3.38 x 10^8 (5.88 x 10^6) |

*Cell counts reported as cells mL^-1, Bacteria, Archaea and Picoeukaryote represent respective 16S and 18S rRNA genes. Adenosine-5’-phosphosulfate reductase A gene (apsA). Dissimilatory sulfite reductase A gene (dsrA). Below detection limit (BDL). Standard deviations are shown in parentheses for qPCR analyses. Comparable data from Ship NF were not collected.
accounted for substantive proportions of the communities within the harbor water and Ships #1, #2 and #3, respectively.

A decrease in number of OTUs was observed from the harbor to Ship NF to Ships #1, 2, and 3 (Supplementary Table 1, analysis of 16S amplicon libraries). All ballast tank samples had much lower Chao1 richness scores than did the harbor sample (Supplementary Table 1). Data from 16S rRNA amplicon libraries gave similar trends in relative abundance for most taxa (Supplementary Fig. 1) as those derived from metagenome sequencing for samples from the harbor and ships containing fuel. The sample from a ship whose ballast tank did not contain fuel (Ship NF) was most similar to the harbor sample but with approximately half the relative abundance of most taxa (Supplementary Fig. 1) as the other ships (Fig. 1). Sequences classified as Archaea were relatively minor and never exceeded 0.3% in any shipboard community.

The Gammaproteobacteria were enriched in the Ship #1 ballast tank assemblage to become the dominant taxonomic class, with the concomitant decline of the Alphaproteobacteria and Flavobacteria. Alteromonadales, Oceanospirillales, and Thiorthricales increased to account for 42%, 35%, and 19% of all observed Gammaproteobacteria, respectively (Fig. 1b). Epsilonproteobacteria were also enriched 10-fold (2.8%) after one week in the ballast tank of Ship #1 and were represented almost exclusively (97%) by members of the Campylobacteriales. Deltaproteobacteria were the dominant Class observed in the Ship #2 community at 21.0%, a ~3.5-fold increase relative to Ship #1. Members of the Desulfomonadales and Desulfovibrionales accounted for the majority of the Deltaproteobacteria at 80 and 10%, respectively (Fig. 1b). Though the total proportion of Gammaproteobacteria was smaller in Ship #2 vs. Ship #1, the Alteromonadales were enriched to represent 71% of this taxon (Fig. 1b). Taxa affiliated with the Clostridia (7.2%) and Mollicutes (4.6%) accounted for substantive portions of the total ballast tank community in Ship #2, with increases of ~3-fold and ~5-fold relative to Ship #1. The Alphaproteobacteria, Flavobacteria, and Epsilonproteobacteria were all slightly higher in relative abundance in the Ship #2 community relative to Ship #1.

The ballast tank community of Ship #3, with the longest residence time, was the most different from the harbor water assemblage (Fig. 1a). Deltaproteobacteria (43.3%) was the most abundant taxon, with members belonging to the Desulfobacteriales, Desulfarculales, and Desulfovibrionales at 66%, 27%, and 6%, respectively (Fig. 1b). This community also contained the largest population of Bacteroidetes (8.2%), represented predominantly by unclassified members of the Bacteroidales (57% of the Bacteroidetes), and the smallest segment of Flavobacteria observed in any sample. Gammaproteobacteria, Alphaproteobacteria, and Alteromonadales at 8.9% and 4.6% total relative abundance, respectively (Fig. 1b). The harbor contained several groups of marine Eukaryota and were dominated by Alveolata (11.0%) and Stramenopiles (4.0%). Despite accounting for ~20% of the total marine community in the harbor, the Eukaryota were substantially reduced in the ballast tank assemblages (Table 2). The decline after only a week of residence time was also reflected in the reduced level of eukaryotic rRNA gene sequences detected in the other ships (Fig. 1). Sequences classified as Archaea were relatively minor and never exceeded 0.3% in any shipboard community. The Gammaproteobacteria were enriched in the Ship #1 ballast tank assemblage to become the dominant taxonomic class, with the concomitant decline of the Alphaproteobacteria and Flavobacteria. Alteromonadales, Oceanospirillales, and Thiorthricales increased to account for 42%, 35%, and 19% of all observed Gammaproteobacteria, respectively (Fig. 1b). Epsilonproteobacteria were also enriched 10-fold (2.8%) after one week in the ballast tank of Ship #1 and were represented almost exclusively (97%) by members of the Campylobacteriales. Deltaproteobacteria were the dominant Class observed in the Ship #2 community at 21.0%, a ~3.5-fold increase relative to Ship #1. Members of the Desulfomonadales and Desulfovibrionales accounted for the majority of the Deltaproteobacteria at 80 and 10%, respectively (Fig. 1b). Though the total proportion of Gammaproteobacteria was smaller in Ship #2 vs. Ship #1, the Alteromonadales were enriched to represent 71% of this taxon (Fig. 1b). Taxa affiliated with the Clostridia (7.2%) and Mollicutes (4.6%) accounted for substantive portions of the total ballast tank community in Ship #2, with increases of ~3-fold and ~5-fold relative to Ship #1. The Alphaproteobacteria, Flavobacteria, and Epsilonproteobacteria were all slightly higher in relative abundance in the Ship #2 community relative to Ship #1.

The ballast tank community of Ship #3, with the longest residence time, was the most different from the harbor water assemblage (Fig. 1a). Deltaproteobacteria (43.3%) was the most abundant taxon, with members belonging to the Desulfobacteriales, Desulfarculales, and Desulfovibrionales at 66%, 27%, and 6%, respectively (Fig. 1b). This community also contained the largest population of Bacteroidetes (8.2%), represented predominantly by unclassified members of the Bacteroidales (57% of the Bacteroidetes), and the smallest segment of Flavobacteria observed in any sample. Gammaproteobacteria, Alphaproteobacteria, and

![Fig. 1 Metagenome analysis of community composition in navy ship ballast systems and the San Diego Harbor seawater. Class-level depiction (a) and Order-level (b) comparison for the Delta- and Gammaproteobacteria.](image-url)
Epsilonproteobacteria also accounted for smaller community abundance in Ship #3 relative to Ship #2.

The trends in relative proportions seen in the metagenome data were supported by the proportions of aerobic alkane degraders, Marinobacter in particular (Supplementary Table 2) as well as of Deltaproteobacteria (Supplementary Table 3) found in the 16S rRNA amplicon libraries. In particular, relative abundances in Ship NF of Marinobacter and Deltaproteobacteria were similar to those in the harbor and less than in the three ships that used fuel in ballast tanks.

Functional genes associated with the utilization of common electron donors and acceptors were characterized by translated Blast analysis (Fig. 2). The nitrification genes, ammonia mono-oxygenase (amoA) and hydroxylamine dehydrogenase (hao) did not exceed 0.3% normalized abundance within any sample library. Genes associated with the oxidation of reduced inorganic sulfur species were present within all of the samples analyzed. Sulfur-oxidizing system (SOX) protein B (soxB) declined from 5.9% in the harbor water metagenome to ~4.2% of the functional capacity to Ships #1 and #2 and to barely detectable levels in Ship #3 (0.5%) (Fig. 2). Nitrogen fixation nitorgenase gene (nifH) generally increased with ballast water residence time from a low in the harbor water (0.9%) to 5.5%, 7.1%, and 8.8% of the libraries for Ships #1, #2, and #3, respectively.

Genes affiliated with the reduction of oxygen and nitrate exhibited a marked increase between the harbor water and Ships #1 and #2. The aerobic respiration genes cytochrome c oxidase subunit I (ccoN and coxA) had the highest levels of any inorganic redox genes assayed in the harbor and ballast tank waters of Ships #1 and #2. However, there was a 50% decrease in cytochrome c oxidase subunit I gene abundance after 8 months in Ship #3. Nitrate reductase genes (cytoplasmic narG and periplasmic napA) had a combined abundance that markedly increased in ballast tank waters of Ships #1 (22.1%) and #2 (38.0%) relative to the harbor water (9.0%) and then declined to 11.3% in the Ship #3 library. The potential for sulfate and thiosulfate reduction was investigated through an assessment of the adenylylsulfate reductase (aprA) and thiosulfate reductase (phsA/prsA) genes, respectively. Total sulfidogenesis potential was marked by the abundance of the dissimilatory sulfite reductase gene, dsrA. All genes associated with the reduction of sulfuric anions remained relatively low levels in all samples excluding Ship #3. Between Ships #2 and #3, there were increases of 4-fold in aprA, 5-fold in phsA, and 5-fold in dsrA genes. These were the highest levels observed for each of these genes within this study and represented increases relative to the harbor water library ranging between 3.6-fold for adenylylsulfate reductase to 30-fold for thiosulfate reductase.

Evidence for hydrocarbon biodegradation

Normalized gene abundances (Fig. 3) and observed metabolic intermediates (Fig. 4) associated with the activation and biodegradation of diesel fuel components were readily evident in the shipboard ballast tank systems. Oxygen-dependent mechanisms for the activation of diesel hydrocarbons were assayed by quantifying the proportion of genes annotated as
alkane 1-monooxygenase (alkB) for aliphatic substrates, ethylbenzene dioxygenase (etbAa), toluene monooxygenase (tmoA), benzene/toluene dioxygenase (todC1), and xylene monooxygenase (xylM) for BTEX hydrocarbons, and naphthalene 1,2-dioxygenase (nahAc) and 1-methylnaphthalene hydroxylase for polyaromatic hydrocarbons. Phenol hydroxylase (dmpL/poxC), phenol 2-monooxygenase, catechol 1,2-dioxygenase (catA) and catechol 2,3-dioxygenase (catE), along with salicylaldehyde dehydrogenase and salicylate hydroxylase were used as metrics for the genetic potential for the aerobic microbial metabolism of mono- and polyaromatic hydrocarbons. Alkyl-, benzyl-, and naphthylmethyl-succinate synthases (assA, bssA, nmsA) served as indicator genes for anaerobic activation of aliphatic, mono-, and polyaromatic hydrocarbons by addition to fumarate. The ethylbenzene dehydrogenase gene ebdA served as a marker for the potential of anaerobic activation of ethylbenzene by dehydrogenation and subsequent hydroxylation of the ethyl-substituent.

Alkane monooxygenase (12.4%) was the only hydrocarbon activation gene that constituted >2.0% of the total harbor water community potential. Genes associated with the anaerobic activation of naphthalenes were <1%. Genes for the alkyl- (2.1%), monooaromatic- (0.3%), and naphthylmethyl- (1.2%) succinate synthases all showed slight increases in Ship #1 samples relative to the harbor water metagenome, but these values were perhaps biased by the presence of other common glycyl-radical enzymes (e.g. pyruvate-formate lyase) that share substantive protein sequence homology. Despite these slight gene enrichments, no succinate-derivative metabolites were detected in Ship #1 samples.

Analysis of the community in Ship #2 revealed a further 11.6% enrichment of alkB possessing taxa relative to Ship #1. In contrast, the genes for ethylbenzene dioxygenase, xylene monooxygenase, and phenol hydroxylase declined sharply and each accounted for less than 2.0% of the total community metagenome. The abundance of the nmsA gene, coding for the anaerobic activation of methylnaphthalene, represented 7.7% of the total gene pool in Ship #2. This was an increase of the same gene abundance relative to Ship #1 of 440.1%. Overall, the total proportion of hydrocarbonoclastic taxa in Ship #2 decreased compared to Ship #1, though there was an enrichment in oxygen-independent activation genes (Fig. 5).

The Ship #3 ballast tank contained the largest hydrocarbon-activating community of any sample investigated. Alkane 1-monooxygenase (6.1%) and catechol dioxygenase (catA/catE) genes (0.4%, 2.0%) decreased to their lowest relative proportion observed in any sample. In contrast, the community possessing assA (5.7%), bssA/hbsA/ibsA (4.7%) and nmsA (20.5%) were found in

Fig. 4  Analysis of fuel metabolites within Ship #1 and Ship #3 ballast tank fluids. Blue boxes indicate compounds identified in Ship #1 only, red in Ship #3 only, and black were observed in both samples. Numbers above arrows correspond to transformations catalyzed by metabolic genes listed in Fig. 3.
the highest respective abundances (Fig. 5). Mono- and polyaromatic succinate metabolites were present corresponding to alkylated aromatic hydrocarbon substrates (Fig. 4). No catechols or aromatic dihydrodiol intermediates were measured in the Ship #3 water sample. Naphthoic and alkanolic acids, aliphatic alcohols, and phenylethanol were also putatively identified in Ship #3 and Ship #1 waters (Fig. 4).

**DISCUSSION**

The study of seawater-compensated fuel ballast tanks holding fluids for varying lengths of time provides insight into the temporal dynamics of microbial community composition changes and associated metabolic activity shifts through metagenomic and metabolite profiling analyses coupled with geochemical assessments. Ribosomal RNA gene analysis revealed the surface waters of San Diego Harbor to be primarily colonized by *Bacteria* belonging to the *Alphaproteobacteria*, *Flavobacteria*, and *Gammaproteobacteria* and marine eukaryotes (Fig. 1a). The harbor receives terrestrial discharges from urban, industrial, and military installations in the surrounding metropolitan area. The bacterioplankton community in San Diego harbor resembles the northern Gulf of Mexico profiles that were dominated by *Alphaproteobacteria* and *Bacteroidetes*, with ratios of *Alpha* to *Gammaproteobacteria* of <1.715. The comparable ratio in the San Diego Bay of 1.4 no doubt reflected the complex natural and anthropogenic inputs enriching for marine heterotrophic taxa that differ substantively from open ocean surveys.

The microbial community composition was significantly altered after a one-week residence time in a Navy ballast tank (Ship #1) relative to the harbor assemblage. Despite the total cell density decrease of ~100-fold (Table 2), there was only a slight decrease in community diversity and an increase in the observed taxa representing the OTU (Table 3). *Gammaproteobacteria* were the dominant community members, predominantly driven by relative increases in the Orders *Alteromonadales*, *Oceanospiralles*, and *Thiotrichales* (Fig. 1b). These bacterial Orders contain numerous well-documented hydrocarbonoclastic aerobic taxa belonging to the genera *Marinobacter*, *Alcanivorax*, and *Cycloclasticus*, respectively. Previous studies noted the enrichment of these groups in marine waters impacted by oil spills across the globe20-23.

*Marinobacter hydrocarbonoclasticus* preferentially metabolizes short to moderate chain length ($C_{n}$-$C_{11}$) n-alkanes and selected monoaromatic hydrocarbons25. Pure cultures of *M. vinificus* and *M. hydrocarbonoclasticus* were able to catalyze the complete oxidation of BTEX hydrocarbons under aerobic conditions24. Marine *Alcanivorax* spp. are known to aerobic biodegrade a wide range of aliphatic petroleum constituents25-28, but are thought incapable of aromatic hydrocarbon decay29,30. However, recent work suggests that *Alcanivorax* may also contribute to the biodegradation of PAHs31. *Cycloclasticus pugetii* is an obligate aerobe with the capacity to metabolize mono- and polynuclear aromatic hydrocarbons (PAHs)32. *Cycloclasticus* strains isolated from marine coastal waters20,21 were also associated with the aerobic biodegradation of a variety of alkylated PAHs and dibenzothiophenes. Genetic studies revealed the presence of distinct naphthalene and biphenyl cluster dioxygenases in several *Cycloclasticus* isolates20. The increased proportions of *Alteromonadales* and *Thiotrichales* likely account for the enrichment of aromatic hydrocarbon degradation genes and the presence of signature aerobic metabolites (e.g. alkylcatechols and dihydroxynaphthalene) observed in Ship #1 (Figs. 3, 4). Phylogenetic, genomic, and metabolic evidence suggests that the ballast tank community of Ship #1 was enriched in taxa responsible for the aerobic oxidation of the more water-soluble aromatic hydrocarbon components of diesel fuel.

Ship #2 was predominantly colonized by members of the *Alteromonadales*, but also had an increase in *Deltaproteobacteria*, specifically the *Desulfuromonadales* (Fig. 1b). In concert with the decreased relative abundance of the *Gammaproteobacteria*, there was also a decrease in the aerobic hydrocarbon-activation genes relative to Ship #1 (Fig. 3). Despite the bloom in *Desulfuromonadales*, there was surprisingly little enrichment in genes associated with sulfido genesis (Fig. 2). Anaerobic hydrocarbon-activation genes exhibited only a slight increase in *nmsA* abundance relative to Ship #1 (Fig. 3). The *Desulfuromonadaeceae* are strictly anaerobic heterotrophs, with some aerotolerance capacity, that typically respire sulfur compounds to sulfide, with the notable exception of several genera that are almost exclusively fermentative29. Members of this Family typically utilize fatty acids and alcohols as growth substrates, but hydrocarbon metabolism is not widely distributed34. However, several species of *Desulfuromonas* are
known to reduce Fe(III) and Mn(IV) as electron acceptors. The oxygen tolerance and ability to utilize metals as electron acceptors, suggests that these taxa bloomed in response to increasing scarcity of oxygen in Ship #2 ballast tank. In the absence of metabolite or genomic evidence suggesting direct fuel biodegradation, it is likely that aerobic catabolites produced by the cohabitating *Alteromonas* helped provide the requisite electron donors and carbon sources.

After 32 weeks aboard ship, the ballast tank community of Ship #3 was predominantly anaerobic taxa affiliated with *Deltaproteobacteria* (42%), specifically the *Desulfo bacteraceae* (27%) and *Desulfurlactaceae* (11%) (Fig. 1a). There was also a concomitant increase in sulfuroxygenic reductive respiration genes to the highest levels observed in this study (Fig. 2). Conversely, the Ship #3 community possessed the smallest population of c-type cytochrome oxidase containing taxa (Fig. 2). Metagenome analysis estimated that 27.0% of the genomes encoded an adenylylsulfate cytochrome oxidase containing taxa (Fig. 2). This community also possessed the highest levels of anaerobic hydrocarbon activation genes, including alkylsuccinate synthase (5.7%), monoaromatic succinate synthases (4.7%), and naphthalene methylsuccinate synthases (20.5%). Pure cultures and enrichments containing *Deltaproteobacteria* able to anaerobically metabolize a variety of normal, iso- and cyclic alkanes, as well as mono- and polycyclic aromatic hydrocarbons were recently reviewed. The cultures were mostly derived from marine environments and could couple the oxidation of their parent hydrocarbon with the reduction of sulfate and production of sulfide. Indeed, the sulfate concentration in Ship #3 was noticeably lower than in the harbor or any of the other ballast tanks (Table 1).

The mechanisms of hydrocarbon activation by *Deltaproteobacteria* are fundamentally different from hydrocarbonoclastic aerobes in that molecular oxygen is not a co-reactant. There are at least four biochemical strategies governing the anaerobic attack on hydrocarbons. The most commonly reported pathway is the activation of the parent substrates via fumarate addition reactions. There was an increase in genes encoding this pathway in Ship #3 and many of the succinyl derivatives associated with such bioconversions were found as signature metabolites in this sample (Figs. 3, 4).

From an ecological perspective, fuel-compensated ballast tanks are microcosms of other hydrocarbon-laden marine ecosystems. After a short exposure (1 week) to petroleum in the absence of light, the coastal marine community based on phototrophic primary production shifted to one supported by aerobic heterotrophic respiration of diesel hydrocarbons and an almost complete loss of the eukaryotic life forms. Ship NF did not show an increase in the relative abundance of hydrocarbonoclastic aerobes. The resulting hydrocarbonoclastic community of Ship #1 is reminiscent of the successional patterns observed during the response of open-ocean surface waters to the influx of petroleum from the Deep Water Horizon oil spill. Here too, surface water assemblages underwent shifts within the first week and experienced strong enrichments in hydrocarbonoclastic aerobes. The shift in community proportions encoding hydrocarbon activation genes, as well as the relative abundance of genes from oxygen-dependent and independent pathways also changed as a function of time (Fig. 5). After one week, the hydrocarbonoclastic community containing aerobic biodegradation genes was enriched 2.4 times that observed in the harbor. At 20 weeks and beyond, the abundance of the same genes decreased to only about 1.5 times that of the harbor assemblage. Within Ship #1, 95% of the activation genes were associated with aerobic mechanisms associated with well-documented aerobic hydrocarbonoclastic taxa, aerobic respiration genes, signature aerobic catabolic intermediates, and a decreased dissolved oxygen content (Figs. 1, 2, 4, and 5; Table 1). As ballast residence time increased, the fuel-degrading activation gene subset was increasingly affiliated with aerobic biodegradation mechanisms. At 32 weeks, the hydrocarbon-degrading community predominantly employed oxygen-independent mechanisms (68%) with key metabolites associated with the fumarate addition-mediated degradation of BTEX and PAH fuel components (Figs. 4, 5).

Biodegradation activity was associated with the corrosion of the ballast tanks. Dissolved metal concentrations within each Navy ballast tank indicate corrosion of the carbon steel and/or the copper-nickel sluice pipes as the values are considerably greater than ship NF or the surrounding harbor waters (Table 1). Dissolved manganese, copper, and nickel were positively correlated with the increase in sulfidogenic taxa between Ships #1 and #2 (Fig. 5). There was also a strong increase in the abundance of sulfide-bearing particulates of each metal species associated with the clearly sulfidogenic community of Ship #3 (Supplementary Fig. 2). The influence of biogenic sulfide in Ship #3 on the dissolved metal concentrations is demonstrated by the inverse correlation between the relative frequency of metals observed in metal-sulfide particles (Cu > Ni > Fe > Mn) and the solubility equilibrium constant ($K_s$) for the reaction:

$$\text{X}_\text{S} + 2\text{H}^+ \leftrightarrow \text{X}^{2+} + \text{H}_2\text{S}$$

where $X = \text{Cu}^{2+}, \text{Ni}^{2+}, \text{Fe}^{2+},$ and $\text{Mn}^{2+},$ with log $K_s$ values ranging from $-15.2$ for CuS, $1.5$ for NiS, $3.0$ for FeS, and $10.0$ and $7.0$ for MnS rammberite and alabandite minerals, respectively. Localized corrosion in low carbon steels in marine environments shows that pit initiation begins at the immediate surroundings of MnS inclusions where strained interfaces act as anodes and the surrounding unstrained iron as cathodes. Hydrolysis in the surrounding region acidifies the local region releasing Mn ions into solution. The increasing dissolved manganese concentration with increasing residence time illustrates ongoing corrosion of carbon steel. Despite increased sulfidic conditions in Ship #3, the predicted solubilities of alabandite and rammberite phase MnS compounds were not exceeded (12.97 mg per liter—alabandite phase; 499.40 mg per liter—rammberite phase) explaining the lack of MnS particles in the samples. Corrosion of carbon steel in Ships #2 and #3 is further demonstrated by the presence of dissolved iron. In Ship #3, the measured dissolved iron concentration (603.0 µg per liter) is nearly equivalent to the calculated equilibrium concentration (441.20 µg per liter) presuming the simultaneous precipitation of FeCO$_3$ and FeS minerals. Copper-nickel alloys are commonly utilized as marine infrastructure, but are highly susceptible to corrosion and leaching by sulfides, organic acids and other products of microbial metabolism. Previous studies have shown strong correlations between hydrocarbon biodegradation-driven sulfidogenesis and both general- and pitting corrosion for carbon steel and copper-nickel alloys in incubations of San Diego Bay seawater and F-76 diesel fuel. Analysis of metal concentrations and particulates suggests that the CuNi-alloy sluice pipes transporting water and fuel between ballast tanks were arguably the primary sites of active biocorrosion in Ships #1 and #2 and that aggressive pitting corrosion of the ballast tank carbon steel was occurring in Ship #3. The concentrations of dissolved copper and nickel in Ship #1 and Ship #3 were greater than the equilibrium solubility of covellite (CuS) (3.88 x 10$^{-7}$ µg per liter) and α-NiS (19.4 µg per liter), suggesting in these ships, the copper-nickel slice pipes initially had an oxidized coating of copper hydroxides and/or carbonates (solubility 9–85 µg per liter) and nickel oxide and/or carbonate (solubility 30–1033 mg per liter). Microbial biofilm formation on copper and copper-nickel alloys can facilitate loss of the passivating layer of copper and nickel oxides and carbonates thereby providing a second mechanism for release of dissolved copper and nickel. Additionally, a thermodynamic model based on the premise of microbial sulfidogenesis in the presence of various metals/alloys suggests that sulfides will
react with metal oxides to produce metallosulfide particles, thus exacerbating the depletion of the protective oxide layers.\(^\text{50}\) Regardless of the mechanisms for corrosion, as sulfidogenic activity increases from Ship #2 to Ship #3, a decrease in the concentration of dissolved copper and nickel occurs due to the formation and precipitation of sulfide particulates.

This study has important implications for the operation and maintenance of fuel compensated ballast tanks of the U.S. Navy. Petroleum-derived diesel fuels are susceptible to biodegradation when in contact with seawater, regardless of oxygenation status or residence time and shipboard infrastructure is negatively impacted by microbiologically influenced corrosion minimally driven by the production of sulfides. Studies conducted on simulated ballast tank systems highlighted the inherent stratified nature of these habitats with zones of variable oxygenation.\(^\text{51,52}\)

The deleterious impacts of microbial activity were not homogeneously distributed within the ballast systems and tank bottoms tended to be sites of aggressive general and localized corrosion.\(^\text{10,15,53,54}\) Ship operators have estimated that a vessel on standard operations consumes \(\sim1000\) gallons of fuel per day, necessitating the introduction of fresh seawater into ballast tanks on a regular basis (personal communication). Cyclic oxygenation/deoxygenation has repeatedly been observed to promote the highest rates of generalized and pitting corrosion.\(^\text{55}\) Furthermore, this condition promotes the persistence of biofilms harboring both aerobic and anaerobic taxa with the capacity for sulfur cycling and the formation of corrosive sulfur granules.\(^\text{15}\)

As treatment of incoming seawater for the removal of oxygen and sulfate is not feasible, a critical consideration might be to eliminate the introduction of sediments (and associated microbial load) into ballast tanks and to ensure all internal shipboard surfaces are coated. Sediments tend to settle onto horizontal surfaces and promote further stratification within ballast tanks and likely serve as a source of sulfide-producing microbes. Given the stratified nature of these systems, further research is warranted to determine how well the sampling of expansion tanks represents the various ballast tank zones. Ultimately, the successional trends in the ecology of ballast tank ecosystems were identified that are critical for understanding the impact of microorganisms on infrastructure and will serve to inform future studies to lessen the biological wastage of naval resources.

**METHODS**

**Sample collection**

Ballast tank fluid samples were collected in sterile polypropylene bottles from valves in the expansion tanks (Fig. 6) on the Navy vessels (Ship #1, 2, and 3). The ballast tank systems on the ships consist of carbon steel tanks with sacrificial anodes and protective coatings connected via copper-nickel alloy sluice pipes. Unlike the Navy ships, the ballast tank system on the Ship NF was not designed to minge fuel and seawater. The sample from Ship NF was pumped from the ballast tank underlying the mess area through an access hatch, preserved with DNAzol\(^\text{TM}\) (Molecular Research Center Inc.; Cincinnati, OH) as indicated below, and shipped by crew members to the laboratory for analysis. For each Navy ship, one 250 mL aliquot of ballast water was preserved with 5 mL of DNAzol for DNA extraction. Another 1 L aliquot was acidified with 6 N HCl (pH < 2) for metabolite profiling. Finally, a separate bottle was filled to capacity (>250 mL) without preservative for chemical and metals analyses. In addition, several liters of seawater were collected from the San Diego Harbor where each ship was docked for comparative analyses. All samples were shipped overnight on ice to the laboratory and stored at 4 °C until use.

**Chemical analyses**

Dissolved oxygen was measured in the field using a Firesting-O2 oxygen meter (Pyro Science GmbH; Aachen, Germany) operated according to manufacturer instructions. Nitrate and sulfate were quantified by ion chromatography as previously described.\(^\text{12}\) For the determination of dissolved metals, each sample was filtered with a 0.45 µm polyethersulfone filter (Whatman Puradisc) and diluted to a metal concentration range between 0 to 40 mg per liter using 0.1% (v/v) nitric acid (Fluka, Ultrapure).

**EDX analysis**

The samples were analyzed with a scanning field emission electron microscope (SUPRA GTA120). Sample volume injected into the graphite furnace was 10 µL with argon as purging gas. Accelerating voltage was set at 90 kV and studied for 60 s, then removed and cooled to 20 °C before going to 2400 °C for 4.8 s. At this final temperature, absorption was measured using a background correction and a spectral bandwidth slit width of 0.2 A six-point calibration curve ranging from 0 to 40 mg per liter was used for each metal examined. All samples were measured in triplicate, and each time after measuring five samples, two known calibration standards and the blank were reanalyzed with recalibration if necessary.

**Microbial enumeration**

Microbial cells were enumerated from unpreserved Navy ballast fluids by direct epifluorescence microscopy after fixing the samples with 1/10 volume of 37% formaldehyde and staining with 4,6-diamidino-2-phenylindole. Fixed cells were then filtered onto 0.22 µm, 25 mm diameter black polycarbonate membrane filters (Sterlitech, Kent WA). At least 30 fields were counted per sample and results were averaged.

**DNA extraction and qPCR analyses**

Samples preserved for DNA extraction were filtered (250 mL; 0.45 µm pore size PES filter) upon receipt, stored at −80 °C until use, and extracted from the filters as previously described.\(^\text{38}\) The same DNA extracts from the harbor sample and from Ships #1, 2, and 3 were used for metagenome, qPCR, and 16S rRNA amplicon libraries. Quantitative PCR was used to enumerate population densities of bacteria, archaea and picoeukaryotes within each sample. Bacterial and archaeal 16S rRNA genes were amplified with Eubac3F/338R and A8F/344R primer pairs.\(^\text{57}\) Marine picoeucaryote 18S rRNA genes were amplified using the 345F/499R primer pair as previously described.\(^\text{49}\) Dissimilatory sulfate reduction gene, ade nosine-5'-phosphosulfate reductase A gene (aprA) and dissimilatory sulfite reductase A (dsrA), were assessed with the RH1apsF/RH2apsR and RH1-dsrF/RH2-dsrR primer pairs, respectively.\(^\text{59}\)

**Metagenome analysis**

Shotgun metagenome libraries were sequenced on an Illumina MiSeq platform (Oklahoma Medical Research Foundation, Oklahoma City, OK). Sequencing adapters were trimmed using Cutadapt\(^\text{40}\) to a minimum average PHRED score of 30 and mate paired with Trimmomatic.\(^\text{61}\) Unpaired reads were not retained for analyses. Community composition was
assessed from unassembled mate-paired reads with significant homology to known small subunit (SSU) rRNA genes. Reads were searched against the Silva SSU Reference database release 108 pre-clustered at 97% sequence similarity using USEARCH. Hits with a minimum of 70% homology to reference SSU gene sequences were extracted for further analysis. Putative SSU reads were analyzed using QiIME 1.9.1 through closed-reference OTU clustering (97% similarity) and taxonomy assignment against the Silva 111 SSU reference sequence and taxonomy databases. Alpha diversity metrics were computed for each sample based upon metagenome reads containing SSU fragments rarefied to 1800 sequences for each library.

Functional gene annotation was performed by translated homology searching of unassembled mate-paired reads against the KEGG database56,66 using DIAMOND51 in the sensitive mode retaining only the top hit (E < 10−5) for each query. Putative anaerobic hydrocarbon activation genes were identified by a translated homology search in USEARCH against the AnHyDeg database62. Acceptance criteria were 60% identity over alignments with at least 35 amino acids. Functional gene abundances were calculated by dividing the number of hits to specific reference gene by the number of hits to the single-copy marker gene, rpoB, to normalize for differential library population size67. A modified approach was used to estimate the total hydrocarbon-activating proportion of each community from that previously used68 by a summation for selected aerobic and anaerobic specific activation genes of the following: (hits to functional gene /length of functional gene)/hits to rpoB. Aerobic activation genes included: alkB, etbAa, toxA, todC1, dmpB, nahAc, and 1-methylnapthalene hydroxylase-like genes. Anaerobic activation genes were ebdA, assA, bsdA, nmsA.

16S rRNA amplicon libraries for high-throughput sequencing
An aliquot of the DNA extracted from filters was used to generate 16S amplicon libraries with distinctive barcodes as previously described69. Briefly, the M13-519F/785R primer set70 was used to amplify the V3-V5 region of the 16S rRNA gene, and a 12 base pair sequence conjugated to M1371 was used to barcode each sample (Supplementary Table 4). Amplicon libraries were sequenced as described above and analyzed as previously described69. Paired-end reads were trimmed and stitched together, and default parameters in QiIME were used to demultiplex, cluster, and remove chimeras.

Metabolite analyses
Mass spectral analysis of diagnostic metabolites associated with oxygen-dependent and independent hydrocarbon activation of diesel fuel components was performed on acidified ballast tank samples. Methods used for the targeted (GC-MS) and global survey of organic intermediates were as reported previously72.

Thermodynamic equilibrium solubility calculations
The equilibrium solubility of the carbonate and sulfide mineral compounds of manganese, copper and nickel in seawater were calculated using MINEQL 5.0, with the following chemical parameters: total carbonates (2mM), ion strength (0.5 M), and a temperature of 25 °C. Since the San Diego Harbor seawater and ballast water ranged in pH from 7.65 to 8.2873 and between 7.71 and 8.3 (data not shown), respectively, a constant pH of 8.2 was used in the MINEQL 5.0 calculations. The small pH differences from the constant value resulted in a <1% change in the metal sulfide equilibrium solubility values. A closed system was modeled for each metal carbonate and sulfide compound contained in the MINEQL 5.0 thermodynamic database, and CO2 (g) and H2S (g) were removed as fixed entities.

DATA AVAILABILITY
Metagenome and 16S amplicon sequence files are available at the NCBI Sequence Read Archive under accession numbers SRR14067720 for Ship #1, SRR14067719 for Ship #2, SRR14067718 for Ship #3, SRR14067717 for Harbor SD, and SRR14121797 for the 16S amplicon library.

Received: 1 June 2021; Accepted: 3 November 2021; Published online: 01 December 2021

REFERENCES
1. Gillings, M. R. & Paulsen, I. T. Microbiology of the anthropocene. Anthropocene 5, 1–8 (2014).
2. Crutzen, P. J. Geology of mankind. Nature 415, 23 (2002).
3. Economist P. World Energy 2016-2050: Annual Report. Peak Oil Barrel [Internet]. Available at: http://peakoilbarrel.com/world-energy-2016-2050-annual-report/ (2016).
4. https://www.worldsteel.org/en/dam/jcr:fc7982217c–fdfe-4fcd-8baa-795ed8075f13/World%2520Steel%2520in%2520Figures%25202020if.pdf (2020).
5. Smil, V. Still the Iron Age: Iron and Steel in the Modern World (Butterworth-Heinemann, 2016).
6. Hazlett, R. N. Examination of Corrosion Deposits from U.S. WO JIMA Aviation Gasoline Storage Tanks Memorandum Report 1779 (U.S. Naval Research Laboratory, Washington, DC, 1967).
7. Hazlett, R. N. Control of Sulfate-reducing Bacteria in Navy Fuel Systems Memorandum Report 1737 (U.S. Naval Research Laboratory, Washington, DC, 1966).
8. Klemme, D. E. & Leonard J. M. Inhibitors for Marine Sulfate-reducing Bacteria in Shipboard Fuel Storage Tanks Memorandum Report 2324 (U.S. Naval Research Laboratory, Washington, DC, 1971).
9. Klemme, D. E. & Neihof R. A. Control of Marine Sulfate-reducing Bacteria in Water-displaced Shipboard Fuel Storage Tanks Memorandum Report 2069 (U.S. Naval Research Laboratory, Washington, DC, 1969).
10. Cleland, J. H. Corrosion risks in ships’ ballast tanks and the IMO pathogen guidelines. Eng. Fail. Anal. 2, 79–94 (1995).
11. Huang, R. T., McFarland, B. L. & Hodgman R. Z. Microbial influenced corrosion in cargo oil tanks of crude oil tankers. In: Paper NACE-9735 Presented at the Corrosion97, New Orleans, Louisiana, March (NACE International, 1997).
12. Liang, R. & Sulfita, J. M. in Hydrocarbon and Lipid Microbiology Protocols. Springer Protocols Handbooks. (eds McGinity, T., Timmis, K. & Nogales, B. J. 211–226 (Springer, Berlin, Heidelberg, 2015). https://doi.org/10.1007/8623_2015_76.
13. Lyles, C. N. et al. Impact of organosulfur content on diesel fuel stability and implications for carbon steel corrosion. Environ. Sci. Technol. 7, 6052–6062 (2013).
14. Aktas, D. F. et al. Anaerobic metabolism of biodiesel and its impact on metal corrosion. Energy Fuels 24, 2924–2928 (2010).
15. Lee, J. S. et al. Sulphide production and corrosion in seawaters during exposure to FAME diesel. Biofuels 28, 465–478 (2012).
16. Liang, R. et al. Anaerobic biodegradation of alternative fuels and associated biocorrosion of carbon steel in marine environments. Environ. Sci. Technol. 50, 4844–4853 (2016).
17. Aktas, D. F. et al. Anaerobic hydrocarbon biodegradation and biocorrosion of carbon steel in marine environments: The impact of different ultra low sulfur diesels and bioaugmentation. Int. Biodeterior. Biodegrad. 118, 45–56 (2017).
18. Liang, R., Harvey, B. G., Quintana, R. L. & Sulfita, J. M. Assessing the biological stability of a terpene-based advanced biofuel and its relationship to the corrosion of carbon steel. Energy Fuels. 29, 5164–5170 (2015).
19. King, G. M., Smith, C. B., Tolar, B. & Hollabaugh, J. T. Analysis of composition and structure of coastal to mesopelagic bacterioplankton communities in the northern Gulf of Mexico. Front. Microbiol. 3, 1–14 (2013). article 438.
20. Geiselbrecht, A. D., Hedlund, B. P., Tichy, M. A. & Staley, J. T. Isolation of marine polyacrylamidic aromatic hydrocarbon (PAH)-degrading Cycloclастicos strains from the Gulf of Mexico and comparison of their PAH degradation ability with that of pure sound Cycloclasticos strains. Appl. Environ. Microbiol. 64, 4703–4710 (1998).
21. Kasai, Y., Kishira, H. & Harayama, S. Bacteria belonging to the genus Cycloclasticus play a primary role in the degradation of aromatic hydrocarbons released in a marine environment. Appl. Environ. Microbiol. 68, 5625–5633 (2002).
22. Dubinsky, E. A. et al. Succession of hydrocarbon-degrading bacteria in the aftermath of the Deepwater Horizon oil spill in the Gulf of Mexico. Environ. Sci. Technol. 47, 10860–10867 (2013).
23. King, G. M., Kostka, J. E., Hazen, C. & Sobecky, P. A. Microbial responses to the Deepwater Horizon oil spill: From coastal wetlands to the deep sea. Ann. Rev. Mar. Sci. 7, 377–401 (2015).
24. Striebel, R. C. et al. Characterization of the F-76 diesel and Jet-A aviation fuel hydrocarbon degradation profiles of Phutomoean aerogenis and Marinobacter hydrocarbonoclasticus. Int. Biodeterior. Biodegrad. 93, 33–43 (2014).
25. Berendis, S. et al. First evidence of aerobic biodegradation of BTEX compounds by pure cultures of Marinobacter. Appl. Biochem. Biotechnol. 160, 1992–1999 (2010).
26. Yakimov, M. M. et al. Alcanivorax borkumensis gen. nov., sp. nov, a new hydrocarbon-degrading and surfactant-producing marine bacterium. Int. J. Syst. Evol. Microbiol. 48, 339–348 (1998).
27. Harayama, S., Kasai, Y. & Hara, A. Microbial communities in oil-contaminated seawater. Curr. Opin. Biotechnol. 15, 205–214 (2004).
ADDITIONAL INFORMATION

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41529-021-00207-z.

Correspondence and requests for materials should be addressed to Joseph M. Sufita.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2021