Data Article

16S rRNA gene sequencing data of the human skin microbiome before and after swimming in the ocean

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Abstract

These data represent the abundance, diversity and predicted function gene profiles of the microbial communities present on human skin before and after swimming in the ocean. The skin microbiome has been shown to provide protection against infection from pathogenic bacteria. It is well-known that exposure to ocean water can cause skin infection, but little is known about how exposure can alter the bacterial communities on the skin. Skin microbiome samples were collected from human participants before and after swimming in the ocean. These data were used to analyze the changes in abundance and diversity of microbial communities on the skin and the changes in the functional profiles of the bacteria, specifically focusing on genes involved in antibiotic resistance and bacterial virulence.

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Specifications Table

| Subject | Microbiology: microbiome |
|---------|--------------------------|
| Specific subject area | Investigation of the changes in human skin microbial communities after ocean water exposure |
| Type of data | Table |
| How data were acquired | 16S rRNA next-generation sequencing of extracted DNA from human skin microbiome swab samples. |
| Instrument | Ion Torrent PGM at MRDNA/Molecular Research LP [1] |
| Data format | Raw |
| Parameters for data collection | Skin swab samples were collected from human volunteers at the beach. Participants had to consent and meet inclusion criteria (detailed in the experimental design section) to enroll. |
| Description of data collection | Skin microbiome samples were collected from human participants before swimming in the ocean, after swimming, and at 6h and 24 h post-swim. |
| Data source location | Institution: University of California, Irvine |
| | City/Town/Region: Irvine, CA |
| | Country: United States |
| | Samples were collected at Huntington Beach, CA |
| Data accessibility | Repository name: NCBI SRA |
| | BioProject ID: PRJNA735376 |
| | Direct URL to data: [http://www.ncbi.nlm.nih.gov/bioproject/735376](http://www.ncbi.nlm.nih.gov/bioproject/735376) |
| | Repository name: Mendeley Data |
| | Direct URL to data: [https://data.mendeley.com/datasets/th7bfgfc6m/1](https://data.mendeley.com/datasets/th7bfgfc6m/1) and [https://data.mendeley.com/datasets/hjnzm949y/1](https://data.mendeley.com/datasets/hjnzm949y/1) [5] |
| Related research article | M.C. Nielsen, N. Wang, S.C. Jiang, Acquisition of antibiotic resistance genes on human skin after swimming in the ocean, Environ. Res. 197 (2021) 110978 [6]. |

Value of the Data

- These data are valuable to marine, environmental and clinical microbiome researchers interested in understanding the role of the environment in skin microbial community function and alteration.
- These data can be used by public health officials interested in investigating mitigation and surveillance strategies for antibiotic resistant infections.
- The microbiome datasets provided can be used to further examine how changes in the microbiome, including additional function gene profiles, are affected by exposure to ocean water.

1. Data Description

The raw datasets contain 16S rRNA gene sequence data of swab samples taken from human skin before and after swimming in the ocean. Sequences are also provided for samples taken 6h and 24h post-swim from most of the participants. These data were used to investigate the changes in the skin microbiome after swimming in the ocean. QIIME was used to analyze changes in microbial communities and PICRUSt was used to analyze the predicted function gene profiles.

Table 1 lists the antibiotic resistance genes (ARGs) and their corresponding Kyoto Encyclopedia of Genes and Genomes (KEGG) Orthologs (KOs) detected in the samples. Table 2 lists the KOs associated with virulence factor genes (VFGs) detected in the samples and their corresponding KEGG descriptions.
Table 1
Antibiotics and their corresponding ARGs and KOs detected in the samples.

| Antibiotic                  | Gene (KO)                                                                 |
|-----------------------------|---------------------------------------------------------------------------|
| Vancomycin                  | vanX (K08641); vanY (K07260); vraR (K07694); vraS (K07681)               |
| Tetracycline                | tetA/tetG/H/J (K08151); tetK (K08168)                                     |
| fluoroquinolone, quinolone, | adeA/cmeA (K03585); catB3 (K00638); basR (K07771); qepA (K08167)         |
| florfenicol, chloramphenicol, |                                                                          |
| and amphenicol (PCA)        |                                                                          |
| beta-lactams                | acrA (K03585); ampC (K01467); ampG (K08218); blal (K02171); blar1 (     |
|                             | (K02172); cfxA (K01624); ftsl (K03587); mecA (K02545); mecR1 (K02547); |
|                             | metallo-beta-lactamase family protein (K07576); mrcA (K05366); mrdA (     |
|                             | (K05515); nagZ (K01207); ompU (K08720); ompC (K09475); ompF (          |
|                             | (K09476); pbpA (K12552); pbp1b (K03693); pbp2A (K12555); pbp2B (        |
|                             | (K00687); pbp2X (K12556); pbp3 (K12553); penA (K03587); tolC (K12340)    |
| multidrug                   | emrE/qac/mmr/smr(K03297); MATE family (K03327); emrB(K03446);           |
|                             | emrA(K03543); marC(K05595); mdtB(K07788); mdtC(K07789); mdtD(K07799);   |
|                             | lmrP(K08152); blt(K08153); mdtA/cmr(K08160); mdtG(K08161); mdtH(K08162);|
|                             | mdtI(K08163); yebQ(K08169); norB(C)(K08170); yitG/yfmD/yfmO(K08221);   |
|                             | oprJ(K08721); ebrB(A)(K11814); ebrF(B)(K11815)                           |
| aminoglycosides             | aacC1(K03395); aacC2(K00662); aacC4(K00663); aadA1(K00984);             |
|                             | aade(K05593); yblC(K08164)                                               |
| macrolide, lincosamide, and | ermA(A)(K00561); ereA_B(K06880); mph(K06979); mef(K08217);              |
| streptogramin B (MLSb)      | macA(K13888)                                                              |

2. Experimental Design, Materials and Methods

2.1. Sample collection

Data collection methods were approved by the University of California, Irvine Institutional Review Board (IRB #2017-3751). Verbal consent was collected from participants before registration. Sample collection events occurred in April and September 2018 in Huntington Beach, CA. In brief, participants were given a detailed sample collection description if they met the participant criteria. Skin microbiome samples were obtained from 9 participants in April and 12 participants in September and included males and females with age ranges from 24–39, with no sunscreen application, infrequent exposure to the ocean and beach (once per month or less), no shower/bath in the past 12 hours, no antibiotic usage in the past 6 months, and no active infections. Rayon-tipped swabs moistened in sterile saline were used to swab an 8cm × 8cm section of skin on the back of the participants’ calves before the individuals swam in the ocean. Participants were then instructed to swim or wade in the ocean for 10 minutes and air dry. At that time, a second set of samples were taken. The before and after samples were collected from the same calf but on different sections of the skin to ensure the sample collection taken before swimming did not remove bacteria from the section of skin swabbed after swimming. The participants were then instructed to leave the beach and not to shower or wash the leg area until after the 6 h and 24 h post-swim samples were collected. The 6 h samples were collected from the right side of the left calf and the 24 h samples were collected from the left side of the left calf. Samples were kept on ice after collection and all samples were processed within 24 h of collection. Ocean water samples (75mL each) were collected from each swim site at the time of experimental sample collection and were analyzed in the same manner as the experimental samples. [7].
Table 2
Descriptions of the KOs associated with VFGs detected in the samples.

| #OTU ID | KEGG Description |
|---------|------------------|
| K12984 | "(heptosyl)LPS beta-1,4-glucosyltransferase [Enzyme commission number (EC):2.4.1.-]" |
| K00655 | "1-acyl-sn-glycerol-3-phosphate acyltransferase [EC:2.3.1.51]" |
| K01771 | "1-phosphatidylinositol phosphodiesterase [EC:4.6.1.13]" |
| K00216 | "2,3-dihydro-2,3-dihydroxybenzoate dehydrogenase [EC:1.3.1.28]" |
| K02510 | "2,4-dihydroxyhept-2-ene-1,7-dioic acid aldolase [EC:4.1.2.-]" |
| K01627 | "2-dehydro-3-deoxyphosphooctonate aldolase (KDO 8-P synthase) [EC:2.5.1.55]" |
| K01626 | "3-deoxy-7-phosphoheptulonate synthase [EC:4.6.1.13]" |
| K11211 | "3-deoxy-D-manno-octulosonic acid kinase [EC:2.7.1.-]" |
| K02527 | "3-deoxy-D-manno-octulosonic-acid transferase [EC:2.-.-.-]" |
| K00979 | "3-deoxy-D-manno-octulosonate cytidylyltransferase (CMP-KDO synthetase) [EC:2.7.7.38]" |
| K00074 | "3-hydroxybutyryl-CoA dehydrogenase [EC:1.1.1.157]" |
| K01704 | "3-isopropylmalate/(R)-2-methylmalate dehydratase small subunit [EC:4.2.1.33 4.2.1.35]" |
| K00659 | "3-oxoacyl-[acyl-carrier protein] reductase [EC:1.1.1.100]" |
| K02372 | "3R-hydroxymyristoyl ACP dehydrase [EC:4.2.1.-]" |
| K00979 | "3-deoxy-D-manno-octulosonate cytidylyltransferase (CMP-KDO synthetase) [EC:2.7.7.38]" |
| K00652 | "8-amino-7-oxononanoate synthase [EC:2.3.1.47]" |
| K01990 | "ABC-2 type transport system ATP-binding protein" |
| K01992 | "ABC-2 type transport system permease protein" |
| K10952 | "accessory chola enterotoxin" |
| K10936 | "accessory colonization factor AcfB" |
| K10939 | "accessory colonization factor AcfC" |
| K10939 | "accessory colonization factor AcfD" |
| K07813 | "accessory gene regulator B" |
| K04072 | "acetaldehyde dehydrogenase / alcohol dehydrogenase [EC:1.2.1.10 1.1.1.1]" |
| K03896 | "acetyl CoA:N6-hydroxylysine acetyl transferase [EC:2.3.1.102]" |
| K00821 | "acetyl-CoA:N6-hydroxylysine acetyl transferase [EC:2.3.1.102]" |
| K01078 | "acid phosphatase [EC:3.1.3.2]" |
| K03191 | "acid-activated urea channel" |
| K02078 | "acyl carrier protein" |
| K13061 | "acyl homoserine lactone synthase [EC:2.3.1.184]" |
| K13060 | "acyl homoserine lactone synthase [EC:2.3.1.184]" |
| K00249 | "acyl-Coa dehydrogenase [EC:13.99.3]" |
| K07116 | "acyl-homoserine-lactone acylase [EC:3.5.1.97], """ |
| K00860 | "adenyllysulfate kinase [EC:2.7.1.85]" |
| K13735 | "adhesin/invasin" |
| K03274 | "ADP-1-glycerol-D-manno-heptose 6-epimerase [EC:5.1.3.20]" |
| K03894 | "aerobactin synthetase subunit alpha [EC:6.3.2.27]" |
| K03895 | "aerobactin synthetase subunit beta [EC:6.3.2.27]" |
| K12678 | "AIDA-I adhesin-like protein" |
| K00697 | "alpha,phi-trehalose-phosphate synthase (UDP-forming) [EC:2.4.1.15]" |
| K12994 | "alpha-1,3-rhamnosyltransferase [EC:2.4.1.-]" |
| K00766 | "anthranilate phosphoribosyltransferase [EC:2.4.2.18]" |
| K01657 | "anthranilate synthase component 1 [EC:4.1.3.27]" |
| K12687 | "antigen 43" |
| K06041 | "arabinose-5-phosphate isomerase [EC:5.3.1.31]" |
| K05372 | "AraC family transcriptional regulator" |
| K06599 | "AraC family transcriptional regulator, chemosensory pilus system protein ChpD" |
| K10923 | "AraC family transcriptional regulator, TCP pilus virulence regulatory protein" |
| K12243 | "AraC family transcriptional regulator, transcriptional activator of the genes for pyochelin and ferrirpyochelin receptors" |
| K01953 | "asparagine synthase (glutamine-hydrolysing) [EC:6.3.5.4]" |
| K01779 | "aspartate 1-decarboxylase [EC:4.1.1.11]" |
| K01779 | "aspartate racemase [EC:5.1.1.13]" |

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| #OTU ID  | KEGG Description                                                                 |
|----------|----------------------------------------------------------------------------------|
| K03224   | \["ATP synthase in type III secretion protein SctN \{EC:3.6.3.14\}\]           |
| K11004   | \["ATP-binding cassette, subfamily B, bacterial HlyB/CyaB\]\]                   |
| K14698   | \["ATP-binding cassette, subfamily B, bacterial IrrA \{EC:3.6.3.\}\]          |
| K14699   | \["ATP-binding cassette, subfamily B, bacterial IrrB \{EC:3.6.3.\}\]          |
| K11085   | \["ATP-binding cassette, subfamily B, bacterial MsbA \{EC:3.6.3.\}\]          |
| K12530   | \["ATP-binding cassette, subfamily B, bacterial RtxB\]\]                       |
| K12531   | \["ATP-binding cassette, subfamily B, bacterial RtxE\]\]                       |
| K06147   | \["ATP-binding cassette, subfamily C, bacterial\]\]                            |
| K12536   | \["ATP-binding cassette, subfamily C, bacterial\]\]                            |
| K03696   | \["ATP-dependent Clp protease ATP-binding subunit ClpC\]\]                     |
| K03697   | \["ATP-dependent Clp protease ATP-binding subunit ClpE\]\]                     |
| K01358   | \["aureolysin \{EC:3.4.24.29\}\]                                              |
| K12688   | \["autotransporter serine protease \{EC:3.4.21.-\}\]                          |
| K00785   | \["beta-galactosamidase-alpha-2,3-sialyltransferase \{EC:2.4.99.-\}\]        |
| K11609   | \["beta-ketoacyl ACP synthase \{EC:2.3.1.-\}\]                                |
| K13714   | \["bifunctional autolysin \{EC:3.5.1.28 3.2.1.96\}\]                        |
| K11936   | \["biofilm PGA synthesis N-glycosyltransferase PgaC \{EC:2.4.-.-\}\]        |
| K11937   | \["biofilm PGA synthesis protein PgaD\]\]                                      |
| K03561   | \["biopolymer transport protein ExbB\]\]                                       |
| K03559   | \["biopolymer transport protein ExbD\]\]                                       |
| K06011   | \["bontoxylisin \{EC:3.4.24.69\}\]                                             |
| K08652   | \["CSa peptidase \{EC:3.4.21.110\}\]                                            |
| K12107   | \["cag pathogenicity island protein 22\]\]                                     |
| K10915   | \["CAI-1 autoinducer synthase \{EC:2.3.-.-\}\]                                |
| K11045   | \["cAMP factor\]\]                                                            |
| K07265   | \["capsular polysaccharide export protein\]\]                                  |
| K07266   | \["capsular polysaccharide export protein\]\]                                  |
| K09689   | \["capsular polysaccharide transport system ATP-binding protein\]\]            |
| K10107   | \["capsular polysaccharide transport system permease protein\]\]               |
| K09688   | \["capsular polysaccharide transport system permease protein\]\]               |
| K03563   | \["carbon storage regulator\]\]                                                |
| K01673   | \["carbonic anhydrase \{EC:4.2.1.1\}\]                                         |
| K06131   | \["cardiolipin synthase \{EC:2.7.8.-\}\]                                      |
| K03781   | \["catalase \{EC:1.11.1.6\}\]                                                 |
| K03782   | \["catalase/peroxidase \{EC:1.11.1.6 1.11.1.7\}\]                             |
| K12956   | \["cation-transporting ATPase V \{EC:3.6.3.-\}\]                              |
| K12950   | \["cation-transporting P-type ATPase C \{EC:3.6.3.-\}\]                       |
| K00523   | \["CDP-4-dehydro-6-deoxyglucose reductase \{EC:1.17.1.1\}\]                   |
| K12452   | \["CDP-6-deoxy-D-xylo-4-hexulose-3-dehydrase\]\]                               |
| K01709   | \["CDP-glucose-4,6-dehydratase \{EC:4.2.1.45\}\]                              |
| K09809   | \["CDP-glycerol glycerophosphotransferase \{EC:2.7.8.12\}\]                   |
| K04095   | \["cell filamentation protein\]\]                                              |
| K05789   | \["chain length determinant protein (polysaccharide antigen chain regulator)\]\|
| K12519   | \["chaperone protein PapD\]\]                                                  |
| K04077   | \["chaperonin GroEL\]\]                                                       |
| K06596   | \["chemosensory pili system protein ChpA (sensor histidine kinase/response regulator)\]\|
| K06597   | \["chemosensory pili system protein ChpB (putative protein-glutamate methyltransferase)\]\|
| K06598   | \["chemosensory pili system protein ChpC\]\]                                   |
| K06600   | \["chemosensory pili system protein ChpE\]\]                                   |
| K03411   | \["chemotaxis protein CheD \{EC:3.5.1.44\}\]                                  |
| K03414   | \["chemotaxis protein CheZ\]\]                                                 |
| K00575   | \["chemotaxis protein methyltransferase CheR \{EC:2.1.1.80\}\]               |
| K02556   | \["chemotaxis protein MotA\]\]                                                 |
| K02557   | \["chemotaxis protein MotB\]\]                                                 |
| K03933   | \["chitin-binding protein\]\]                                                   |
| K10928   | \["cholera enterotoxin subunit A \{EC:2.4.2.36\}\]                           |
| K10929   | \["cholera enterotoxin subunit B\]\]                                           |
| #OTU ID | KEGG Description |
|---------|------------------|
| K01442  | ["choloylglycine hydrolase [EC:3.5.1.24]""] |
| K03496  | ["chromosome partitioning protein""] |
| K03497  | ["chromosome partitioning protein, ParB family""] |
| K01644  | ["citrate lyase subunit beta [EC:4.1.3.6]", "citrate lyase subunit beta / citrly-CoA lyase [EC:4.1.3.6 4.1.3.34]""] |
| K14201  | ["clumping factor A""] |
| K14192  | ["clumping factor B""] |
| K08605  | ["coccusin [EC:3.4.24.30]""] |
| K02237  | ["competence protein ComEA""] |
| K14446  | ["crotonyl-CoA carboxylase/reductase""] |
| K10914  | ["CRP/FNR family transcriptional regulator, cyclic AMP receptor protein""] |
| K07796  | ["Cu(I)/Ag(I) efflux system outer membrane protein CusC""] |
| K04565  | ["Cu/Zn superoxide dismutase [EC:1.15.1.1]""] |
| K04337  | ["cycle production assembly/transport component CsgE""] |
| K04338  | ["cycle production assembly/transport component CsgF""] |
| K06214  | ["cycle production assembly/transport component CsgG""] |
| K04336  | ["cycle production protein""] |
| K02500  | ["cyclase HisF [EC:4.1.3.-]", "cyclase [EC:4.1.3.-]""] |
| K13688  | ["cyclic beta-1,2-glucan synthetase [EC:2.4.1.-]""] |
| K00574  | ["cyclopropane-fatty-acyl-phospholipid synthase [EC:2.1.1.79]""] |
| K11049  | ["C01G protein""] |
| K01697  | ["cystathionine beta-synthase [EC:4.2.1.22]""] |
| K01738  | ["cysteine synthase A [EC:2.5.1.47]""] |
| K02424  | ["cysteine transport system substrate-binding protein""] |
| K02197  | ["cytochrome c-type biogenesis protein CcmE""] |
| K02198  | ["cytochrome c-type biogenesis protein CcmF""] |
| K11013  | ["cytolethal distending toxin subunit A""] |
| K11014  | ["cytolethal distending toxin subunit B""] |
| K11015  | ["cytolethal distending toxin subunit C""] |
| K03367  | ["D-alanine-poly(phosphoribitol) ligase [EC:6.1.11.13], "D-alanine–poly(phosphoribitol) ligase subunit 1 [EC:6.1.11.13]""] |
| K07262  | ["D-alyglycine-d-alanine endopeptidase (penicillin-binding protein 7) [EC:3.4.99.-]", "D-alyglycine-d-alanine endopeptidase (penicillin-binding protein 7) [EC:3.4.21.-]""] |
| K03272  | ["D-beta-H-heptose 7-phosphate kinase / D-beta-H-heptose 1-phosphate adenosyltransferase [EC:2.7.1.- 2.7.7.1]""] |
| K01494  | ["dCTP deaminase [EC:3.5.4.13]""] |
| K12202  | ["defect in organelle trafficking protein DotA""] |
| K12203  | ["defect in organelle trafficking protein DotB""] |
| K12204  | ["defect in organelle trafficking protein DotC""] |
| K12205  | ["defect in organelle trafficking protein DotD""] |
| K11039  | ["delta-hemolysin""] |
| K00689  | ["dextranucrase [EC:2.4.1.5]""] |
| K03273  | ["D-glycerol-d-manno-heptose 1,7-bisphosphate phosphatase [EC:3.1.3.-]""] |
| K00836  | ["diaminobutyrate-2-oxoglutarate transaminase [EC:2.6.1.76]""] |
| K01586  | ["diaminopimelate decarboxylase [EC:4.1.1.20]""] |
| K12239  | ["dihydroaeruginoic acid synthetase""] |
| K02073  | ["D-methionine transport system substrate-binding protein""] |
| K01972  | ["DNA ligase (NAD+) [EC:6.5.1.12]""] |
| K04483  | ["DNA repair protein RadA""] |
| K03631  | ["DNA repair protein RecN (Recombination protein N)""] |
| K03466  | ["DNA segregation ATPase PtsK/SpolIIe, S-DNA-T family""] |
| K03746  | ["DNA-binding protein H-NS""] |
| K03530  | ["DNA-binding protein HU-beta""] |
| K01790  | ["dTDP-4-dehydrohamnose 3,5-epimerase [EC:5.1.3.13]""] |
| K00067  | ["dTDP-4-dehydrohamnose reductase [EC:1.1.13.3]""] |
| K01710  | ["dTDP-glucose 4,6-dehydratase [EC:4.2.1.46]""] |

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Table 2 (continued)

| #OTU ID | KEGG Description |
|---------|------------------|
| K03709  | "DtxR family transcriptional regulator, Mn-dependent transcriptional regulator" |
| K13461  | "effector protein HopM1" |
| K02358  | "elongation factor EF-Tu [EC:3.6.5.3], "elongation factor Tu" |
| K01689  | "enolase [EC:4.2.1.11]" |
| K00209  | "enoyl-[acyl-carrier-protein] reductase (NADPH2, B-specific) [EC:1.3.1.10]" |
| K02363  | "enterobactin 2,3-dihydroxybenzoate-AMP ligase / S-dihydroxybenzoyltransferase [EC:2.7.7.58 2.3.1.-]" |
| K01252  | "enterobactin isochorismatase [EC:3.3.2.1]" |
| K02356  | "enterobactin synthetase component D [EC:2.7.8.-]" |
| K02364  | "enterobactin synthetase component F [EC:2.7.7.-]" |
| K01066  | "esterase / lipase [EC:3.1.1.-]" |
| K11041  | "exfoliative toxin A/B" |
| K01142  | "exodeoxyribonuclease III [EC:3.1.11.2]" |
| K12423  | "fatty acid CoA ligase FadD21" |
| K12427  | "fatty acid CoA ligase FadD28" |
| K12421  | "fatty acid CoA ligase FadD9" |
| K00666  | "fatty-acyl-CoA synthase [EC:6.2.1.-]" |
| K13255  | "ferric iron reductase protein FhuF" |
| K01772  | "ferrochelatase [EC:4.99.1.1]" |
| K04758  | "ferrous iron transport protein A" |
| K04759  | "ferrous iron transport protein B" |
| K13782  | "fibrinogen-binding protein" |
| K13734  | "fibronecin-binding protein 1" |
| K13732  | "fibronecin-binding protein A" |
| K13733  | "fibronecin-binding protein B" |
| K07346  | "fimbrial chaperone protein" |
| K08087  | "fimbrial protein FimW" |
| K08088  | "fimbrial protein FimY" |
| K07351  | "fimbrial protein" |
| K03773  | "FKBP-type peptidyl-prolyl cis-trans isomerase FklB [EC:5.2.1.8]" |
| K02386  | "flagella basal body P-ring formation protein FlgA" |
| K02399  | "flagella synthesis protein FlgN" |
| K13626  | "flagellar assembly factor FliW" |
| K02411  | "flagellar assembly protein FlhB" |
| K02389  | "flagellar basal-body rod modification protein FlgD" |
| K02387  | "flagellar basal-body rod protein FlgB" |
| K02388  | "flagellar basal-body rod protein FlgC" |
| K02391  | "flagellar basal-body rod protein FlgF" |
| K02392  | "flagellar basal-body rod protein FlgG" |
| K02400  | "flagellar biosynthesis protein FlhA" |
| K02404  | "flagellar biosynthesis protein FlhF" |
| K04562  | "flagellar biosynthesis protein FlhG" |
| K04061  | "flagellar biosynthesis protein" |
| K02401  | "flagellar biosynthetic protein FlhB" |
| K02419  | "flagellar biosynthetic protein FlhP" |
| K02420  | "flagellar biosynthetic protein FlIQ" |
| K02421  | "flagellar biosynthetic protein FlIR" |
| K13920  | "flagellar biosynthetic protein FlIR/FlhB" |
| K02413  | "flagellar FliJ" protein |
| K02415  | "flagellar FliJ protein" |
| K02390  | "flagellar hook protein FlgE" |
| K02396  | "flagellar hook-associated protein 1 FlgK" |
| K02407  | "flagellar hook-associated protein 2" |
| K02397  | "flagellar hook-associated protein 3 Flgl" |
| K02408  | "flagellar hook-basal body complex protein FliE" |
| K02414  | "flagellar hook-length control protein FliK" |
| K02393  | "flagellar L-ring protein precursor FliH" |
| K02410  | "flagellar motor switch protein FliC" |
| K02416  | "flagellar motor switch protein FliM" |

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| #OTU ID | KEGG Description |
|---------|------------------|
| K02417  | ["flagellar motor switch protein FliN/FliY"] |
| K02409  | ["flagellar M-ring protein FliF"] |
| K02394  | ["flagellar P-ring protein precursor FlgI"] |
| K06602  | ["flagellar protein FlaF"] |
| K06603  | ["flagellar protein FlaG"] |
| K02385  | ["flagellar protein FliD"] |
| K06601  | ["flagellar protein FliT"] |
| K02395  | ["flagellar protein FlgI"] |
| K03516  | ["flagellar protein FlhE"] |
| K02418  | ["flagellar protein FliO/FliZ"] |
| K02422  | ["flagellar protein Fls"] |
| K02423  | ["flagellar protein Flt"] |
| K06604  | ["flagellar rod protein Flal"] |
| K02402  | ["flagellar transcriptional activator FlhC"] |
| K02403  | ["flagellar transcriptional activator FlhD"] |
| K02406  | ["flagellin"] |
| K02412  | ["flagellum-specific ATP synthase [EC:3.6.3.14]"] |
| K02425  | ["Fliz protein"] |
| K07533  | ["foldase protein PrsA [EC:5.2.1.8]"] |
| K01622  | ["fructose 1,6-bisphosphate aldolase/phosphatase [EC:4.1.2.13 3.1.3.11]"] |
| K13007  | ["Fuc2NAc and GlcNAc transferase [EC:2.4.1.-]"] |
| K03711  | ["Fur family transcriptional regulator, ferric uptake regulator"] |
| K00681  | ["gamma-glutamyltranspeptidase [EC:2.3.2.2]"] |
| K02377  | ["GDP-L-fucose synthase [EC:1.1.1.271]"] |
| K01711  | ["GDP-mannose 4,6-dehydratase [EC:4.2.1.47]"] |
| K00066  | ["GDP-mannose 6-dehydrogenase [EC:1.1.1.132]"] |
| K03285  | ["general bacterial porin, GBP family"] |
| K02450  | ["general secretion pathway protein A"] |
| K02451  | ["general secretion pathway protein B"] |
| K02452  | ["general secretion pathway protein C"] |
| K02453  | ["general secretion pathway protein D"] |
| K02454  | ["general secretion pathway protein E"] |
| K02455  | ["general secretion pathway protein F"] |
| K02456  | ["general secretion pathway protein G"] |
| K02457  | ["general secretion pathway protein H"] |
| K02458  | ["general secretion pathway protein I"] |
| K02459  | ["general secretion pathway protein J"] |
| K02460  | ["general secretion pathway protein K"] |
| K02461  | ["general secretion pathway protein L"] |
| K02462  | ["general secretion pathway protein M"] |
| K02463  | ["general secretion pathway protein N"] |
| K02464  | ["general secretion pathway protein O [EC:3.4.23.43 2.1.1.-]"] |
| K02465  | ["general secretion pathway protein S"] |
| K01178  | ["glucoamylase [EC:3.2.1.3]"] |
| K00973  | ["glucose-1-phosphate thymidylyltransferase [EC:2.7.7.24]"] |
| K01810  | ["glucose-6-phosphate isomerase [EC:5.3.1.9]"] |
| K12998  | ["glucosyltransferase [EC:2.4.1.-]"] |
| K01845  | ["glutamyl endopeptidase [EC:3.4.21.19]"] |
| K02492  | ["glutamyl-tRNA reductase [EC:1.2.1.70]"] |
| K00134  | ["glyceraldehyde 3-phosphate dehydrogenase [EC:1.2.1.12]"] |
| K00613  | ["glycine amidinotransferase [EC:2.4.2.-]"] |
| K01915  | ["glutamine synthetase [EC:6.3.1.2]"] |
| K01318  | ["glutamate-1-semialdehyde 2,1-aminomutase [EC:5.4.3.8]"] |
| K02501  | ["glutathione peroxidase [EC:1.1.1.9]"] |
| K01915  | ["glutathione peroxidase [EC:1.1.1.9]"] |
| K02501  | ["glutamine amidotransferase [EC:2.4.2.-]"] |
| K01915  | ["glutamine synthetase [EC:6.3.1.2]"] |
| K01845  | ["glutamate-1-semialdehyde 2,1-aminomutase [EC:5.4.3.8]"] |
| K00134  | ["glyceraldehyde 3-phosphate dehydrogenase [EC:1.2.1.12]"] |
| K00613  | ["glycine amidinotransferase [EC:2.4.2.-]"] |
| K07270  | ["glycosyltransferase, family 25"] |
| K13002  | ["glycosyltransferase [EC:2.4.1.-]"] |
| K01495  | ["GTP cyclohydrolase I [EC:3.5.4.16]"] |
| K00951  | ["GTP pyrophosphokinase [EC:2.7.6.5]"] |
| #OTU ID  | KEGG Description                                      |
|----------|-------------------------------------------------------|
| K12545   | "heme acquisition protein HasA"                      |
| K02193   | "heme exporter protein A [EC:3.6.3.41]"               |
| K02194   | "heme exporter protein B"                            |
| K02196   | "heme exporter protein D"                            |
| K07215   | "heme oxygenase"                                     |
| K11005   | "hemolysin A"                                        |
| K07326   | "hemolysin activation/secretion protein"             |
| K11017   | "hemolysin activation/secretion protein??"           |
| K11035   | "hemolysin BL binding component"                     |
| K11037   | "hemolysin BL lytic component L1"                     |
| K11036   | "hemolysin BL lytic component L2"                     |
| K11003   | "hemolysin D"                                        |
| K11139   | "hemolysin E"                                        |
| K11032   | "hemolysin II"                                       |
| K11068   | "hemolysin III"                                      |
| K11016   | "hemolysin"                                          |
| K10948   | "hemolysin"                                          |
| K02498   | "HemY protein"                                       |
| K02848   | "heptose (I) phosphotransferase [EC:2.7.1.-]"        |
| K02841   | "heptosyltransferase I [EC:2.4.-.-]"                 |
| K12982   | "heptosyltransferase I [EC:2.4.-.-]"                 |
| K02843   | "heptosyltransferase II [EC:2.4.-.-]"                |
| K02849   | "heptosyltransferase III [EC:2.4.-.-]"               |
| K01590   | "histidine decarboxylase [EC:4.1.1.22]"             |
| K10016   | "histidine transport system permease protein"        |
| K13993   | "HSP20 family protein"                               |
| K00752   | "hyaluronan synthase [EC:2.4.1.212]"                |
| K01727   | "hyaluronate lyase [EC:4.2.2.1]"                     |
| K01197   | "hyaluronoglucosaminidase [EC:3.2.1.35]"            |
| K10814   | "hydrogen cyanide synthase HcnA [EC:1.4.99.5]"      |
| K10815   | "hydrogen cyanide synthase HcnB [EC:1.4.99.5]"      |
| K10816   | "hydrogen cyanide synthase HcnC [EC:1.4.99.5]"      |
| K01749   | "hydroxymethylbilane synthase [EC:2.5.1.61]"        |
| K09117   | "hypothetical protein"                               |
| K09790   | "hypothetical protein"                               |
| K09153   | "hypothetical protein"                               |
| K07226   | "hypothetical protein"                               |
| K09824   | "hypothetical protein"                               |
| K09860   | "hypothetical protein"                               |
| K07227   | "hypothetical protein"                               |
| K09152   | "hypothetical protein"                               |
| K01390   | "IgaA-specific metalloendopeptidase [EC:3.4.24.13]"  |
| K01347   | "IgaA-specific serine endopeptidase [EC:3.4.21.72]" |
| K09607   | "immune inhibitor A [EC:3.4.24.-]"                   |
| K14196   | "immunoglobulin G-binding protein A"                 |
| K14197   | "immunoglobulin G-binding protein Sbi"                |
| K11021   | "insecticidal toxin complex protein TccC"            |
| K13730   | "internalin A"                                       |
| K12206   | "intracellular multiplication protein lcmB"           |
| K12207   | "intracellular multiplication protein lcmC"           |
| K12208   | "intracellular multiplication protein lcmD"           |
| K12209   | "intracellular multiplication protein lcmE"           |
| K12211   | "intracellular multiplication protein lcmG"           |
| K12212   | "intracellular multiplication protein lcmJ"           |
| K12213   | "intracellular multiplication protein lcmK"           |
| K12214   | "intracellular multiplication protein lcmL"           |
| K12216   | "intracellular multiplication protein lcmN"           |
| K12217   | "intracellular multiplication protein lcmO"           |

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| #OTU ID  | KEGG Description                                                                 |
|----------|----------------------------------------------------------------------------------|
| K12218   | ["intracellular multiplication protein IcmP"]                                   |
| K12219   | ["intracellular multiplication protein IcmQ"]                                   |
| K12221   | ["intracellular multiplication protein IcmS"]                                   |
| K12222   | ["intracellular multiplication protein IcmT"]                                   |
| K12223   | ["intracellular multiplication protein IcmV"]                                   |
| K12224   | ["intracellular multiplication protein IcmW"]                                   |
| K12225   | ["intracellular multiplication protein IcmX"]                                   |
| K13285   | ["invasin B"]                                                                   |
| K13287   | ["invasin D"]                                                                   |
| K13745   | ["L-2,4-diaminobutyrate decarboxylase [EC:4.1.1.86]"]                            |
| K02014   | ["iron complex outermembrane receptor protein"]                                 |
| K02013   | ["iron complex transport system ATP-binding protein [EC:3.6.3.34]"]               |
| K02015   | ["iron complex transport system permease protein"]                               |
| K02016   | ["iron complex transport system substrate-binding protein"]                     |
| K02010   | ["iron(III) transport system ATP-binding protein [EC:3.6.3.30]"]                 |
| K02011   | ["iron(III) transport system permease protein"]                                  |
| K02506   | ["leader peptidase HopD [EC:3.4.23.43]"]                                         |
| K02517   | ["leader peptidase (prepilin peptidase)/ N-methyltransferase [EC:3.4.23.43 2.1.1-]""] |
| K02654   | ["LacI family transcriptional regulator"]                                        |
| K02529   | ["L.2,4-diaminobutyrate decarboxylase [EC:4.1.1.86]"]                            |
| K02560   | ["leukocidin/hemolysin toxin family protein"]                                    |
| K111038  | ["leader peptidase (prepilin peptidase)/ N-methyltransferase [EC:3.4.23.43 2.1.1-]""] |
| K02560   | ["leukocidin/hemolysin toxin family protein"]                                    |
| K02517   | ["leukocidin/hemolysin toxin family protein"]                                    |
| K025790  | ["Lipoate-protein ligase A [EC:2.7.7.63]""]                                       |
| K07271   | ["lipopolysaccharide cholinephosphotransferase [EC:2.7.8.-]""]                   |
| K09691   | ["lipopolysaccharide transport system ATP-binding protein""]                    |
| K09690   | ["lipopolysaccharide transport system permease protein""]                        |
| K11038   | ["long chain fatty acid CoA FadD26""]                                            |
| K01909   | ["long-chain fatty acid [acyl-carrier-protein] ligase [EC:6.2.1.20]""]            |
| K10112   | ["major pilin subunit PapA""]                                                    |
| K07345   | ["major type 1 subunit fimbrin (pilin)""]                                         |
| K12989   | ["mannosylglycoprotein endo-beta-N-acetylglucosaminidase [EC:3.2.1.96]""]        |
| K00971   | ["mannosylglycosyltransferase [EC:2.4.1.-]""]                                     |

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| #OTU ID | KEGG Description |
|---------|------------------|
| K13001  | "mannosyltransferase [EC:2.4.1.-]" |
| K05375  | "MbtH protein" |
| K01273  | "membrane dipeptidase [EC:3.4.13.19]" |
| K07058  | "membrane protein" |
| K14743  | "membrane-anchored mycosin MYCP [EC:3.4.21.-]" |
| K08305  | "membrane-bound lytic murein transglycosylase B [EC:3.2.1.-]" |
| K08307  | "membrane-bound lytic murein transglycosylase D [EC:3.2.1.-]" |
| K13640  | "MerR family transcriptional regulator, heat shock protein HspR" |
| K00604  | "methionyl-tRNA formyltransferase [EC:2.1.2.9]" |
| K05874  | "methyl-accepting chemotaxis protein I, serine sensor receptor" |
| K05875  | "methyl-accepting chemotaxis protein II, aspartate sensor receptor" |
| K03406  | "methyl-accepting chemotaxis protein" |
| K07522  | "MFS transporter, DHAI family, bicyclomycin/chloramphenicol resistance protein" |
| K02429  | "MFS transporter, FHS family, L-fucose permease" |
| K02575  | "MFS transporter, NNP family, nitrate/nitrite transporter" |
| K07784  | "MFS transporter, OPA family, hexose phosphate transport protein UhpT" |
| K05373  | "MFS transporter, putative signal transducer" |
| K01531  | "Mg2+-importing ATPase [EC:3.6.3.2]" |
| K01387  | "microbial collagenase [EC:3.4.24.3]" |
| K01174  | "micrococcal nuclease [EC:3.1.31.1]" |
| K04335  | "minor curlin subunit" |
| K07349  | "minor fimbrial subunit" |
| K07348  | "minor fimbrial subunit" |
| K07350  | "minor fimbrial subunit" |
| K04043  | "molecular chaperone DnaK" |
| K04080  | "molecular chaperone LbpA" |
| K03455  | "monovalent cation:H+ antiporter-2, CPA2 family" |
| K12276  | "MSHA biogenesis protein MshE" |
| K12277  | "MSHA biogenesis protein MshF" |
| K12278  | "MSHA biogenesis protein MshG" |
| K12279  | "MSHA biogenesis protein MshH" |
| K12280  | "MSHA biogenesis protein MshI" |
| K12281  | "MSHA biogenesis protein MshK" |
| K12282  | "MSHA biogenesis protein MshL" |
| K12283  | "MSHA biogenesis protein MshM" |
| K12284  | "MSHA biogenesis protein MshN" |
| K12285  | "MSHA biogenesis protein MshO" |
| K12286  | "MSHA biogenesis protein MshP" |
| K12287  | "MSHA biogenesis protein MshQ" |
| K10924  | "MSHA pilin protein MshA" |
| K10925  | "MSHA pilin protein MshB" |
| K10926  | "MSHA pilin protein MshC" |
| K10927  | "MSHA pilin protein MshD" |
| K02026  | "multiple sugar transport system permease protein" |
| K02025  | "multiple sugar transport system permease protein" |
| K02027  | "multiple sugar transport system substrate-binding protein" |
| K04793  | "mycobactin lysine-N-oxygenase" |
| K04789  | "mycobactin peptide synthetase MbtE" |
| K04792  | "mycobactin peptide synthetase MbtF" |
| K04788  | "mycobactin phenylazoxazoline synthetase" |
| K04790  | "mycobactin polyketide synthetase MbtC" |
| K04791  | "mycobactin polyketide synthetase MbtD" |
| K04787  | "mycobactin salicyl-AMP ligase [EC:6.3.2.-]" |
| K08068  | "N-acetylglucosamine-6-phosphate 2-epimerase and phosphatase [EC:5.1.3.-]" |
| K05946  | "N-acetylglucosaminylidophosphodecaprenol [EC:2.4.1.187]" |
| K01447  | "N-acetylmuramoyl-L-alanine amidase [EC:3.5.1.28]" |
| K01654  | "N-acetylneuraminate synthase [EC:2.5.1.56]" |
| K00983  | "N-acetylneuraminate cytidylyltransferase [EC:2.7.7.43]" |
| K00343  | "NADH dehydrogenase I subunit N [EC:1.6.5.3], "NADH-quinone oxidoreductase subunit N [EC:1.6.5.3]" |

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| #OTU ID | KEGG Description |
|---------|------------------|
| K00336  | "NADH-quinone oxidoreductase subunit G [EC:1.6.5.3]", "NADH dehydrogenase I subunit G [EC:1.6.5.3]" |
| K02398  | "negative regulator of flagellin synthesis FlgM" |
| K01389  | "neprilysin [EC:3.4.24.11]" |
| K00370  | "nitrate reductase 1, alpha subunit [EC:1.7.99.4]" |
| K00371  | "nitrate reductase 1, beta subunit [EC:1.7.99.4]" |
| K00373  | "nitrate reductase 1, delta subunit [EC:1.7.99.4]" |
| K00374  | "nitrate reductase 1, gamma subunit [EC:1.7.99.4]" |
| K02003  | "None" |
| K02004  | "None" |
| K00257  | "None" |
| K07001  | "None" |
| K07145  | "None" |
| K06998  | "None" |
| K07043  | "None" |
| K07154  | "None" |
| K07126  | "None" |
| K07011  | "None" |
| K06867  | "None" |
| K07018  | "None" |
| K07164  | "None" |
| K01795  | "None" |
| K07017  | "None" |
| K00786  | "None" |
| K06921  | "None" |
| K07031  | "None" |
| K06900  | "None" |
| K06887  | "None" |
| K06882  | "None" |
| K01732  | "None" |
| K11033  | "non-hemolytic enterotoxin A" |
| K11034  | "non-hemolytic enterotoxin B/C" |
| K12237  | "nonribosomal peptide synthetase VibF" |
| K00940  | "nucleoside-diphosphate kinase [EC:2.7.4.6]" |
| K02428  | "nucleoside-triphosphate pyrophosphatase [EC:3.6.1.19]" |
| K13012  | "O-antigen biosynthesis protein WbpP" |
| K03286  | "OmpA-OmpF porin, OOP family" |
| K07268  | "opacity associated protein" |
| K00611  | "ornithine carbamoyltransferase [EC:2.1.3.3]" |
| K01750  | "ornithine cyclodeaminase [EC:4.3.1.12]" |
| K12340  | "outer membrane channel protein TolC" |
| K12686  | "outer membrane lipase/esterase" |
| K02494  | "outer membrane lipoprotein LolB" |
| K12538  | "outer membrane protein HasF" |
| K10940  | "outer membrane protein OmpT" |
| K08720  | "outer membrane protein OmpU" |
| K07277  | "outer membrane protein" |
| K12518  | "outer membrane usher protein PapC" |
| K0347   | "outer membrane usher protein" |
| K02495  | "oxygen-independent coproporphyrinogen III oxidase [EC:1.3.99.22]" |
| K12973  | "palmitoyl transferase [EC:2.3.1.-]" |
| K00954  | "pantheine-phosphate adenylyltransferase [EC:2.7.7.3]" |
| K01918  | "pantoate-beta-alanine ligase [EC:6.3.2.1]" |
| K12267  | "peptide methionine sulfoxide reductase msrA/msrB [EC:1.8.4.11 1.8.4.12]" |
| K02032  | "peptide/nickel transport system ATP-binding protein" |
| K03640  | "peptidoglycan-associated lipoprotein" |
| K03767  | "peptidyl-prolyl cis-trans isomerase A (cyclophilin A) [EC:5.2.1.8]" |
| K03832  | "periplasmic protein TonB" |
| K13010  | "perosamine synthetase" |
| K03386  | "peroxiredoxin (alkyl hydroperoxide reductase subunit C) [EC:1.11.1.15]" |
| K12681  | "pertactin" |

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Table 2 (continued)

| #OTU ID   | KEGG Description                                                                 |
|-----------|----------------------------------------------------------------------------------|
| K13063    | ['phenazine biosynthesis protein phzE [EC:2.6.1.86]']                           |
| K12440    | ['phenolthiocerol synthesis type-I polyketide synthase A']                      |
| K12441    | ['phenolthiocerol synthesis type-I polyketide synthase B']                      |
| K12443    | ['phenolthiocerol synthesis type-I polyketide synthase D']                      |
| K12444    | ['phenolthiocerol synthesis type-I polyketide synthase E']                      |
| K04750    | ['PhnB protein']                                                                |
| K00981    | ['phosphatidate cytidyltransferase [EC:2.7.7.41]']                              |
| K13292    | ['phosphatidylglycerol:prolipoprotein diacylglycerol transferase [EC:2.-.-.-]']  |
| K03760    | ['phosphoethanolamine transferase']                                             |
| K01835    | ['phosphoglucomutase [EC:5.4.2.2]']                                             |
| K03431    | ['phosphoglucomutase mutase [EC:5.4.2.10]']                                     |
| K03271    | ['phosphoheptose isomerase [EC:5.-.-.-]']                                       |
| K01114    | ['phospholipase C [EC:3.1.4.3]']                                                |
| K01840    | ['phosphomannomutase [EC:5.4.2.8]']                                             |
| K01923    | ['phthiobiosylaminomimidazole-succinocarboxamide synthase [EC:6.3.2.6]']       |
| K14728    | ['phthiobiolone/phenolphthiobiolone dimycocerosates ketoreductase [EC:1.2.-.-]'] |
| K02279    | ['pilus assembly protein CpaB']                                                 |
| K02280    | ['pilus assembly protein CpaC']                                                 |
| K02281    | ['pilus assembly protein CpaD']                                                 |
| K02282    | ['pilus assembly protein CpaE']                                                 |
| K02283    | ['pilus assembly protein CpaF']                                                 |
| K02651    | ['pilus assembly protein Flp/PilA']                                             |
| K13925    | ['plasmin and fibronectin-binding protein A']                                   |
| K08566    | ['plasminogen activator [EC:3.4.23.48]']                                        |
| K01729    | ['poly(beta-D-mannuronate) lyase [EC:4.2.2.3]']                                  |
| K07282    | ['poly-gamma-glutamate synthesis protein (capsule biosynthesis protein)']      |
| K12430    | ['polyketide synthase 1/15']                                                   |
| K12436    | ['polyketide synthase 12']                                                      |
| K12433    | ['polyketide synthase 5']                                                       |
| K12434    | ['polyketide synthase 7']                                                       |
| K00937    | ['polyphosphate kinase [EC:2.7.4.1]']                                           |
| K01991    | ['polysaccharide export outer membrane protein']                                |
| K03328    | ['polysaccharide transporter, PST family']                                     |
| K01698    | ['porphobilinogen synthase [EC:4.2.1.24]']                                     |
| K02278    | ['prepilin peptidase CpaA [EC:3.4.23.43]']                                     |
| K02682    | ['prepilin peptidase dependent protein D']                                     |
| K03070    | ['preprotein translocase subunit SecA']                                         |
| K12537    | ['protease secretion protein HasE']                                            |
| K13571    | ['proteasome accessory factor A [EC:6.3.2.-]']                                  |
| K13527    | ['proteasome-associated ATPase']                                                |
| K11030    | ['protective antigen']                                                          |
| K13743    | ['protein IpgB2']                                                               |
| K14204    | ['protein Map']                                                                 |
| K02504    | ['protein transport protein HofB']                                             |
| K02505    | ['protein transport protein HofC']                                             |
| K02507    | ['protein transport protein HofQ']                                             |
| K01104    | ['protein-tyrosine phosphatase [EC:3.1.3.48]']                                  |
| K03305    | ['proton-dependent oligopeptide transporter, POT family']                      |
| K00230    | ['protoporphyrinogen oxidase [EC:1.3.3.4]']                                     |
| K01399    | ['pseudolysin [EC:3.4.24.26]']                                                  |
| K03408    | ['purine-binding chemotaxis protein CheW']                                     |
| K07804    | ['putative virulence related protein PagC']                                     |
| K02021    | ['putative ABC transport system ATP-binding protein']                           |
| K02067    | ['putative ABC transport system substrate-binding protein']                    |
| K06160    | ['putative ATP-binding cassette transporter']                                   |
| K06132    | ['putative cardiolipin synthase [EC:2.7.8.-]']                                  |
| K03606    | ['putative colanic acid biosynthesis UDP-glucose lipid carrier transferase']    |
| K06994    | ['putative drug exporter of the RND superfamily']                               |
| K07386    | ['putative endopeptidase [EC:3.4.24.-]']                                        |
| K07010    | ['putative glutamine amidotransferase']                                         |
| K10039    | ['putative glutamine transport system substrate-binding protein']              |

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Table 2 (continued)

| #OTU ID | KEGG Description |
|---------|------------------|
| K07225  | "putative hemin transport protein" |
| K06442  | "putative hemolysin" |
| K03699  | "putative hemolysin" |
| K07507  | "putative Mg2+ transporter-C (MgtC) family protein" |
| K07497  | "putative transposase" |
| K07498  | "putative transposase" |
| K12238  | "pyochelin biosynthesis protein PchD" |
| K12242  | "pyochelin biosynthetic protein PchC" |
| K12240  | "pyochelin synthetase" |
| K03474  | "pyridoxine 5-phosphate synthase [EC:2.6.99.2]" |
| K06137  | "pyrroloquinoline-quinone synthase [EC:1.3.3.11]" |
| K03789  | "ribosomal RNA small subunit methyltransferase E [EC:2.1.1.1]" |
| K03087  | "RNA polymerase nonessential primary-like sigma factor" |
| K03086  | "RNA polymerase primary sigma factor" |
| K02405  | "RNA polymerase sigma factor for flagellar operon FlIA" |
| K03092  | "RNA polymerase sigma-54 factor" |
| K03090  | "RNA polymerase sigma-B factor" |
| K01851  | "salicylate biosynthesis isochorismate synthase [EC:5.4.4.2]" |
| K04781  | "salicylate synthetase [EC:5.4.4.2 4.2.99.21], "salicylate synthetase [EC:5.4.4.2 4.13.-]" |
| K06024  | "segregation and condensation protein B" |
| K00640  | "serine O-acetyltransferase [EC:2.3.1.30]" |
| K12684  | "serine protease autotransporter [EC:3.4.21.-]" |
| K04691  | "serine protease DegS [EC:3.4.21.-]" |
| K04771  | "serine protease Do [EC:3.4.21.107]" |
| K08884  | "serine/threonine protein kinase, bacterial [EC:2.7.11.1]" |
| K11915  | "serine/threonine protein phosphatase Stp1 [EC:3.1.3.16]" |
| K14949  | "serine/threonine-protein kinase PknG [EC:2.7.11.11]" |
| K11912  | "serine/threonine-protein kinase PpkA [EC:2.7.11.1]" |
| K11916  | "serine/threonine-protein kinase Stk1 [EC:2.7.11.-]" |
| K14194  | "serine-aspartate repeat-containing protein C(D/E)" |
| K01406  | "serralysin [EC:3.4.24.40]" |
| K12683  | "serum resistance protein" |
| K01186  | "sialidase-1 [EC:3.2.1.18]" |
| K11914  | "sigma-54 dependent transcriptional regulator" |
| K11908  | "sigma-54 specific transcriptional regulator" |
| K11917  | "sigma-54 specific transcriptional regulator" |
| K10941  | "sigma-54 specific transcriptional regulator, flagellar regulatory protein A" |
| K07315  | "sigma-B regulation protein RsBl (phosphoserine phosphatase)" |
| K03597  | "sigma-E factor negative regulatory protein RseA" |
| K03598  | "sigma-E factor negative regulatory protein RseB" |
| K03803  | "sigma-E factor negative regulatory protein RseC" |
| K03101  | "signal peptidase II [EC:3.4.23.36]" |
| #OTU ID | KEGG Description |
|--------|------------------|
| K05813 | 
K07284 | ["sn-glycerol 3-phosphate transport system substrate-binding protein"] |
| K08600 | ["sortase A", "sortase A [EC:3.4.22.70]"] |
| K01117 | ["sphingomyelin phosphodiesterase [EC:3.1.4.12]"] |
| K07173 | ["S-ribosylhomocysteine lyase [EC:4.4.1.21]"] |
| K08258 | ["staphopain A [EC:3.4.22.48]"] |
| K13715 | ["staphopain B [EC:3.4.22.70]"] |
| K17040 | ["staphylococcal enterotoxin"] |
| K04047 | ["starvation-inducible DNA-binding protein"] |
| K12685 | ["streptolysin S associated protein"] |
| K00300 | ["sphingomyelin phosphodiesterase [EC:3.4.22.70]"] |
| K06019 | ["superoxide dismutase, Fe-Mn family [EC:1.15.1.1]"] |
| K08693 | ["teichoic acid transport system ATP-binding protein [EC:3.6.3.40]"] |
| K08692 | ["teichoic acid transport system permease protein"] |
| K00912 | ["tetracyclylacylcarbamide 4-kinase [EC:2.7.1.130]"] |
| K11018 | ["thiol-activated cytolysin"] |
| K12510 | ["tight adherence protein B"] |
| K12511 | ["tight adherence protein C"] |
| K12512 | ["tight adherence protein D"] |
| K12513 | ["tight adherence protein E"] |
| K12514 | ["tight adherence protein F"] |
| K12515 | ["tight adherence protein G"] |
| K11043 | ["toxic shock syndrome toxin-1"] |
| K11063 | ["toxin A/B"] |
| K10930 | ["toxin co-regulated pilin"] |
| K10932 | ["toxin co-regulated pilus biosynthesis outer membrane protein C"] |
| K10931 | ["toxin co-regulated pilus biosynthesis protein B"] |
| K10933 | ["toxin co-regulated pilus biosynthesis protein D"] |
| K10934 | ["toxin co-regulated pilus biosynthesis protein E"] |
| K10935 | ["toxin co-regulated pilus biosynthesis protein F"] |
| K10919 | ["toxin co-regulated pilus biosynthesis protein H"] |
| K10919 | ["toxin co-regulated pilus biosynthesis protein I"] |
| K10966 | ["toxin co-regulated pilus biosynthesis protein J [EC:3.4.23.43 2.1.1.-]"] |
| K10920 | ["toxin co-regulated pilus biosynthesis protein P"] |
| K10962 | ["toxin co-regulated pilus biosynthesis protein Q"] |
| K10963 | ["toxin co-regulated pilus biosynthesis protein R"] |
| K10964 | ["toxin co-regulated pilus biosynthesis protein S"] |
| K10965 | ["toxin co-regulated pilus biosynthesis protein T"] |
| K07165 | ["transmembrane sensor"] |
| K01046 | ["triacylglycerol lipase [EC:3.1.1.3]"] |
| K03545 | ["trigger factor"] |
| K02659 | ["twitching motility protein PilL"] |
| K02660 | ["twitching motility protein PilJ"] |
| K02669 | ["twitching motility protein PilT"] |
| K02670 | ["twitching motility protein PilU"] |
| K02657 | ["twitching motility two-component system response regulator PilG"] |
| K02658 | ["twitching motility two-component system response regulator PilH"] |
| K10943 | ["two-component system, response regulator FlrC"] |
| K07707 | ["two-component system, AgrA family, response regulator AgrA"] |
| K07706 | ["two-component system, AgrA family, sensor histidine kinase AgrC [EC:2.7.13.-]"] |
| K03412 | ["two-component system, chemotaxis family, response regulator CheB [EC:3.1.1.61]"] |
| K03415 | ["two-component system, chemotaxis family, response regulator CheV"] |
| K03413 | ["two-component system, chemotaxis family, response regulator CheY"] |
| K03407 | ["two-component system, chemotaxis family, sensor kinase CheA [EC:2.7.13.3]"] |
| K08083 | ["two-component system, LytF family, response regulator AlgR"] |
| K08082 | ["two-component system, LytF family, sensor histidine kinase AlgZ [EC:2.7.13.3]"] |
| K07689 | ["two-component system, NarL family, invasion response regulator UvrY"] |

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| #OTU ID  | KEGG Description                                                                 |
|---------|----------------------------------------------------------------------------------|
| K07695  | [“two-component system, NarL family, response regulator DevR”]                   |
| K07690  | [“two-component system, NarL family, response regulator EvgA”]                  |
| K02479  | [“two-component system, NarL family, response regulator”]                       |
| K07688  | [“two-component system, NarL family, response regulator, fimbrial Z protein, FimZ”] |
| K07678  | [“two-component system, NarL family, sensor histidine kinase BarA [EC:2.7.13.3]”] |
| K07682  | [“two-component system, NarL family, sensor histidine kinase DevS [EC:2.7.13.3]”] |
| K07679  | [“two-component system, NarL family, sensor histidine kinase EvgS [EC:2.7.13.3]”] |
| K11384  | [“two-component system, NtrC family, response regulator AlgB”]                 |
| K02667  | [“two-component system, NtrC family, response regulator PilR”]                 |
| K02481  | [“two-component system, NtrC family, response regulator”]                      |
| K07710  | [“two-component system, NtrC family, sensor histidine kinase AtoS [EC:2.7.13.3]”] |
| K02668  | [“two-component system, NtrC family, sensor histidine kinase PilS [EC:2.7.13.3]”] |
| K02482  | [“two-component system, NtrC family, sensor kinase [EC:2.7.13.3]”]             |
| K07669  | [“two-component system, OmpR family, response regulator MprA”]                 |
| K07660  | [“two-component system, OmpR family, response regulator PhoP”]                 |
| K07671  | [“two-component system, OmpR family, response regulator RssA”]                |
| K07776  | [“two-component system, OmpR family, response regulator RegX3”]                |
| K07661  | [“two-component system, OmpR family, response regulator RstA”]                |
| K02483  | [“two-component system, OmpR family, response regulator”]                     |
| K07653  | [“two-component system, OmpR family, sensor histidine kinase MprB [EC:2.7.13.3]”] |
| K07637  | [“two-component system, OmpR family, sensor histidine kinase PhoQ [EC:2.7.13.3]”] |
| K07655  | [“two-component system, OmpR family, sensor histidine kinase PrrB [EC:2.7.13.3]”] |
| K07639  | [“two-component system, OmpR family, sensor histidine kinase RstB [EC:2.7.13.3]”] |
| K07768  | [“two-component system, OmpR family, sensor histidine kinase SenX3 [EC:2.7.13.3]”] |
| K02484  | [“two-component system, OmpR family, sensor kinase [EC:2.7.13.3]”]             |
| K10942  | [“two-component system, sensor histidine kinase FlrB [EC:2.7.13.3]”]           |
| K07357  | [“type 1 fimbriae regulatory protein FimB”]                                     |
| K07358  | [“type 1 fimbriae regulatory protein FimE”]                                     |
| K04049  | [“type III secretion protein SctB”]                                             |
| K03219  | [“type III secretion protein SctC”]                                             |
| K03220  | [“type III secretion protein SctD”]                                             |
| K04050  | [“type III secretion protein SctE”]                                             |
| K03221  | [“type III secretion protein SctF”]                                             |
| K04051  | [“type III secretion protein SctG”]                                             |
| K04052  | [“type III secretion protein SctH”]                                             |
| K04053  | [“type III secretion protein SctI”]                                             |
| K03222  | [“type III secretion protein SctJ”]                                             |
| K04054  | [“type III secretion protein SctK”]                                             |
| K03223  | [“type III secretion protein SctL”]                                             |
| K04056  | [“type III secretion protein SctO”]                                             |
| K04057  | [“type III secretion protein SctP”]                                             |
| K03225  | [“type III secretion protein SctQ”]                                             |
| K03226  | [“type III secretion protein SctR”]                                             |
| K03227  | [“type III secretion protein SctS”]                                             |
| K03228  | [“type III secretion protein SctT”]                                             |
| K03229  | [“type III secretion protein SctU”]                                             |
| K03230  | [“type III secretion protein SctV”]                                             |
| K04058  | [“type III secretion protein SctW”]                                             |
| K04059  | [“type III secretion protein SctX”]                                             |
| K04060  | [“type III secretion protein SctY”]                                             |
| K08084  | [“type IV fimbrial biogenesis protein FimT”]                                     |
| K08085  | [“type IV fimbrial biogenesis protein FimU”]                                     |
| K02487  | [“type IV pilus sensor histidine kinase and response regulator”]               |
| K02650  | [“type IV pilus assembly protein PilA”]                                          |
| K02652  | [“type IV pilus assembly protein PilB”]                                          |
| K02653  | [“type IV pilus assembly protein PilC”]                                          |
| K02655  | [“type IV pilus assembly protein PilE”]                                          |
| K02656  | [“type IV pilus assembly protein PilF”]                                          |
| K02661  | [“type IV pilus assembly protein PilK”]                                          |
| K02662  | [“type IV pilus assembly protein PilM”]                                          |
| #OTU ID   | KEGG Description                                                                 |
|---------|----------------------------------------------------------------------------------|
| K02663  | ["type IV pilus assembly protein PilN"]                                         |
| K02664  | ["type IV pilus assembly protein PilO"]                                         |
| K02665  | ["type IV pilus assembly protein PilP"]                                         |
| K02666  | ["type IV pilus assembly protein PilQ"]                                         |
| K02671  | ["type IV pilus assembly protein PilV"]                                         |
| K02672  | ["type IV pilus assembly protein PilW"]                                         |
| K02673  | ["type IV pilus assembly protein PilX"]                                         |
| K02674  | ["type IV pilus assembly protein PilY1"]                                        |
| K02676  | ["type IV pilus assembly protein PilZ"]                                         |
| K03194  | ["type IV secretion system protein VirB1"]                                      |
| K03195  | ["type IV secretion system protein VirB10"]                                     |
| K03196  | ["type IV secretion system protein VirB11"]                                     |
| K03197  | ["type IV secretion system protein VirB2"]                                      |
| K03198  | ["type IV secretion system protein VirB3"]                                      |
| K03199  | ["type IV secretion system protein VirB4"]                                      |
| K03200  | ["type IV secretion system protein VirB5"]                                      |
| K03201  | ["type IV secretion system protein VirB6"]                                      |
| K03202  | ["type IV secretion system protein VirB7"]                                      |
| K03203  | ["type IV secretion system protein VirB8"]                                      |
| K03204  | ["type IV secretion system protein VirB9"]                                      |
| K03205  | ["type IV secretion system protein VirD4"]                                      |
| K11919  | ["type VI secretion system lysozyme-related protein"]                           |
| K11902  | ["type VI secretion system protein ImpA"]                                       |
| K11901  | ["type VI secretion system protein ImpB"]                                       |
| K11900  | ["type VI secretion system protein ImpC"]                                       |
| K11899  | ["type VI secretion system protein ImpD"]                                       |
| K11898  | ["type VI secretion system protein ImpE"]                                       |
| K11897  | ["type VI secretion system protein ImpF"]                                       |
| K11896  | ["type VI secretion system protein ImpG"]                                       |
| K11895  | ["type VI secretion system protein ImpH"]                                       |
| K11894  | ["type VI secretion system protein ImpI"]                                       |
| K11893  | ["type VI secretion system protein ImpJ"]                                       |
| K11892  | ["type VI secretion system protein ImpK"]                                       |
| K11891  | ["type VI secretion system protein ImpL"]                                       |
| K11890  | ["type VI secretion system protein ImpM"]                                       |
| K11889  | ["type VI secretion system protein ImpN [EC:2.7.11.1]"                         |
| K11906  | ["type VI secretion system protein VasD"]                                       |
| K11907  | ["type VI secretion system protein VasG"]                                       |
| K11909  | ["type VI secretion system protein VasI"]                                       |
| K11910  | ["type VI secretion system protein VasJ"]                                       |
| K11911  | ["type VI secretion system protein VasL"]                                       |
| K11905  | ["type VI secretion system protein"]                                            |
| K11913  | ["type VI secretion system protein"]                                             |
| K11918  | ["type VI secretion system protein"]                                             |
| K11903  | ["type VI secretion system secreted protein Hcp"]                               |
| K11904  | ["type VI secretion system secreted protein VgrG" ]                              |
| K03269  | ["UDP-2,3-diacylglucosamine hydrolase [EC:3.6.1.-]" ]                             |
| K13017  | ["UDP-3-keto-D-GlcNAc aminotransferase [EC:2.6.1.-]" ]                           |
| K02536  | ["UDP-3-O-[3-hydroxyxymristoyl] glucosamine N-acyltransferase [EC:2.3.1.-]" ]    |
| K02535  | ["UDP-3-O-[3-hydroxyxymristoyl] N-acetylgulcosamine deacetylase [EC:3.5.1.-]" ]  |
| K13018  | ["UDP-D-GlcNAc3NA acetyltransferase [EC:2.3.1.-]" ]                              |
| K13020  | ["UDP-D-GlcNAc oxidase [EC:1.1.1.-]" ]                                           |
| K13016  | ["UDP-D-GlcNAc oxidase [EC:1.1.1.-]" ]                                           |
| K01854  | ["UDP-galactopyranose mutase [EC:5.4.99.9]" ]                                   |
| K13019  | ["UDP-GlcNAc3NAc epimerase [EC:5.1.3.23]" ]                                     |
| K10011  | ["UDP-GlcUA decarboxylase/UDP-L-Ara4N formyltransferase [EC:1.1.1.-]" ]          |
| K01784  | ["UDP-glucose 4-epimerase [EC:5.1.3.2]" ]                                        |
| K00012  | ["UDPGlucose 6-dehydrogenase [EC:1.1.1.22]" ]                                  |

(continued on next page)
Table 2 (continued)

| #OTU ID  | KEGG Description                                                                                                                                 |
|----------|-------------------------------------------------------------------------------------------------------------------------------------------------|
| K03279   | ['UDP-glucose:(galactosyl)LPS alpha-1,2-glucosyltransferase [EC:2.4.1.58]']                                                                     |
| K02844   | ['UDP-glucose:(heptosyl)LPS alpha-1,3-glucosyltransferase [EC:2.4.1.-]']                                                                         |
| K02474   | ['UDP-N-acetyl-D-galactosamine dehydrogenase [EC:1.1.1.-]']                                                                                   |
| K13015   | ['UDP-N-acetyl-D-glucosamine dehydrogenase [EC:1.1.1.-]']                                                                                     |
| K02472   | ['UDP-N-acetyl-D-mannosaminuronic acid dehydrogenase [EC:1.1.1.-]']                                                                         |
| K01791   | ['UDP-N-acetylglucosamine 2-epimerase [EC:5.1.3.14]']                                                                                         |
| K02473   | ['UDP-N-acetylglucosamine 4-epimerase [EC:5.1.3.7]']                                                                                          |
| K00677   | ['UDP-N-acetylglucosamine acyltransferase [EC:2.3.1.129]']                                                                                      |
| K00075   | ['UDP-N-acetylmuramate dehydrogenase [EC:1.1.1.158]']                                                                                            |
| K01138   | ['uncharacterized sulfatase [EC:3.1.6.-]']                                                                                                 |
| K00806   | ['undecaprenyl pyrophosphate synthetase [EC:2.5.1.31]”, “undecaprenyl diphosphate synthase [EC:2.5.1.31]”']                                   |
| K02851   | ['undecaprenyl-phosphate alpha-N-acetylglucosaminytransferase [EC:2.7.8.-]']                                                                   |
| K00996   | ['undecaprenyl-phosphate galactose phosphotransferase [EC:2.7.8.6]']                                                                         |
| K03188   | ['urease accessory protein']                                                                                                                  |
| K03190   | ['urease accessory protein']                                                                                                                  |
| K03189   | ['urease accessory protein']                                                                                                                  |
| K03187   | ['urease accessory protein']                                                                                                                  |
| K01428   | ['urease alpha subunit [EC:3.5.1.5]”, “urease subunit alpha [EC:3.5.1.5]”']                                                                   |
| K14048   | ['urease subunit gamma/beta [EC:3.5.1.5]”']                                                                                                  |
| K02496   | ['uroporphin-III C-methyltransferase [EC:2.1.1.107]”']                                                                                         |
| K01599   | ['uroporphinogen decarboxylase [EC:4.1.1.37]”']                                                                                               |
| K01719   | ['uroporphinogen-III synthase [EC:4.2.1.75]”']                                                                                                |
| K00963   | ['UTP-glucose-1-phosphate uridylyltransferase [EC:2.7.7.9]”']                                                                                 |
| K04778   | ['vibriobactin synthetase”']                                                                                                                  |
| K08604   | ['vibriolysin [EC:3.4.24.25]”']                                                                                                                |
| K03980   | ['virulence factor”']                                                                                                                        |
| K04784   | ['yersiniabactin nonribosomal peptide synthetase”']                                                                                           |
| K04786   | ['yersiniabactin nonribosomal peptide/polyketide synthase”']                                                                                   |
| K04783   | ['yersiniabactin salicyl-AMP ligase [EC:6.3.2.-]”']                                                                                           |
| K04785   | ['yersiniabactin synthetase, thiazolinyl reductase component”']                                                                               |
| K05374   | ['yersiniabactin synthetase, thioesterase component”']                                                                                         |
| K08598   | ['YopJ protease family”']                                                                                                                     |
| K08643   | ['zinc metalloprotease ZmpB [EC:3.4.24.-]”']                                                                                                   |
| K09815   | ['zinc transport system substrate-binding protein”']                                                                                           |
| K02074   | ['zinc/manganese transport system ATP-binding protein”']                                                                                       |
| K02075   | ['zinc/manganese transport system permease protein”']                                                                                          |
| K02077   | ['zinc/manganese transport system substrate-binding protein”']                                                                                |
| K10954   | ['zona occludens toxin”']                                                                                                                     |

2.2. 16S rRNA gene sequencing

Samples were vortexed to loosen the bacteria from the swab, and then centrifuged to form a bacteria pellet. Cell pellets were frozen at −80 °C within 24 h of collection and submitted for analysis at MRDNA/Molecular Research LP (Shallowater, TX.). There, genomic DNA was extracted from the cell pellets using DNeasy PowerSoil Extraction Kit (Qiagen, USA) according to manufacturer instructions. A single-step 30 cycle PCR was performed for the 16S rRNA gene V4 variable region using PCR primers 515F/806R (515F: 5’-GTGCCAGCMGCCGCGGTAA-3’; and 806R: 5’-GGACTACVGGGTATCTAAAT-3’). PCR conditions were: 94 °C for 3 minutes, followed by 28 cycles of 94 °C for 30 seconds, 53 °C for 40 seconds and 72 °C for 1 minute, and a final elongation step at 72 °C for 5 minutes. NextGen Sequencing (NGS) for the 16S rRNA gene V4 variable region was performed on an Ion Torrent PGM at MRDNA/Molecular Research LP [1].

2.3. Data analysis

The raw sequencing data were analyzed using QIIME version 1.9.1 [2]. Raw sequencing reads were demultiplexed using split_libraries.py. The sequences were filtered using a cut-off
quality score of 25 and were then clustered into an open reference operational taxonomic unit (OTU) table using pick_open_reference_ots.py and taxonomically classified. The sequences were filtered using filter_taxa_from_otu_table.py to remove chloroplasts, and diversity analyses were performed using core_diversity_analyses.py. These data were further analyzed using PICRUSt version 1.1.4 to predict functional profiles of the bacteria and compared to shotgun metagenomic sequencing to reveal changes in ARGs, VFGs and antibiotic biosynthesis genes (ABSGs) after swimming. The following PICRUSt scripts were used to acquire gene counts: normalize_by_copy_number.py, predict_metagenomes.py, categorize_by_function.py and metagenome_contributions.py. Weighted nearest sequenced taxon index (NSTI) scores for each sample were calculated to assess prediction accuracy using the predict_metagenomes.py with the -a option. In order to obtain OTU-specific gene counts for ARGs and VFGs, we used the metagenome_contributions.py script with -l option for each gene of interest detected in the predicted profiles [8,9]. These gene counts, referred to as KOs [10], were compared to previously published literature on ARGs [8,11] and virulence factors [12], and with the KO database to determine which ARGs (Table 1) and VFGs (Table 2) were detected in the samples.

Ethics Statement

Data collection methods were approved by the University of California, Irvine Institutional Review Board (IRB #2017-3751). Participants gave verbal consent to enroll.

CRediT Author Statement

Marisa Nielsen: Conception, Data Curation, Formal Analysis, Visualization, Investigation, Writing-Original Draft Preparation; Sunny Jiang: Conception, Writing-Reviewing and Editing, Investigation, Supervision. All authors read and approve the manuscript.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships which have or could be perceived to have influenced the work reported in this article.

Acknowledgements

Funding for this research was partially provided by NSF CBET 1806066, 1804166 and UCI Oceans Fellowship.

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