1	Distinctive Gene and Protein Characteristics of Extremely Piezophilic Colwellia
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24 Background

The deep ocean is characterized by low temperatures, high hydrostatic pressures, and low
concentrations of organic matter. While these conditions likely select for distinct genomic
characteristics within prokaryotes, the attributes facilitating adaptation to the deep ocean are
relatively unexplored. In this study, we compared the genomes of seven strains within the genus *Colwellia*, including some of the most piezophilic microbes known, to identify genomic features
that enable life in the deep sea.

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32 **Results**

33 Significant differences were found to exist between piezophilic and non-piezophilic strains of 34 Colwellia. Piezophilic Colwellia have a more basic and hydrophobic proteome. The piezophilic 35 abyssal and hadal isolates have more genes involved in replication/recombination/repair, cell 36 wall/membrane biogenesis, and cell motility. The characteristics of respiration, pilus generation, 37 and membrane fluidity adjustment vary between the strains, with operons for a *nuo* 38 dehydrogenase and a *tad* pilus only present in the piezophiles. In contrast, the piezosensitive 39 members are unique in having the capacity for dissimilatory nitrite and TMAO reduction. A 40 number of genes exist only within deep-sea adapted species, such as those encoding d-alanine-d-41 alanine ligase for peptidoglycan formation, alanine dehydrogenase for NADH/NAD⁺ 42 homeostasis, and archaeal methyltransferase for tRNA modification. Many of these piezophile-43 specific genes are in variable regions of the genome near genomic islands, transposases, and 44 toxin-antitoxin systems.

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46 Conclusions

47	We identified a number of adaptations that may facilitate deep-sea radiation in members of the
48	genus Colwellia, as well as in other piezophilic bacteria. An enrichment in more basic and
49	hydrophobic amino acids could help piezophiles stabilize and limit water intrusion into proteins
50	as a result of high pressure. Variations in genes associated with the membrane, including those
51	involved in unsaturated fatty acid production and respiration, indicate that membrane-based
52	adaptations are critical for coping with high pressure. The presence of many piezophile-specific
53	genes near genomic islands highlights that adaptation to the deep ocean may be facilitated by
54	horizontal gene transfer through transposases or other mobile elements. Some of these genes are
55	amenable to further study in genetically tractable piezophilic and piezotolerant deep-sea
56	microorganisms.
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70 Background

71 The deep biosphere makes up one of the largest biomes on earth. An inherent 72 environmental parameter present throughout deep-oceanic and subsurface habitats is high 73 hydrostatic pressure. Elevated hydrostatic pressure influences many aspects of biochemistry and 74 requires adaptations throughout the cell (e.g. Somero, 1992). One well-studied adaptation is the 75 incorporation of unsaturated fatty acids into the membrane to combat physical changes such as 76 decreased fluidity (e.g. DeLong & Yayanos, 1985; DeLong & Yayanos, 1986; Allen et al., 77 1999). Additional membrane-associated adaptations are linked to porin-mediated nutrient 78 transport (Bartlett et al., 1989; Bartlett & Chi 1994), respiration (e.g. Yamada et al., 2000; Vezzi 79 et al., 2005; Xiong et al., 2016), and flagellar function (Eloe et al., 2008). Within the cell 80 changes in DNA replication, DNA structure, protein synthesis, and compatible solutes are also 81 important (El-Hajj et al., 2009; Martin et al., 2002; Lauro et al., 2008; Yancey et al., 2014). 82 Pressure-induced changes in transcription implicate additional functions (e.g. Campanaro et al., 83 2012; Michoud & Jebbar, 2016). Despite the fact that pressure exerts a profound influence on the 84 nature of life at depth, it is largely ignored in studies of deep-ocean biomes, and in marked 85 contrast to microbial adaptation to temperature or salinity, a robust description of biochemical 86 adaptation to high pressure is lacking.

87 Only a modest number of psychrophilic (cold-loving) and piezophilic (high-pressure 88 loving) species have been isolated to date, in large part due to the constraints imposed by 89 culturing under under *in situ*, high hydrostatic pressure conditions. However, metagenomic 90 sequencing of deep-ocean communities, and additional analyses of individual microbial 91 genomes, have provided insights. Metagenomic investigations have included locations within the 92 North Pacific subtropical gyre, the Mediterranean and the Puerto Rico Trench (DeLong *et al.*,

93 2006; Martin-Cuadrado et al., 2007; Konstantinidis et al., 2009; Eloe et al., 2011; Smedile et al., 94 2013). Genomic studies include those on Pseudoalteromonas (Qin et al., 2011), Alteromonas 95 (Ivars-Martinez et al., 2008), Shewanella (Wang et al., 2008; Aono et al., 2010), Photobacterium 96 (Campanaro et al., 2005; Vezzi et al., 2005; Lauro et al., 2014), SAR11 (Thrash et al., 2014), 97 and members of the *Thaumarchaeota* (Luo et al., 2014; Swan et al., 2014). One picture that has 98 emerged from the examinations at this level is that deep-sea microbes are enriched in mobile 99 elements, such as phage and transposases (DeLong et al., 2006; Ivars-Martinez et al., 2008; Eloe 100 et al., 2011; Qin et al., 2011; Lauro et al., 2013; Smedile et al., 2013; Léon-Zayas et al., 2015). 101 This has been attributed to the relaxation of purifying selection as an adaptive mechanism 102 (Konstantinidis et al., 2009), either to deep-ocean conditions or to the conditions found on 103 particles (Ganesh et al., 2014). Additional properties include an enrichment in heavy metal 104 resistance genes (Ivars-Martinez et al., 2008; Eloe et al., 2011; Qin et al., 2011; Smedile et al., 105 2013; Fontanez et al., 2015), the ability to use persistent dissolved organic material under 106 oligotrophic conditions (e.g. Martin-Cuadrado et al., 2007; Ivars-Martinez et al., 2008; Arrieta et 107 al., 2015; Landry et al., 2017), and widespread ability for chemoautotrophy (Swan et al., 2011; 108 Swan et al., 2014; Dyksma et al., 2016; Mußmann et al., 2017; Pachiadaki et al., 2017). The 109 small number of genome sequences of experimentally-confirmed deep-ocean piezophiles include 110 hyperthermophilic archaea (Pyrococcus and Thermoccus; Vannier et al., 2011; Jun et al., 2015; 111 Dalmasso et al., 2016), a thermophilic bacterium (Marinitoga; Lucas et al., 2012), a mesophilic 112 bacterium (Desulfovibrio; Pradel et al., 2013) and psychrophilic bacteria (Photobacterium, 113 Psychromonas, and Shewanella; Vezzi et al., 2005; Aono et al., 2010; Lauro et al., 2013a; Lauro 114 et al., 2013b; Zhang et al., 2019b). The genomic adaptations of these microorganisms to the deep 115 ocean or high hydrostatic pressure have not been fully explored (e.g. reviewed in Simonato et al.,

116 2006; Lauro et al., 2008; Oger & Jebbar, 2010; Peoples & Bartlett, 2017). Thus far the genome 117 characteristics of only one experimentally-confirmed obligately psychropiezophilic bacterial 118 species, Shewanella benthica (Lauro et al., 2013a; Zhang et al., 2019b), and one species of 119 obligately thermopiezophilic archaeon, Pyrococcus yayanosii (Jun et al., 2011), have been 120 described. 121 Most known psychropiezophilic strains belong to phylogenetically narrow lineages of 122 Gammaproteobacteria, including members of the Colwellia, Shewanella, Moritella, 123 Photobacterium, and Psychromonas (reviewed in Jebbar et al., 2015; Nogi et al., 2017). The 124 genus *Colwellia* contains some of the most psychrophilic and piezophilic species currently 125 known. Members of this genus are heterotrophic and facultatively anaerobic (Bowman et al., 126 2014). This genus has been of recent interest because of its association with the Deepwater 127 Horizon oil spill, where members of the *Colwellia* became some of the most abundant taxa 128 present because of their ability to degrade hydrocarbons (Redmond & Valentine, 2012; Mason et 129 al., 2014; Kleindienst et al., 2015). Although Colwellia do not appear to be abundant members of 130 deep-ocean or hadal (typically < 1%; e.g. Eloe *et al.*, 2011; Tarn *et al.*, 2016; Peoples *et al.*, 131 2018) communities, they can become dominant members under mesocosm conditions 132 (Hoffmann et al., 2017; Boeuf et al., 2019; Peoples et al., 2019a). At least four piezophiles have 133 been successfully isolated and described from this genus. The first known obligate 134 psychropiezophile, originally designated *Colwellia* sp. MT41, was isolated from the amphipod 135 *Hirondellea gigas* from the Mariana Trench at a depth of 10,476 m (Yayanos *et al.*, 1981). Strain 136 MT41 shows optimum growth at 103 MPa and does not grow at a pressure below 35 MPa, 137 approximately the pressure at average ocean depths (Yayanos *et al.*, 1981; Yayanos, 1986; 138 DeLong et al., 1997). Recently, Colwellia marinimaniae MTCD1, the most piezophilic microbe

139 known to date, was isolated from an amphipod from the Mariana Trench (Kusube *et al.*, 2017). 140 This strain displays an optimum growth pressure of 120 MPa and a growth range from 80 to 140 141 MPa, higher than the pressure found at full ocean depth. Based on 16S rRNA gene similarity 142 both strains MT41 and MTCD1 strains were determined to belong to the species Colwellia 143 marinimaniae (Kusube et al., 2017). Other psychropiezophiles within the genus include C. 144 hadaliensis (Deming et al., 1988) and C. piezophila (Nogi et al., 2004), isolated from the Puerto 145 Rico and Japan trenches, respectively. While the growth characteristics and fatty acid profiles of 146 these piezophilic species of *Colwellia* have been reported, other adaptations of these strains for 147 dealing with high hydrostatic pressure and deep-ocean environmental conditions have not been 148 investigated in great detail. 149 In this study, we compared the genomes of members of the *Colwellia* to identify 150 attributes that confer adaptation to the deep ocean. We report the genome sequences of three 151 obligately piezophilic Colwellia; Colwellia marinimaniae MT41, C. marinimaniae MTCD1, and 152 a new isolate obtained from sediment in the Tonga Trench, *Colwellia* sp. TT2012. We compared 153 these genomes, along with the publicly-available genome of C. piezophila ATCC BAA-637 154 (isolated as strain Y223G; Nogi et al., 2004), against three piezosensitive strains of C. 155 *psychrerythraea*. The piezosensitive strains include the most well-studied member of the 156 Colwellia, C. psychrerythraea 34H, a psychrophile isolated from Arctic ocean sediments 157 (Huston *et al.*, 2000) whose adaptations to low temperature have been investigated at multiple 158 levels (e.g. Marx et al., 2009; Showalter & Deming, 2018), including with genomics (Methé et 159 al., 2005). The two other comparison strains are C. psychrerythraea GAB14E and ND2E, 160 obtained from the Great Australian Bight at a depth of 1472 m and the Mediterranean Sea from 161 495 m, respectively (Figure 1A; Techtmann et al., 2016). While the C. psychrerythraea strains

162	share 99% identical 16S rRNA sequences, they have very divergent average nucleotide identities
163	(ANI; Techtmann et al., 2016). Because low temperatures and high pressures have similar effects
164	on biochemical processes, these three microbes were selected as comparison strains because they
165	all show growth at low temperatures, reducing the impact of temperature as a confounding
166	factor. Through the comparison of these seven strains depth and pressure-associated shifts were
167	identified in protein amino acid distributions and isoelectric points, as well as in gene
168	abundances, including the discovery of piezophile-specific genes.
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170	Results
171	General characteristics
172	We first evaluated the influence of high hydrostatic pressure on the growth of the seven
173	strains of Colwellia. The growth characteristics of Colwellia marinimaniae MT41, C.
174	marinimaniae MTCD1, and C. piezophila have been previously reported, showing growth
175	optima at 103 MPa (Yayanos, 1986; DeLong et al., 1997), 120 MPa (Kusube et al., 2017), and
176	60 MPa (Nogi et al., 2004), respectively. Colwellia sp. TT2012 is obligately piezophilic,
177	showing growth at 84 and 96 MPa but not at atmospheric pressure. We report the optimum
178	growth pressure in this manuscript as 84 MPa because strain TT2012 was not able to be revived
179	after repeated cryopreservation for growth rate analyses at lower or higher pressures. The three
180	C. psychrerythraea strains displayed different growth patterns from one another, but similarly
181	showed no growth at a pressure of 40 MPa after 10 days regardless of temperature (4°C or 16°C;
182	Supplementary Figure 1). Based on these growth characteristics, we classified the microbes as
183	either piezophilic (C. marinimaniae MT41, C. marinimaniae MTCD1, Colwellia sp. TT2012,

184 and C. piezophila) or piezosensitive (C. psychrerythraea strains 34H, GAB14E, and ND2E). 185 These terms are used to describe these groupings for the remainder of the manuscript. 186 To identify genomic attributes that facilitate growth at high pressure in the deep sea, we 187 compared the genomes of the piezophilic and piezosensitive strains (Table 1). We report here for 188 the first time the genome sequences of Colwellia marinimaniae MT41, C. marinimaniae 189 MTCD1, and *Colwellia* sp. TT2012. The remaining genomes are either publicly available (C. 190 *piezophila*, Kyrpides *et al.*, 2014) or have been previously reported (strain 34H, Methé *et al.*, 191 2005; strains ND2E and GAB14E, Techtmann *et al.*, 2016). The piezophiles are more closely 192 related to one another than to the piezosensitive strains based on a whole genome marker tree 193 and average nucleotide identity (Figure 1). This is also true when the strains are compared using 194 a ribosomal 16S RNA gene phylogenetic tree (Supplementary Figure 2). Colwellia marinimaniae 195 MT41, C. marinimaniae MTCD1, and Colwellia sp. TT2012 share approximately 96% 16S 196 rRNA gene sequence similarity and formed a monophyletic clade with an isolate from the 197 Kermadec Trench. Despite being isolated 34 years apart, strains MT41 and MTCD1 share 198 greater than 99% 16S rRNA gene sequence similarity and ANI. In contrast, the ANI of these 199 strains are only 95% similar to TT2012, indicating that TT2012 represents a distinct species. C. 200 piezophila does not appear to belong to this 16S rRNA gene tree piezophile-only monophyletic 201 clade (Supplementary Figure 2), although this relationship could not be confirmed with a whole 202 genome marker tree due to a lack of related genomes. Despite showing greater than 98% 16S 203 rRNA gene sequence similarity, the ANI of C. psychrerythraea strains 34H, GAB14E, and 204 ND2E is less than 90%, indicating that they have highly variable genome sequences. 205

206 GC content and amino acid features

207	We first compared general genomic attributes of the piezophilic and piezosensitive
208	strains, including genome size, GC content, isoelectric point, and amino acid distribution.
209	Genome sizes ranged between 4.3 and 5.7 Mbp in size (Table 1). The three piezophiles isolated
210	from the deepest depths (strains MT41, MTCD1, TT2012) have smaller genomes than the
211	piezosensitive strains (T-test, $p < .031$), but no correlation between genome size and optimum
212	growth pressure was found when considering C. piezophila and other members of the Colwellia
213	(Supplementary Figure 3). Coding density is lower in the piezophilic Colwellia. This is true even
214	when including all sequenced members of the <i>Colwellia</i> (T-test, $p < .01$). GC content ranged
215	between \sim 38 and 39%, with slightly higher GC present in the piezophiles. This is also true when
216	compared with other Colwellia strains with the exception of C. chukchiensis (Supplementary
217	Figure 3; T-test, $p < .08$). However, when examining members of the genera <i>Colwellia</i> ,
218	Psychromonas, and Shewanella, no correlation was apparent between % GC and growth
219	pressure. No correlation was found between optimum growth pressure and % GC within full
220	length 16S rRNA genes in the Colwellia.
221	Next, we evaluated the isoelectric point distributions of the Colwellia. Both piezophilic
222	and piezosensitive strains show a similar bimodal distribution of protein isoelectric points.
223	However, the piezophiles have a higher number of basic proteins (Figure 2; T-test, $p < .01$). This
224	shift is also seen when comparing within a broader number of <i>Colwellia</i> (T-test, $p < .01$) with the
225	exception of C. chukchiensis (Supplementary Figure 4). Piezophilic strains within the genera
226	Psychromonas and Shewanella also show a higher number of basic proteins compared to their
227	piezosensitive counterparts (Supplementary Figure 4; T-test, Psychromonas, $p < .03$; T-test,
228	Shewanella, clade 3, $p < .34$), with obligate piezophiles such as <i>Shewanella benthica</i> KT99,
229	Psychromonas sp. CNPT3, and an uncultured Psychromonas single-amplified genome from a

230 hadal amphipod (Leon-Zayas et al., 2015) having dramatically more basic proteins. GC content 231 or optimum growth temperature does not appear to be responsible for this shift in pI bias, even 232 when taking into account within-genus phylogenetic clades (Supplementary Figure 4, 233 Supplementary Figure 5). 234 Comparisons of amino acid abundances within conserved, orthologous proteins showed 235 that certain amino acids are more abundant in the piezophilic proteins when compared to those in 236 C. psychrerythraea 34H (Figure 2). Amino acids that are specifically enriched in the piezophiles 237 included tryptophan (W), tyrosine (Y), leucine (L), phenylalanine (F), histidine (H), and 238 methionine (M). In contrast, amino acids enriched in the piezosensitive strain included glutamic 239 acid (E), aspartic acid (D), asparagine (N), and serine (S). Specific amino acid asymmetrical 240 substitutions in which one amino acid consistently replaced another, including substitutions that 241 were also conserved in comparisons within members of the Shewanella, from piezosensitive to 242 piezophilic amino acid were: glutamic acid \rightarrow alanine (A), proline (P) \rightarrow alanine, threonine (T) 243 \rightarrow isoleucine (I), valine (V) \rightarrow isoleucine (I), glutamic acid \rightarrow lysine (K), asparagine (N) \rightarrow 244 lysine, glutamic acid \rightarrow glutamine (Q; Figure 2). Further asymmetrical substitutions specific to 245 the genus *Colwellia* include, from non-piezophile to piezophile, aspartic acid \rightarrow alanine, glycine (G) \rightarrow alanine, serine \rightarrow alanine, asparagine \rightarrow histidine, valine \rightarrow leucine, and glutamic acid \rightarrow 246 247 threonine.

248

249 *Gene differences*

We compared the predicted gene complements of the piezophilic and piezosensitive strains. When comparing relative abundances of clusters of orthologous genes (COGs; Figure 3), piezophilic *Colwellia* have a higher percentage of genes for replication/recombination/repair 253 (Category L), cell wall/membrane biogenesis (Category M), cell motility (Category N), 254 extracellular structures (Category W), and translation and ribosomal structure (Category J). The 255 piezosensitive strains have higher percentages of genes for transcription (Category K), secondary 256 metabolite biosynthesis/transport/metabolism (Category Q), and general function prediction 257 (Category R). Transposable elements are notably more abundant in the piezophiles, with the 258 exception of C. piezophila, having almost twice as many transposases as their piezosensitive 259 counterparts (Figure 3). Toxin-antitoxin genes are also enriched in the piezophiles, with 260 piezophilic strains containing 24-33 toxin-antitoxin genes while the piezosensitive Colwellia 261 have 9-18 copies. We found that strain MT41 and C. psychrerythraea 34H have approximately 262 11 and 9 genomic islands (GIs), respectively, as determined using Island Viewer (Bertelli *et al.*, 263 2017). We do not report the total number of GIs in the other strains because the fragmentation of 264 their genomes likely leads to GI misidentification. 265 All of the strains analyzed are heterotrophic. However, potential differences in carbon 266 metabolism exist (Figure 3). Genes for sarcosine oxidase (soxBDAG), which function in the 267 catabolism of glycine betaine in Colwellia (Collins & Deming, 2013), are present in 34H and 268 ND2E but not in the piezophiles. Transporters and permeases for putrescine are enriched in 34H 269 and GAB14E, strains where putrescine has been experimentally shown to be used as a sole 270 carbon source (Techtmann et al., 2016). In contrast, we identified genes involved in chitin 271 degradation, such as a chitin binding protein and chitinase (family 10 and 18), in the piezophiles 272 and GAB14E but not in the other piezosensitive strains. 273 Members of the *Colwellia* are facultative anaerobes capable of respiration and 274 fermentation. While all the *Colwellia* compared here use both the rnf (*rnfABCDGE*) and Na⁺-ngr

275 (*nqrABCDEF*) respiratory complexes, the NADH dehydrogenase I complex

276 (*nuoABCEFGHIJKLMN*) is only present in the three hadal piezophiles. These genes show 277 similarity to those in the piezophiles Shewanella benthica and S. violacea and to metagenomic 278 sequences from hadal sediments (Peoples, 2018). While all seven strains may have the capacity 279 for assimilatory nitrite (such as *nirBD*, *nasA*) and nitrate reduction (*napCBADFE*), genes for 280 dissimilatory nitrite reduction (*nirSCFNTB*) are only present in C. psychrerythraea strains 34H 281 and ND2E. The gene *nirK* is present in *C. piezophila*, although this strain was shown to reduce 282 nitrate but not nitrite (Nogi et al., 2004). The gene cluster for nitrous oxide reduction, 283 nosRZDFYL, is present in strains 34H, ND2E, and C. piezophila. This operon is flanked by 284 conserved regions found in the other strains, suggesting an insertion or deletion event. 285 Furthermore, the capacity for trimethylamine-N-oxide reduction using torSTRECAD is present in 286 strains 34H and ND2E but not in any of the piezophiles. 287 The seven strains of *Colwellia* compared are psychrophilic or psychrotolerant and have 288 adaptations to low temperatures. For example, all contain *pfaABCD* to produce polyunsaturated 289 fatty acids to counteract decreases in membrane fluidity because of low temperatures. In the case 290 of the deep-sea Colwellia this system will also optimize membrane phospholipid physical state at 291 high pressure. However, a number of genes involved in membrane adaptation are differentially 292 present among the two Colwellia groups. All piezophilic Colwellia have genes encoding delta-9 293 acyl-phospholipid desaturase, another enzyme promoting unsaturated fatty acid synthesis by 294 introducing double bonds directly into membrane phospholipid saturated fatty acids. In contrast, 295 a fatty acid cis/trans isomerase that alters the ratio of cis- and trans- phospholipids by 296 isomerizing -cis to -trans double bonds, is encoded within all piezosensitive *Colwellia* but is 297 notably absent in the piezophilic *Colwellia*. Furthermore, the piezophilic strains encode almost 298 twice as many glycosyltranferases, enzymes involved in extracellular polysaccharide synthesis.

299 Stress-response genes are also differentially present in the genomes.

300 Deoxyribopyrimidine photolyase (DNA photolyase; *phrB*), which is involved in repairing DNA 301 damaged by ultraviolet light, is found in strains 34H and ND2E but notably absent in all 302 piezophilic Colwellia. Both piezophilic and piezosensitive strains contain superoxide dismutase 303 and catalase for responding to oxidative stress. The genes araC and lvsR, whose products control 304 the expression of a variety of stress response systems, are more abundant in the piezosensitive 305 Colwellia. The piezophilic Colwellia are distinct in having multicopper oxidases and copper 306 chaperones for coping with heavy metal damage and maintaining copper homeostasis. 307 Phenotypic analysis of the *Colwellia* showed that the piezophiles appear more resistant to copper 308 exposure compared to their non-piezophilic counterparts (Supplementary Figure 6). Some of the 309 genes which putatively confer heavy metal resistance are similar to other piezophiles and are 310 located near genomic islands or other horizontally transferred elements, consistent with the 311 hypothesis that heavy metal genes can be horizontally transferred (e.g. Orellana & Jerez, 2011; 312 Navarro et al., 2013; Chen et al., 2017). 313 We identified other unique genes that differ not only between Colwellia strains but show 314 biased distributions towards additional piezophilic microbes and deep-ocean metagenomic 315 datasets (Table 2; Dombrowski et al., 2018; Hu et al. 2018; Tully et al., 2018; Peoples, 2018). 316 For example, a putative archaeal S-adenosyl-l-methionine (SAM) dependent methyltransferase 317 (pfam13659) is present in the piezophiles and strain GAB14E. This protein is similar to those 318 present in bacterial and archaeal piezophiles, including members of the genera Colwellia, 319 Shewanella, Moritella, Psychromonas, Methanocaldococcus, Thermococcus, and Pyrococcus. 320 The related methyltransferase isolated from *Pyrococcus abysii* (39% similar to MT41 protein) 321 functions in tRNA modification (Guelorget et al., 2010). Piezophilic Colwellia have two copies

322	of d-alanine-d-alanine ligase (pfam07478), a gene which may be involved in peptidoglycan
323	synthesis. Unlike the situation in piezophilic Shewanella (Zhang et al., 2019b), this gene is not
324	present near flagellar assembly components. While all strains have operons for a Type II
325	secretion system and a Type IV pilus, a <i>tad</i> pilus involved in adhesion is found only in the
326	piezophiles and related to that in Shewanella violacea. This operon is also found in GAB14E;
327	however, this strain lacks a number of putative tadE-like genes that are present in the piezophile
328	operons. Two alanine dehydrogenases are also present in the piezophilic strains while only one is
329	present in the piezosensitive members. The piezophile-specific dehydrogenase (pfam05222) is
330	thought to catalyze the NAD-dependent reversible amination of pyruvate to alanine. It is similar
331	to a dehydrogenase present in other piezophilic species, including Shewanella benthica,
332	Moritella yayanosii, Photobacterium profundum SS9, and binned genomes from a deep subsea
333	aquifer (Tully et al., 2018) and trench sediments (Peoples, 2018).
334	A number of the genes specific to piezophiles are present near one another, rather than
335	individually spread throughout the genome (Table 2). Many of these genes are near variable
336	regions containing genomic islands, phage genes, transposases, and toxin-antitoxin system genes
337	(Supplementary Figure 7). For example, the d-alanine-d-alanine ligase in strain MT41 is next to
338	two putative genomic island regions, one of which is different than that present in strain TT2012
339	(Figure 4). Because genomic islands are identified based on nucleotide bias across the genome
340	and the Colwellia sp. TT2012 genome is fragmented into short contigs, the lack of predicted
341	genomic islands does not preclude their presence. In the piezophile Moritella yayanosii this gene
342	is near a gene encoding a predicted phage integrase protein, while in Shewanella benthica KT99
343	it is present in a flagellar operon that also contains a transposase embedded within it. Similarly,
344	the piezophile-specific alanine dehydrogenase is present near a number of phage and

348 not specific to piezophiles, display low similarity to members of the genus *Vibrio*. The similarity

349 of variable genes within *Colwellia* to species of *Vibrio* has been previously noted (Collins &

350 Deming, 2013). Horizontal gene transfer has been shown to be important in the evolution of

351 *Vibrio* species (Faruque & Mekalanos, 2013).

352

353 Discussion

354 In this study we compared the genomes of members of piezophilic *Colwellia* – including 355 the most high pressure-adapted species known to date – with their piezosensitive counterparts to 356 search for features that could confer adaptation to the deep sea. These microbes were isolated 357 from surface and bathyal waters to abyssal and hadal depths. Both 16S rRNA gene sequence-358 based phylogenetic analyses and phylogenomic analyses indicate that the piezophilic Colwellia 359 are closely related. While the piezophiles appear to form a single cluster based on the 360 phylogenomic tree, in the 16S rRNA gene phylogenetic tree C. piezophila appears basal to not 361 only the piezophiles but also a clade that includes piezosensitive lineages. Therefore, it is 362 possible that piezophily has evolved multiple times within the *Colwellia*. Further whole genome 363 sequencing will be needed to determine if all piezophilic *Colwellia* form a single clade 364 independent from other piezosensitive microbes as has been reported for Shewanella (Aono et 365 al., 2010). Piezophilic Colwellia have now been isolated from five different trenches, including 366 the Mariana (strains MT41, MTCD1), Puerto Rico (C. hadaliensis), Japan (C. piezophila), Tonga 367 (strain TT2012), and Kermadec (Bartlett laboratory unpublished; Lauro et al., 2007; Peoples et

al., 2019a). Piezophilic members of the genus *Colwellia* are therefore widespread within deepocean and hadal environments.

370	While the piezophiles have lower coding density than their non-piezophilic counterparts,
371	no correlation was found between genome size and optimum pressure of growth. This is in
372	contrast to comparisons between shallow and deep pelagic datasets showing that deeper lineages
373	appear to have larger genomes (e.g. Konstantinidis et al., 2009; Beszteri et al., 2010; Eloe et al.,
374	2011; Thrash et al., 2014). Instead, the three piezophiles with the deepest collection depths
375	represented some of the smallest Colwellia genomes examined. One possibility is that these
376	differences reflect different selective pressures operating within seawater, sediments and
377	amphipods. It is remarkable that strain MT41 and MTCD1, two piezophiles isolated from
378	amphipod material in the Mariana Trench 34 years apart, share over 99% ANI. Perhaps this
379	reflects strong selection for a particular Colwellia strain within the microbiome of Mariana
380	Trench Hirondellea gigas amphipods, such as that seen within symbionts of deep-sea
381	anglerfishes (Baker et al., 2019). Their consistent isolation from amphipods (e.g. Yayanos et al.,
382	1981; Kusube et al., 2017) suggests that some members can be associated with hosts, and host-
383	microbe relationships can lead to genome streamlining and smaller genome sizes (McCutcheon
384	& Moran, 2012). Nearly all known piezophilic genera have been found in conjunction with hosts
385	(e.g. Nakayama et al., 2005) and the microbial activity of the gut contents of deep-sea animals
386	shows high levels of piezophily (Tabor et al., 1982). However, the genus Colwellia is not present
387	in recognizable abundances within hadal amphipod metagenomes (Zhang et al., 2019a), their
388	high % GC is not indicative of an endosymbiont (McCutcheon & Moran, 2012), and the obligate
389	piezophile Colwellia sp. TT2012 was isolated from sediments rather than amphipods. An
390	alternative hypothesis is that Colwellia may be undergoing genome reduction because of a

specialized lifestyle within the deep sea, as hypothesized for some psychrophiles within sea ice (Feng *et al.*, 2014). Members of this genus may instead be isolated in conjunction with amphipods because of their ability to degrade nutrient-rich decaying amphipod material, for example using genes for chitin degradation. *Colwellia* may also be ingested by amphipods as a byproduct of the feeding of these deep-sea scavenging macrofauna because of the preference of *Colwellia* for nutrient-rich particulate organic material (Hoffmann *et al.*, 2017; Peoples *et al.*, 2018; Boeuf *et al.*, 2019).

398 The isoelectric point (pI) distribution of proteins within a proteome can correlate with the 399 ecological niche of an organism (Kiraga et al., 2007). Here we found that piezophilic Colwellia 400 have a more basic proteome than their piezosensitive counterparts. This pattern is conserved in 401 comparisons between piezophilic and piezosensitive members of the genera Shewanella and 402 *Psychromonas*, indicating it is a property that may be widespread amongst piezophiles within the 403 Gammaproteobacteria. Although intracellular microorganisms also have more basic proteomes 404 than free-living species (Kiraga et al., 2007), this is associated with an AT base pair enrichment 405 not present in the piezophilic Colwellia. A basic proteome may be the result of the accumulation 406 of mutations (Kiraga et al., 2007), consistent with the low coding density and high numbers of 407 transposable elements within the piezophiles. Alternatively it could arise to help with charge 408 balance within the cytoplasm, analogous to the role of the more acidic proteome of haloarchaea, 409 which counters the high intracellular potassium ion levels present at high osmotic pressures (Paul 410 et al., 2008). The intracellular inorganic and organic solute levels within piezophiles are not well 411 known, but could be important to the maintenance of macromolecule function at high pressure 412 (Martin et al., 2002; Yancey et al., 2001; Yancey, 2005). Among orthologous proteins 413 piezophiles are also enriched in hydrophobic residues, including tryptophan, tyrosine, leucine,

414 phenylalanine, histidine, and methionine. This finding has been noted in metagenomes from 415 4,000 m (Konstantinidis et al., 2009) and may be important in maintaining protein structure 416 against water penetration at high pressure (Hummer et al., 1998; Somero, 2003). Specific amino 417 acid substitutions where one amino acid is consistently replaced by another indicate that small 418 nonpolar compounds (alanine, isoleucine), amine-containing polar compound (glutamine), and a 419 positively charged basic compound (lysine) are selected for in piezophiles, while negatively 420 charged acidic compounds (glutamate), polar compounds (threonine, asparagine), and non-polar 421 compounds (valine, proline) are selected against. Similar shifts were also seen in Desulfovibrio 422 piezophilus (Pradel et al., 2013), although different amino acids were preferentially abundant in 423 piezothermophilic archaea (Di Giulio, 2005). 424 We identified a number of gene abundance characteristics that could confer adaptation to 425 the deep ocean. Enrichments in COG J (translation), L (replication and repair), M (cell 426 wall/membrane biogenesis), and N (cell motility) appear enriched in the piezophiles. An 427 enrichment of category M and L has previously been observed within deep ecotypes of 428 Alteromonas (Ivars-Martinez et al., 2008). The enrichment within the piezophiles of COG M is 429 in part due to higher abundances of glycosyltransferases, which appear to correlate with depth 430 within metagenome datasets (DeLong et al., 2006). Glycosyltransferases have been predicted to 431 contribute to low temperature-adaptation (Methé et al., 2005) and could be more abundant in the 432 psychropiezophiles because they are more stenothermic. In contrast, a fatty acid cis/trans 433 isomerase was present only in the piezosensitive strains. The rapid cis-to-trans isomerization of 434 unsaturated fatty acids via this isomerase has been observed in *Pseudomonas putida* P8 in 435 response to changes in temperature and salinity (Loffeld & Keweloh, 1996; Holtwick et al., 436 1997). Furthermore, the COG category for transcription (K) is significantly enriched in non-

437 piezophiles compared to piezophiles. This is in part due to an enrichment in the transcription 438 factors AraC and LysR, which have a wide variety of regulatory functions including carbon 439 metabolism and stress response (Gallegos et al., 1997; Maddocks & Oyston, 2008). The 440 enrichment of COG category K in shallow-water organisms has been observed in the surface-441 water ecotype of Alteromonas macleodii (Ivars-Martinez et al., 2008). These findings could 442 reflect the adaptation of non-piezophilic shallow-water microbes to a more dynamic 443 environment, such as rapid salinity or temperature shifts associated with sea-ice or surface 444 seawater. In contrast, autochthonous, obligate deep-ocean microbes would not be expected to 445 experience similar rates or magnitudes of these changes. 446 Other specific genes biased towards piezophiles within COG M include delta-9 acyl-447 phospholipid desaturase and a CDP-alcohol phosphatidyltransferase. While the desaturase is 448 upregulated at high pressure in *Photobacterium profundum* SS9 (Campanaro et al., 2005), this 449 gene is present in other non-piezophilic strains of the *Colwellia* not examined here, indicating it 450 may not be pressure-specific. An extra copy of d-alanine-d-alanine ligase is present in the 451 piezophiles and may function in peptidoglycan biosynthesis. While this gene was reported within a flagellar operon in Shewanella benthica (Zhang et al., 2019b), in strain MT41 it is present next 452 453 to a putative genomic island (Figure 5). The non-piezophile-specific copy of d-alanine-d-alanine 454 ligase is upregulated in the proteome of strain 34H after incubation at -1°C (Nunn et al., 2015), 455 perhaps reflecting a role in low temperature acclamation. Overall, the enrichment in piezophiles 456 of genes involved in COG category M is consistent with a wealth of experimental evidence 457 demonstrating that changes in membrane structure are critical for adapting to high hydrostatic 458 pressure. Unsaturated fatty acids help maintain membrane structure under high pressure (Chi and 459 Bartlett, 1995; Yano et al., 1998; Allen et al., 1999; Usui et al., 2012; Abe, 2013), with strain

460 MT41 able to produce more than 15% of its total membrane fatty acids as docosahexaenoic acid461 (22:6; Delong & Yayanos, 1986).

462	Another adaptation associated with the membrane involves energetics and respiration.
463	We identified an additional NADH ubiquinone oxidoreductase (nuo) gene cluster in a number of
464	piezophiles. This unique NADH dehydrogenase, which translocates four protons per two
465	electrons (Pinchuk et al., 2010), may help with energy acquisition under in situ, high pressure
466	conditions. We also identified an alanine dehydrogenase specific to the piezophiles that may
467	function in the reversible amination of pyruvate to alanine coupled with the oxidation of NADH
468	to NAD ⁺ . This may act as an adaptive strategy under inhibited respiratory conditions by
469	maintaining NADH/NAD ⁺ homeostasis (Jeong & Oh, 2019), such as during shifts to anoxic
470	conditions (Hutter & Dick, 1998; Feng et al., 2002) or after exposure to physical stressors
471	impeding electron flow. Alanine dehydrogenases in Listeria are insensitive to inactivation up to
472	pressures of 550 MPa (Simpson & Gilmour, 1997), transcriptionally upregulated in
473	Desulfovibrio piezophilus at high pressure (Pradel et al., 2013), and abundant in the proteomes of
474	strain 34H at sub-zero temperatures (Nunn et al., 2015). We speculate that the piezophilic
475	alanine dehydrogenase functions in NADH/NAD ⁺ homeostasis under high hydrostatic pressure
476	conditions. In contrast, we found that TMAO reductase (torECAD), which reduces TMAO to
477	TMA, was not present in any of the piezophilic Colwellia. A similar finding has been noted in
478	genomes of Psychromonas from the guts of hadal amphipods, where the lack of TMAO
479	reductase was attributed to the preferential need for TMAO as a piezolyte in the host amphipod
480	over its use as an electron acceptor by the microbe (Zhang et al., 2018). An alternative
481	hypothesis is that TMAO is used by microbial piezophiles as a piezolyte as it is in deep-sea
482	metazoans (Yancey et al., 2001; Yancey et al., 2014). Finding differences in respiratory

483	capacity within piezophiles is not unexpected. Others have previously noted the influence of
484	collection depth and pressure on the presence and regulation of respiratory membrane-bound
485	cytochrome c oxidases and hydrogenases (Yamada et al., 2000; Vezzi et al., 2005; Chikuma et
486	al., 2007; Tamegai et al., 2013; Leon-Zayas et al., 2015; Vannier et al., 2015; Michoud &
487	Jebbar, 2016; Xiong et al., 2016; Zhang et al., 2018). These changes could stem directly from
488	pressure influences or from a greater reliance on the colonization of reduced oxygen niches
489	associated with particles or animals (Boeuf et al., 2019; Peoples et al., 2019a). This latter
490	possibility could be facilitated by the tad pilus present in the piezophilic Colwellia (Planet et al,
491	2003; Tomich et al., 2007; Pu et al., 2018).
492	Horizontal gene transfer (HGT) can provide genetic material that enhances fitness in new
493	environments. An experimental demonstration of this impact is the introduction of a DNA
494	photolyase gene, missing in piezophilic Colwellia and other deep-sea species (Delong et al.,
495	2006; Lauro & Bartlett, 2008; Konstantinidis et al., 2009; Peoples et al., 2019b), into the
496	piezophile Photobacterium profundum SS9 to generate a UV resistant phenotype (Lauro et al.,
497	2014). It is striking that many of the Colwellia genes most similar to those in other piezophiles
498	appear in clusters within variable regions that include genomic islands, putative phage genes,
499	transposases, and toxin-antitoxin systems. Despite their smaller genome sizes, laterally
500	transferred elements such as transposase and toxin-antitoxin genes are more abundant in the
501	piezophilic Colwellia examined here, consistent with their lower coding densities. Another
502	notable feature of these variable regions is that they differ even between closely-related strains,
503	such as between Colwellia marinimaniae MT41 and C. marinimaniae MTCD1.
504	Mobile genetic elements have been suggested to confer adaptations to extreme conditions
505	(e.g. Anderson et al., 2011; Pradel et al., 2013; Feng et al., 2014; Lossouarn et al., 2015; Mao &

506	Grogan, 2017), such as in the known piezophile Photobacterium profundum SS9 (Campanaro et
507	al., 2005). Deep-sea specific toxin-antitoxin systems have been identified in members of the
508	Shewanella (Zhang et al., 2019b) and have been shown to influence the growth of Pyrococcus
509	yayanosii at different pressures (Li et al., 2016; Li et al., 2018). Mobile genetic elements may
510	provide new metabolisms within strains of Colwellia psychrerythraea, including the transfer of
511	sox genes involved in sarcosine metabolism (Collins & Deming, 2013; Techtmann et al., 2016).
512	Because of the similarity of many genomic island-associated genes in members of the
513	piezophilic Colwellia to those in other gammaproteobacterial piezophiles, we suggest that HGT
514	is a significant evolutionary process governing high pressure adaptation. Future studies should
515	evaluate these regions and their associated genes for their importance in piezophily.
516	
517	Conclusions
518	In this study we compared the genomes of piezophilic and piezosensitive Colwellia to
519	
517	identify adaptations to extreme deep ocean conditions. Differences in amino acid composition,
520	identify adaptations to extreme deep ocean conditions. Differences in amino acid composition, membrane and cell wall structure, respiratory capacity, tRNA modification, and complex organic
520	membrane and cell wall structure, respiratory capacity, tRNA modification, and complex organic
520 521	membrane and cell wall structure, respiratory capacity, tRNA modification, and complex organic carbon utilization appear to be important for life at hadal depths. Many piezophile-enriched
520 521 522	membrane and cell wall structure, respiratory capacity, tRNA modification, and complex organic carbon utilization appear to be important for life at hadal depths. Many piezophile-enriched genes are located near areas of genomic variability and could be shared among piezophiles by
520521522523	membrane and cell wall structure, respiratory capacity, tRNA modification, and complex organic carbon utilization appear to be important for life at hadal depths. Many piezophile-enriched genes are located near areas of genomic variability and could be shared among piezophiles by horizontal gene transfer. Some of the adaptations identified may not be for high pressure
 520 521 522 523 524 	membrane and cell wall structure, respiratory capacity, tRNA modification, and complex organic carbon utilization appear to be important for life at hadal depths. Many piezophile-enriched genes are located near areas of genomic variability and could be shared among piezophiles by horizontal gene transfer. Some of the adaptations identified may not be for high pressure adaptation per se, but for lifestyles favored in hadal trenches such as affiliation with particulate

528

529 Sample collection and high-pressure cultivation conditions

530	Colwellia sp. TT2012 was isolated from sediments collected via gravity core in the
531	Tonga Trench (16° 38.505' S, 172° 12.001' W) at a depth of 9161 m on September 2, 2012
532	aboard the R/V Roger Revelle. Sediment from the upper three cm sediment depth horizon was
533	mixed with filter-sterilized trench seawater and maintained at a pressure of 84 MPa and 4°C. A
534	subset of this material was inoculated into ZoBell 2216 Marine Medium (BD Difco, Thermo
535	Fisher, Waltham, MA, USA) under the same pressure and temperature conditions. Colwellia sp.
536	TT2012 was eventually isolated as a pure culture following a number of dilution to extinction
537	inoculations.
538	The isolation of both strains of Colwellia marinimaniae have been previously described.
539	Colwellia marinimaniae MTCD1was isolated from amphipods at a depth of 10,918 m in the
540	Challenger Deep (Kusube et al, 2017). Colwellia marinimaniae MT41 was also isolated from
541	amphipods at a depth of 10,476 m (Yayanos et al., 1981). Both strains were maintained in
542	pressurizable polyethylene transfer pipette bulbs (Samco Scientific, USA) with Zobell 2216
543	Marine Medium broth at 4°C and high pressure prior to sequencing.
544	
545	Pressure Sensitivity and Heavy Metal Sensitivity Testing
546	The growth of the strains was evaluated under different pressure and temperature
547	conditions. Cultures of Colwellia strains 34H, GAB14E, and ND2E were incubated in Zobell
548	2216 marine medium supplemented with 100 mM HEPES and 20 mM glucose at 4°C. Growth
549	under high hydrostatic pressure was evaluated by incubating cultures at 20 MPa increments
550	between 0.1-80 MPa at 4°C and 16°C in triplicate. The OD600 was measured every 2.5 days for
551	ten days. Growth rates of <i>Colwellia</i> sp. TT2012 were conducted at 0.1, 84, and 96 MPa at 4°C.

552 Copper sensitivity tests were also performed on the piezophilic (strains MT41, MTCD1, and 553 TT2012) and non-piezophilic *Colwellia* strains (strains 34H, GAB14E, ND2E). Copper (II) 554 chloride dihydrate in concentrations ranging from 0 - 1.5 mM in 0.3 mM increments were added 555 to inoculated 2216 media and the cultures were incubated at 4°C for 1-4 weeks with weekly 556 inspection. 557 558 Genome sequencing and assembly Genomic DNA from C. marinimaniae MTCD1 was extracted from 100 mL of liquid 559 560 culture after 4 weeks of incubation at 110 MPa. DNA was isolated using the Mo-Bio Ultraclean Microbial DNA Isolation Kit (Mo-Bio, USA). Genomic DNA was obtained from Colwellia sp. 561 562 TT2012 after growth at 84 MPa and 4°C for 3 weeks. Cells were filtered onto a 0.22 um 563 Millipore Sterivex filter cartridge (Fischer Scientific, USA) and first subjected to a lysis buffer 564 (50mM Tris-HCl at pH 8.3, 40mM EDTA at pH 8.0, 0.75 M sucrose) and R1804M Ready-Lyse 565 lysozyme solution (Illumina, USA). After 15 minutes of incubation at 37°C, proteinase K and 566 sodium dodecyl sulfate were added to a final concentration of 0.5mg/ml and 1% respectively. 567 The mixture was then incubated at 55°C for 25 minutes, followed by 70°C for 5 minutes. The 568 lysate was treated two times with phenol-chloroform-isoamyl alcohol (24:24:1) and 569 chloroform: isoamyl alcohol (24:1) and further purified using a Mo-Bio Utraclean DNA Isolation 570 Kit spin column. The genomes of C. marinimaniae and Colwellia sp. TT2012 were sequenced at 571 the Institute for Genomic Medicine (IGM) at UCSD using the MiSeq sequencing platform 572 (Illumina, San Diego). The raw forward and reverse reads were merged using FLASH version 573 1.2.10 (Magoč & Salzberg, 2011) and assembled with SPAdes version 3.1.0 (Bankevich et al., 574 2012).

575	The genome of strain MT41 was sequenced to closure by whole random shotgun
576	sequencing. Briefly, one small insert plasmid library (2–3 kb) and one medium insert plasmid
577	library (10-15 kb) were constructed by random nebulization and cloning of genomic DNA. The
578	sequences were assembled using the TIGR Assembler (Sutton et al., 1995). All sequence and
579	physical gaps were closed by editing the ends of sequence traces, primer walking on plasmid
580	clones, and combinatorial PCR followed by sequencing of the PCR product.
581	
582	Genomic completeness, phylogenetic analysis, and annotation
583	The genomes were evaluated for their completeness and phylogenetic relationships.

584 Genome completeness and contamination was estimated using CheckM (Parks et al., 2015). A

585 whole-genome phylogenetic tree was built using RAxML (Stamatakis *et al.*, 2014) on the

586 CIPRES science gateway (Miller et al., 2010) using the single-copy marker genes identified

587 within CheckM. Ribosomal 16S RNA gene trees were also built by aligning sequences using the

588 SINA Aligner (Pruesse *et al.*, 2012) and built using RAxML All trees were visualized using the

589 Interactive Tree of Life (Letunic & Bork, 2016). Genomes were annotated using the Integrated

590 Microbial Genomes pipeline (IMG/ER; Markowitz *et al.*, 2014). Pairwise average nucleotide

591 identity between the genomes was evaluated within both the IMG interface and with orthoANI

592 (Lee *et al.*, 2016).

593

594 *Comparative genomic analysis*

A comparative genomic analysis was performed between the piezophilic and nonpiezophilic strains of *Colwellia* to identify whole-genome changes and specific genes unique to piezophiles. The isoelectric point (pI) of each predicted proteome was calculated using the

598 compute pI/MW tool in the ExPASy Bioinformatics Resource Portal (Artimo *et al.*, 2012). 599 Isoelectric point values from ExPASy were rounded to the nearest tenth and the frequency of 600 each protein pI was plotted in Figure 2a as a percent of the total proteome. Each proteome was 601 divided into an acidic set of proteins (pI<7; N_a) and a basic set (pI>7; N_b) and the bias quantified 602 using the formula ((N_b-N_a)/(N_b+N_a) × 100). The pI bias percentage is calculated such that 100% 603 means the proteins in the entire proteome are basic, -100% means all the proteins are acidic, and 604 0% means equal percentage of basic and acidic proteins.

605 To identify specific amino acid substitutions that may correlate with piezophily, amino 606 acid asymmetry was calculated using the procedure and software described in McDonald et al. 607 (McDonald et al., 1999). First, proteins from the genomes were clustered using TribeMCL 608 (Enright et al., 2002; scripts available at https://github.com/juanu/MicroCompGenomics) with a 609 Blastp cutoff of 1e-5 and an inflation value of 1.4. Orthologous single-copy gene clusters present 610 in both the piezophiles and Colwellia psychrerythraea 34H were aligned using MAFFT (Katoh 611 & Toh, 2008) and then processed with the Asymmetry programs AmbiguityRemover (using a 612 value of 2 for the number of adjacent sites), AsymmetryCounter, and AsymmetryScaler (with 613 three decimal places and 100 replicates; McDonald et al., 1999). Approximately 346,000 aligned 614 amino acid sites were examined in each comparison. Comparisons were also performed between 615 the Shewanella strains S. benthica KT99, S. violacea DSS12, and S. piezotolerans WP3 against 616 the piezosensitive S. sediminis EB3.

617 Protein abundances from the genomes were compared to identify attributes preferentially 618 enriched in either the piezophiles or piezosensitive strains. General COG category distributions 619 were evaluated using IMG/ER annotations. For the identification of differentially-abundant 620 specific proteins, protein clusters were generated using the TribeMCL analysis as described above. These identified protein clusters were further screened using blastp (Altschul *et al.*, 1990)
against the nr database for their prevalence in other *Colwellia* genomes, other piezophile
genomes, or other metagenomes. This manual curation allowed for the identification of both
genes differentially abundant within the groups of genomes immediately discussed here but also
allowed for a culled, smaller dataset of genes that may be present in other deep-ocean isolates
and datasets.

627 Certain genomic features within the genomes were also identified. Genomic islands were 628 identified using IslandViewer (Bertelli et al., 2017). Regions that may represent genomic islands 629 were also identified using the Mean Shift Genomic Island Predictor (MSGIP; de Brito et al., 630 2016). As incomplete genomes appeared to give spurious results, the total number of genomic 631 islands are reported only for the complete genomes of *Colwellia marinimaniae* MT41 and *C*. 632 psychrerythraea 34H. However, genomic islands for some of the partial genomes are shown here 633 (e.g. Figure 4, Figure 5) only when IslandViewer or MSGIP identified a region as a genomic 634 island, it was in a similar region as a genomic island found in either of the 34H or MT41 635 genomes, and it appeared to be a region of variability based on IMG/ER annotations. The 636 homology of these variable regions was analyzed using blastn and visualized with the R package 637 genoPlotR (Guy et al., 2010) and Kablammo (Wintersinger et al., 2015). Putative transposases 638 and toxin/antitoxin genes were identified based on IMG/ER annotations. Putative viral regions of 639 each genome were also identified based predominantly on IMG/ER annotations with a functional 640 search using the terms 'phage' and 'virus,' but also with VirFinder (Ren et al., 2017) and 641 VirSorter (Roux et al., 2015). Different types of flagella and pili were annotated using 642 MacSyFinder and TXSScan (Abby et al., 2016; https://galaxy.pasteur.fr/#forms::txsscan) with

643	default parameters. Carbohydrate-active enzymes within each genome were identified using
644	dbCAN (Yin et al., 2012).
645	
646	Declarations
647	Ethics approval and consent to participate
648	Not applicable.
649	
650	Consent for publication
651	Not applicable.
652	
653	Availability of data and material
654	The genome sequences of strains MT41, MTCD1, and TT2012 have been deposited at
655	GenBank under the accessions CP013145, GCA_001432325, and GCA_001440345,
656	respectively. The assembled and annotated genomes of strains MT41, MTCD1, and TT2012 can
657	be located in IMG/JGI under the IMG taxon IDs 2501651205, 2585427605, and 2585428047
658	respectively.
659	
660	Competing interests
661	The authors declare that they have no competing interests.
662	
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670	
671	Authors' contributions
672	LMP, TSK, JAU, KM, RAC, AAY, BAM performed experimental and bioinformatics
673	work. LMP, TSK, DHB performed data analysis. LMP, TSK, DHB wrote the manuscript. All
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675	
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- obligate piezophilic bacterium isolated from the deepest Mariana Trench sediment. Mar Gen. 2019b;44:52-56.

1287 Table 1. Genome characteristics of strains of *Colwellia* compared in this study.

1207	
1288	

GC Coding region Predicted genes Protein coding genes with function prediction	39.34% 83.68% 3895 2826	39.40% 83.88% 4057 2933	39.55% 83.00% 4071 2897	38.84% 83.65% 4598 3362	37.97% 85.76% 4790 3484	38.08% 85.73% 4479 3379	-
Contamination	1.47%	0.73%	2.61%	1.01%	0.68%	2.38%	
<u>Completeness</u>	100%	100%	99.33%	100%	99.49%	100%	
Isolation location Isolation source Genome size (Mb) DNA scaffold count Completeness Contamination	184	1	250	38	77	57	
Genome size (Mb)	4.37	4.34	4.44	5.48	5.72	5.15	
Isolation source	Amphipod	Amphipod	Sediment	Sediment	Water	Water	
Isolation depth	10918 m	10476 m	9161 m	6278 m	1472 m	495 m	
Isolation location	Mariana Trench	Mariana Trench	Tonga Trench	Japan Trench	GAB14E Great Australian Bight	Mediterranean Sea	
Strain	MTCD1	<u>MT41</u>	TT2012	C. piezophila	GAB14E	ND2E	

Table 2. Genes identified in piezophilic Colwellia but not the piezosensitive strains and which

show a biased presence within other known piezophilic microbes and deep-ocean datasets.

IMG Gene ID MT41	Start MT41 (bp)	End MT41 (bp)	<u>Similar to:</u>
2501712773-2501712774	738561	741622	P. hadalis , S. benthica , S. violacea , M. yayanosii , Moritella sp. PE36, Peoples 2018, Hu et al . 2018, Tully et al . 2018
2501712781	748798	749364	S. benthica, Tully et al. 2018, Dombrowski et al. 2018
2501712785	751307	751420	P. hadalis , Peoples 2018
2501713024-2501713025	1002524	1003568	M. yayanosii , Moritella sp. PE36, Tully et al. 2018, Dombrowski et al. 2018
2501713028-2501713043	1004921	1020893	P. hadalis , S. benthica , S. violacea , M. yayanosii , Moritella sp. PE36, Hu et al. 2018, Tully et al. 2018
2501713628	1635614	1636453	P. hadalis, S. benthica, S. violacea, M. yayanosii, Moritella sp. PE36, piezophilic archaea, Hu et al. 2018, Tully et al. 2018
2501713976	1995082	1995321	S. benthica
2501714033	2052442	2052666	S. benthica
2501714084	2101280	2101915	P. hadalis , S. benthica , S. violacea , M. yayanosii , Moritella sp. PE36, Tully et al. 2018
2501714124-2501714126	2137413	2141565	P. hadalis, S. benthica, M. yayanosii, Tully et al. 2018, Dombrowski et al. 2018
2501714471-2501714485	2514635	2530350	S. benthica , S. violacea , Peoples 2018
2501714619	2663589	2663918	S. benthica , Peoples 2018, Tully et al . 2018
2501714669	2714988	2715770	M. yayanosii, Moritella sp. PE36, SAR324, Peoples 2018, Tully et al. 2018
2501715698	3869630	3871057	Photobacterium profundum SS9, S. benthica , M. yayanosii , Moritella sp. PE36, Peoples 2018, Tully et al. 2018
2501715722	3894109	3895707	P. hadalis , Tully et al. 2018
2501715931-2501715932	4122279	4122819	S. benthica , M. yayanosii
2501716002-2501716003	4182966	4183371	S. benthica , M. yayanosii , Tully et al. 2018

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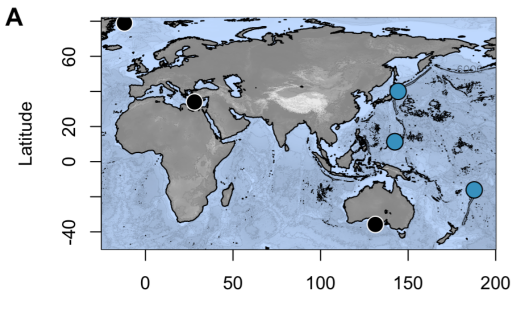
1316

Figure 1. A; Approximate sample collection locations for the *Colwellia* strains compared in this study. B; Whole genome phylogenetic tree and shared average nucleotide identities among the seven strains of interest.

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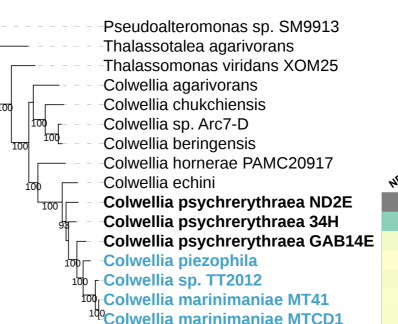
1330 Figure 2. A; Isoelectric point distribution of proteins within piezophilic (blue points) or

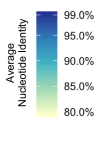
- piezosensitive (black) strains, with an average line of fit within each group. B; Isoelectric point
- 1332 protein bias within each strain as a function of their growth pressure. C; Asymmetry index values
- indicating preference of amino acids in the piezophiles or *C. psychrerythraea* 34H within
- 1334 orthologous proteins present in all strains. D; Specific amino acid substitutions from *C*.
- 1335 *psychrerythraea* 34H to the piezophiles within orthologous proteins. The substitutions shown
- 1336 were also identified within comparisons between piezophilic and piezosensitive *Shewanella*.
- 1337
- 1338 Figure 3. A; The percent abundance of proteins within each COG category within piezophilic or
- piezosensitive *Colwellia*. B) Specific genomic attributes that were differentially present in
 piezophilic or piezosensitive strains. Present, grey; absent, white.
- 1341
- 1342 Figure 4. A; The location of a d-ala d-ala ligase gene in strains MT41 and TT2012, with
- 1343 surrounding genes labeled. B; An amino acid tree of the d-ala-d-ala ligase with sequences 1344 approximately > 50% similar shown.
- 1345
- 1346 Figure 5. A; The location of alanine dehydrogenase genes in strains MT41 and TT2012, with
- 1347 surrounding genes labeled. B; An amino acid tree of the alanine dehydrogenase with sequences
- 1348 approximately > 50% similar shown.



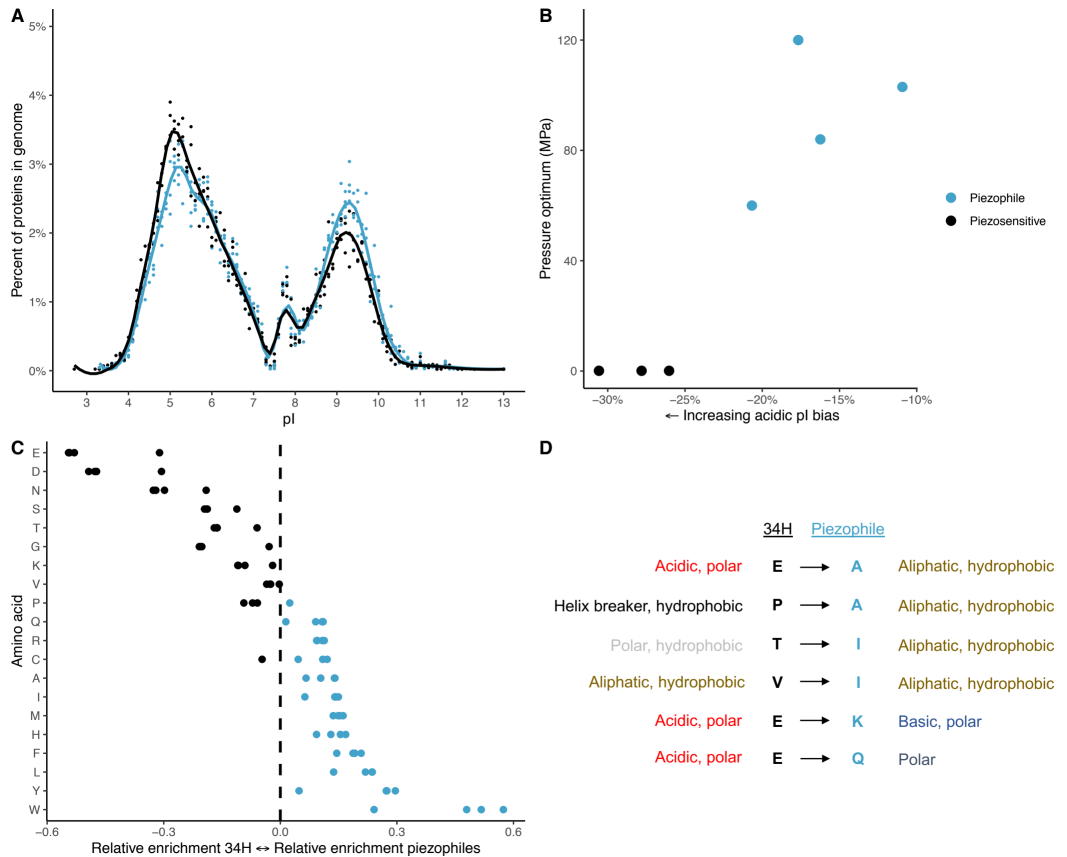
Longitude

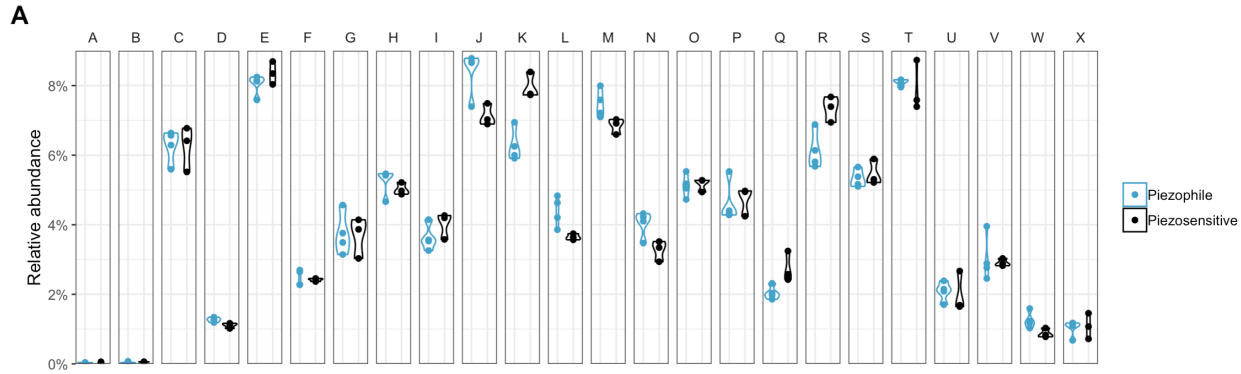
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T41			24	22								2		2				35	30	66		

MT41		24	22				2	2		35	30	66	
MTCD1		21	23				2	2		24	33	57	
TT2012		24	22				2	2		25	24	61	
C. piezophila		16	41				2	2		26	25	12	
GAB14E		15	60				1	1		30	18	39	
ND2E		12	70				1	1		36	12	21	
34H		11	92				1	1		24	9	32	

В

