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Microbial diversity of extreme habitats in human homes

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High-throughput sequencing techniques have opened up the world of microbial diversity to scientists, and a flurry of studies in the most remote and extreme habitats on earth have begun to elucidate the key roles of microbes in ecosystems with extreme conditions. These same environmental extremes can also be found closer to humans, even in our homes. Here, we used high-throughput sequencing techniques to assess bacterial and archaeal diversity in the extreme environments inside human homes (e.g., dishwashers, hot water heaters, washing machine bleach reservoirs, etc.). We focused on habitats in the home with extreme temperature, pH, and chemical environmental conditions. We found a lower diversity of microbes in these extreme home environments compared to less extreme habitats in the home. However, we were nonetheless able to detect sequences from a relatively diverse array of bacteria and archaea. Habitats with extreme temperatures alone appeared to be able to support a greater diversity of microbes than habitats with extreme pH or extreme chemical environments alone. Microbial diversity was lowest when habitats had both extreme temperature and one of these other extremes. In habitats with both extreme temperatures and extreme pH, taxa with known associations with extreme conditions dominated. Our findings highlight the importance of examining interactive effects of multiple environmental extremes on microbial communities. Inasmuch as taxa from extreme environments can be both beneficial and harmful to humans, our findings also suggest future work to understand both the threats and opportunities posed by the life in these habitats.

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1	Microbial diversity of extreme habitats in human homes
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27 Abstract:

High throughput sequencing techniques have opened up the world of microbial diversity to 28 29 scientists, and a flurry of studies in the most remote and extreme habitats on earth have begun to 30 elucidate the key roles of microbes in ecosystems with extreme conditions. These same 31 environmental extremes can also be found closer to humans, even in our homes. Here, we used 32 high throughput sequencing techniques to assess bacterial and archaeal diversity in the extreme environments inside human homes (e.g., dishwashers, hot water heaters, washing machine bleach 33 34 reservoirs, etc.). We focused on habitats in the home with extreme temperature, pH and chemical 35 environmental conditions. We found a lower diversity of microbes in these extreme home environments compared to less extreme habitats in the home. However, we were nonetheless 36 able to detect sequences from a relatively diverse array of bacteria and archaea. Habitats with 37 extreme temperatures alone appeared to be able to support a greater diversity of microbes than 38 habitats with extreme pH or extreme chemical environments alone. Microbial diversity was 39 40 lowest when habitats had both extreme temperature and one of these other extremes. In habitats with both extreme temperatures and extreme pH, taxa with known associations with extreme 41 conditions dominated. Our findings highlight the importance of examining interactive effects of 42 43 multiple environmental extremes on microbial communities. Inasmuch as taxa from extreme environments can be both beneficial and harmful to humans, our findings also suggest future 44 work to understand both the threats and opportunities posed by the life in these habitats. 45

<u>Keywords:</u> Community Structure, Extreme environments, Human Homes, Interactive effects,
Microbial Diversity

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51 **Introduction:**

52 The innovation of culture-independent, high-throughput sequencing techniques has facilitated the 53 discovery of high microbial diversity in many habitats once considered inhospitable to life (Rothschild and Mancinelli 2001). The species in these environments are frequent targets for the 54 55 discovery of useful enzymes (Niehaus et al. 1999, van den Burg 2003, Elleuche et al. 2014), and studies of microbes living in extreme environments have provided key insights into the evolution 56 of microbial metabolism (Valentine 2007, Hoehler and Jorgensen 2013). Often overlooked, 57 however, is that the attributes that define many of the most extreme habitats on Earth, such as 58 extremes of temperature, pH, water activity, or low nutrient levels, can also be found more 59 immediate to everyday experience. Human homes, for example, contain microhabitats as hot, 60 acidic, basic or salty as any encountered elsewhere on Earth (Martin et al. 2015). 61 We know of only two extreme habitats within homes where microbial diversity has been studied 62

to date, and in both cases culture-dependent techniques were used. In 1973, Brock and Boylen discovered a species of the genus *Thermus* (*T. aquaticus*) living in hot water heaters. Species of this genus had previously been known only from hot springs (Brock and Boylen 1973). In addition, studies have considered the biology of tap water. Tap water is hospitable in terms of its abiotic conditions (e.g. temperature, pH, toxicity) but is very low in nutrients and so was long

assumed to be relatively devoid of life; until, that is, it was studied. Tap water has now been
shown to contain many species of bacteria capable of surviving in low nutrient environments
(Kalmbach et al. 1997, Szewzyk et al. 2000, Boe-Hansen et al. 2002). If life exists in hot water
heaters and tap water, it seems possible and even likely that many extreme habitats in homes
sustain life. That the environmental extremes imposed by these conditions in homes (cold, hot,
acidic, alkaline, wet or dry) delineate which species are present seems inevitable. That they are
lifeless is unlikely.

Here, we used culture-independent, high-throughput sequencing to address the following 75 76 questions: (1) What is the relative diversity of microbes (specifically, Bacteria and Archaea) under extremes of temperature, pH and chemical environments of southeast US homes and how 77 does it compare to habitats without each extreme conditions? Harrison et al. (2013) recently 78 argued that because many extreme environments include simultaneous extremes in multiple 79 environmental factors, interactive effects of these multiple sources of extreme conditions are 80 81 likely to be important determinants of microbial diversity in extreme environments. Therefore, we additionally asked (2) how do multiple, simultaneous extreme conditions influence microbial 82 diversity in human homes? Finally, we asked (3) which bacterial and archaeal genera from the 83 84 broader home (Dunn et al. 2013) fail to persist in extreme home habitats, and which microbial genera persist only in these extreme habitats? 85

86 Methods:

87 Sampling extreme home environments

88 We sampled extreme environments in six houses in the Raleigh-Durham metropolitan area

89 (Supp. Fig 1). In each house, we used dual-tipped sterile BBL^{TM} CultureSwabsTM or 50 ml

conical tubes to collect water from each of 10 standardized extreme locations in homes. The sites 90 sampled in all six houses included environments that were extreme in terms of their temperature, 91 pH and/or chemical environments (Supp. Table 1). Our assumptions concerning these sampling 92 locations are based upon publicly available consumer resources regarding certain commercial 93 and industrial requirements (e.g. http://www.nsf.org/consumer-resources/health-and-safety-94 95 tips/home-product-appliance-tips/sanitizing-dishwasher, http://energy.gov/energysaver/projects/savings-project-lower-water-heating-temperature). For 96 example, our sampling of dishwashers was influenced by the NSF/ANSI 184 standard for 97 residential dishwashers to provide a final rinse at a temperature of at least 150 °F (65.6 °C). 98 Additionally, temperature ranges for residential water heaters are 90 to 150 F (32-65.6 °C), 99 depending on the manufacturer. Bleach receptacles in clothes washing machines would also be 100 assumed to have a pH of 12 when bleach is present. Although the pH and chemical composition 101 of laundry detergent and dishwasher detergent can be quite variable, manufacturing standards are 102 103 generally within the 7-10 pH range. While measurements, opposed to assumptions, would be very useful, taking measurements of all the potential extreme axes under various sample sites in 104 multiple homes was not feasible. All samples were preserved at -20° C immediately after 105 collection. 106

107 Isolating and identifying microbes in extreme home environments

Genomic DNA was extracted from all samples using the MoBio Power Soil DNA extraction kit
(MoBio, Carlsbad, CA) as described previously (Fierer et al., 2008; Lauberet al., 2009). For
swabs, the tips were placed in PowerBead tubes containing solution C1 and swirled vigorously
for approximately 10 seconds to release contents and removed. Water samples were thawed and
filtered using Corning 50 ml 0.22um cellulose acetate filters after which the filters were added to

the PowerBead tubes. The extractions were subsequently performed as directed by the 113 manufacturer, except that the final elution was performed in 50µl of 70° C C6 elution buffer. 114 Because the water samples were frozen prior to filtering and extraction, the results reported for 115 the water samples likely under-represents the true diversity of taxa in those environments. 116 We used methods described in Bates et al (2011) to amplify bacterial and archaeal DNA from the 117 118 samples collected from homes and six negative controls. Briefly, amplicons were produced by PCR with universal bacterial/archaeal 515F and 806R primers to which Roche 454 B 119 pyrosequencing adapters had been added, as described in Hulcr et al. (2012). The 515F primer 120 contained an additional 12-bp barcode sequence for individual sample identification. All the 121 samples were amplified by triplicate PCR reactions, cleaned using the UltraClean-htp 96-well 122 PCR Clean-up kit (MoBio), and quantified with a Quant-iT PicoGreen dsDNA Assay kit 123 (Invitrogen). Equimolar amounts of each sample were pooled into a single sample to sequence. 124 125 DNA pyrosequencing was performed at Selah Clinical Genomics Center at Innovista (University 126 of South Carolina, USA) using a Roche Genome Sequencer 454 FLX system to facilitate comparison to previous related work that utilized this platform (Dunn et al, 2013). Though these 127 methods here do not distinguish living from recently dead cells, with the comparative approach 128 129 used here, we presume that taxa frequently identified in one habitat but rare or absent in most others are likely surviving in the habitat from which they are frequently identified. The 130 sequences were submitted to NCBI (SRA accession number SRP071677). 131 The QIIME analysis package (Caporaso et al, 2010a) was used to process and analyze the 132 barcoded microbial (bacterial and archaeal) amplicon sequences. Sequences were quality filtered 133 134 to a minimum quality score of 25 with no unambiguous bases and sorted to each sample by the 12 bp barcodes. The 454 pyrosequencing produced 197,305 reads that passed the quality 135

screening. The sequences were grouped into Operational Taxonomic Units (OTUs) that shared at 136 least 97% sequence similarity. A representative sequence was taken for each OTU group and 137 PyNAST (Caporaso et al, 2010b) was employed to align these representative sequences to the 138 Greengenes database (DeSantis et al, 2006) and the taxonomic identity of each OTU was 139 determined using the RDP Classifier (Wang et al, 2007). Phylotypes were considered to be 140 141 contaminants if they were seen in at least two of the six negative control samples. There were 152 OTUs at the genus level present in more than one negative sample, representing 9% of the 142 143 total OTUs at this level. After removing contaminant sequences and singletons, the number of quality-filtered reads per sample was between 6 and 5861 (median=2306). Finally, we removed 144 any OTU's represented by 20 or fewer reads to reduce the possibility of spurious results from the 145 sequencing process. For among samples comparisons we rarefied to each to a depth of 1000 146 sequences. Thus, our assessments of microbial diversity are conservative. 147

148

Analysis of the relative diversity of microbes in extreme temperature, pH and chemical 149 environments of homes and how it compares to habitats without each extreme condition 150 151 We compared microbial species accumulation among three extreme variables in homes: temperature, pH, and chemical extremes. Temperature was classified on a scale of 1-5, with 1 152 representing the coldest environments and 5 representing the hottest environments. We then 153 154 binned 1 and 5 into an extreme temperature category and 2-4 into an intermediate temperature category. Similarly, environments were classified as acidic, basic or neutral and then binned into 155 extreme pH (acidic or basic environments) versus neutral environments. Finally, chemical 156 157 extremes were those environments characterized by the presence of detergent, bleach, metals, ammonia, or natural gas (Supp. Table 2). 158

We used EstimateS v. 9.1.0 (Colwell 2013) to construct individual-based species accumulations for all three extreme environments and their non-extreme counterparts. For these curves, reads were used as individuals and the curves were constructed using 1000 iterations. To formally assess differences in accumulated species by read, we used \pm 95% confidence intervals for each curve. Non-overlapping 95% confidence intervals are considered formal evidence of significance (Knezevic 2008).

Assessing how multiple, simultaneous extreme conditions influence microbial diversity in human
homes

We were interested in testing the hypothesis that interactive effects of multiple, simultaneously 167 extreme environmental conditions are important determinants of microbial diversity in extreme 168 home environments (Harrison et al. 2013). Our study included multiple samples with more than 169 one environmental extreme (Supp. Table 1); however, we only had sufficient replication to 170 assess this hypothesis for 2-way interactions between extremely high temperatures and extreme 171 pH as well as high temperature and chemical environments. Because number of reads varied 172 173 significantly among different environmental extremes, we could not use a standard 2-way ANOVA. Instead, we assessed these effects using an ordination framework. 174

We visualized the composition of bacteria and archaea from extreme habitats in homes using
non-metric multidimensional scaling ordination (NMDS) in Primer-E v.7.0.9 with

177 PERMANOVA +1 (Clarke & Gorley, 2015). To do this, we first constructed NMDS plots with

178 100 restarts and a Type I Kruskal fit scheme based on a Dissimilarity matrix of Bray-Curtis

179 distances. To assess the relationship between temperature (extreme vs. intermediate) and the

- 180 other extremes (pH: extreme vs. neutral; chemicals: extreme vs. none) for a-diversity of
- 181 microbes, we conducted a permuted multivariate analysis of variance (PERMANOVA) test with

temperature class and either pH or chemical class and their interaction as factors, 9,999 iterations 182 and Type III sums of squares. When interactions were significant (Anderson et al. 2008), we 183 conducted pairwise PERMANOVA to determine which treatment combinations significantly 184 differed from one another. Similarly, we assessed these relationships in terms of β -diversity 185 using a permuted dispersion (PermDisp) test of a presence/absence matrix of OTU occurrences. 186 187 When these tests were significant, we conducted pairwise tests of extreme vs. non-extreme chemical and pH environments in habitats with intermediate and extreme temperatures (thus 2 188 tests per treatment combination). Finally, we conducted SIMPER analyses for each significant 189 treatment combination to determine the OTUs that contributed the most to pairwise between-190 group differences in ordination space. Because we conducted two separate analyses for each 191 level of diversity, we accounted for the additional error associated with multiple tests, using a 192 revised a=0.05/2=0.025 as our cut-off for statistical significance for the results of each test. This 193 conservative a is particularly important because we did not have equal sample sizes in all groups 194 195 for these analyses, which can increase the risk of Type I error (Anderson & Walsh 2013).

196

197 Determining which microbial genera differentiate extreme home habitats from the rest of the198 home

We compared the occurrences of microbes in our samples to those reported in less extreme home environments (Dunn et al. 2013). Human-associated microbes were common to both datasets, and we were particularly interested in those taxa unique to our dataset, relative to the broader home (Dunn et al. 2013). Therefore, we removed human-associated OTU's from our dataset. We identified these human-associated OTU's using databases that identified human gut (Flores et al. 2014) and skin (Urban et al. 2016) microbiomes. OTU's that occurred in at least 80% of the

samples in those databases were considered human-associates and excluded from our analyses of the microbial diversity of extreme habitats in human homes. We then determined the identity of microbes that were absent from the broader homes dataset, but present in extreme environments and then tabulated the extreme habitat(s) in which they were present. Likewise, we identified the non-human associated microbes that were present in the broader home environment, but absent from all extreme environments in our samples.

211

212 **Results and Discussion:**

What is the relative diversity of microbes in extreme temperature, pH and chemical environments
of southeast US homes and how does it compare to habitats without each extreme condition?

The cumulative diversity (OTU richness) in habitats with extreme temperatures was more 215 than twice as high as in habitats with extreme pH (maximum of 73 vs. 33, Fig. 1) and almost 216 three times as high as habitats with extreme chemical environments (27.6; Fig. 1). Habitats with 217 extreme temperatures also had higher OTU richness than habitats with intermediate temperatures 218 (Fig. 2a). Conversely, previous research indicates that the diversity in habitats with either 219 extremely high or extremely low temperatures is generally low, and dominated by a small 220 number of abundant bacterial species (Lewin et al. 2013). For example, Sharp et al. (2014) 221 recently found that OTU richness in hydrothermal vents peaked at intermediate temperatures 222 (24°C), with reduced OTU richness in extremely hot or cold environments (Sharp et al. 2014). 223 We did not detect significant differences in the rarefied species richness of bacterial and archaeal 224 225 microbes in extreme vs. neutral pH conditions; however, the marginally non-significant trend suggests that extreme pH environments also had higher microbial diversity than neutral habitats 226

(Fig. 2b). Recent studies have demonstrated that pH is a key predictor of microbial diversity in 227 both extreme environments, such as acid mine drainage sites (Kuang et al. 2013), and less 228 extreme environments, such as tropical soils (Tripathi et al. 2012). In both cases, habitats with 229 neutral pH had higher microbial diversity than those with a pH higher or lower than neutral. 230 Thus, we again found different patterns in extreme home environments compared to other studies 231 232 comparing extreme and non-extreme habitats. One possible explanation for the difference between our findings and these recent studies is that human-associated microbes are present in 233 home environments with intermediate temperatures. Perhaps these species are able to dominate 234 habitats with intermediate, but not extreme, conditions. Alternatively, the lower diversity in 235 habitats with intermediate temperatures and neutral pH in our study could be due to the 236 occurrence of extreme conditions along different axes (e.g. intermediate temperature, but 237 extreme pH or chemical habitats). We examine potential interactive effects of these polyextreme 238 habitats in the next section. 239

In contrast, habitats with extreme chemicals had significantly lower accumulated OTU richness than did habitats without these extreme conditions (Fig. 2c). Extreme chemical environments are poorly studied and understood (Rothschild and Mancinelli 2001). Our data suggest that they could act as strong filters in extreme environments.

244 How do multiple, simultaneous extreme conditions influence microbial diversity in extreme home245 environments?

Many of the habitats in this study were characterized by more than one extreme environmental condition. Therefore, we also examined the potential for interactive effects of multiple, simultaneous extreme conditions on microbial diversity. Due to limited replication across all environmental extremes, we were only able to examine extreme pH and chemical

habitats with and without extreme temperatures. We used an ordination framework to examinethese interactive effects (see methods).

252	We found significant interactions between extreme temperature and both extreme pH
253	(PERMANOVA: Pseudo- $F_{1, 82}$ = 2.53, <i>P</i> =0.0001; Figure 3A) and extreme chemical
254	(PERMANOVA: Pseudo- $F_{1, 82}$ = 3.16, <i>P</i> =0.0001; Figure 3C) environments for OTU
255	composition. When temperatures were intermediate, there were no significant differences in
256	microbial composition in extreme vs. neutral pH habitats (pairwise PERMANOVA: $t_{1, 38} = 1.02$,
257	P = 0.40). However, when temperatures were extreme, there was a very large difference between
258	the composition of microbes in extreme pH habitats, compared to neutral habitats (pairwise
259	PERMANOVA: $t_{1,38} = 1.70$, <i>P</i> =0.0001; Fig. 3A). The five genera that contributed the most to
260	differences between these two habitat types (from SIMPER analysis) were Parascardovia,
261	Micrococcus, Rothia, Brachybacterium, and an unknown genus from Sphingomonadaceae. Most
262	of these genera are associated with humans (Oshima et al. 2015, Gueimonde et al. 2012, Kloos et
263	al. 1975, Kocur et al. 2006, Vaccher et al. 2007, Uchibori et al. 2012). Sphingomonadaceae are
264	widespread in aquatic habitats, including drinking water (Vaz-Moreira et al. 2011), but also other
265	aquatic environments (e.g. tree holes-Xu et al. 2008). Brachybacterium is usually associated with
266	marine environments (Ward and Boru 2006), including Antarctic sea ice (Junge et al. 1998).
267	However, it was recently detected in an urban shopping center (Tringe et al. 2008). All of these
268	genera were more common in habitats with extreme temperatures and neutral pH than they were
269	in habitats with both extreme temperatures and extreme pH. While different houses had
270	significantly different microbial composition (3-way PERMANOVA, P=0.0001), there were no
271	significant 2-or 3-way interactions with house (Supp. Table 3).

The interaction between temperature and chemical extremes was similar. Microbial composition 272 was indistinguishable between the habitats that only had one extreme condition-regardless of 273 whether it was temperature or chemicals that were extreme. There were also no significant 274 differences between habitats with neither extreme temperatures nor extreme chemical conditions 275 and habitats that had a single extreme condition. However, habitats with both extreme 276 277 temperatures and extreme chemicals had significantly different microbial composition compared to all other groups (pairwise PERMANOVA; $t_{1,38} = 1.75$, P=0.0001; Fig. 3C). The five genera 278 279 that contributed the most to compositional difference between these two habitats (from SIMPER analysis) were Methylobacterium, an unknown genus of Moraxellaceae, Sejonia, an unknown 280 genus of Sphingomonadaceae, and Flavobacterium. With the exception of the unknown genus 281 of Moraxellaceae, which was more common in extreme chemical and temperature environments, 282 all of these genera were more common in the habitats without temperature and chemical 283 extremes. Moraxallaceae have been found in other extreme environments, including deep sea 284 285 sediments (Maruyama et al. 1997). Although it was more common in our less extreme environments, Sejonia is better known from Antarctic ice (Yi et al. 2005). Sphingomonadaceae 286 as described above are common to aquatic habitats. *Methylobacterium* is a widespread habitat 287 288 generalist that is facultatively methyltrophic (Green 2006). Finally, *Flavobacterium* is common in freshwater and marine ecosystems but tends to flourish in cold environments with high salinity 289 290 (Bernardet and Bowman 2006).

There were also significant differences in the β -diversity in home environments with more than one extreme condition. When temperatures were intermediate, there were no significant differences between neutral and extreme pH environments (Figure 4A; PermDisp: *P* = 0.3864). However when temperatures were also extreme, habitats with extreme pH conditions had

significantly higher β -diversity than those with neutral pH conditions (Figure 4A; PermDisp: 295 P=0.0014). Similarly, at intermediate temperatures, there was a non-significant trend (Figure 4B; 296 PermDisp: P=0.03, Bonferroni-corrected a = 0.025) in which habitats without extreme 297 chemicals present had higher β -diversity than those with extreme chemicals present. However, 298 when temperatures were also extreme, habitats with extreme chemicals present had higher β -299 300 diversity than those without extreme chemicals (Figure 4B; PermDisp: P = 0.0006). This increase in β-diversity in extreme pH and chemical environments when temperatures were also 301 extreme suggests that polyextreme conditions may support a higher diversity of extremophiles 302 and/or reduced occurrences of numerically dominant genera compared to environments with a 303 single extreme condition, at least among habitats (in contrast to within habitats). The 5 genera 304 that contributed the most to differences in β -diversity between neutral and extreme pH conditions 305 when temperatures were also extreme were: Veillonella, Kocuria, Peptoniphilus, Parascardovia, 306 and Anaerococcus. Interestingly, these were also the top 5 genera contributing to differences 307 308 between habitats with and without extreme chemicals that also had extreme temperatures. All of these genera were less common in habitats with 2 extremes than they were in habitats with only 309 extreme temperatures. They are also genera that include human-associated species (Bhatti & 310 311 Frank 2000, Fadda et al. 2001, Song et al. 2007, Gueimonde et al. 2012).

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313 Which microbial genera differentiate extreme home habitats from the rest of the home?

After removing all human-associated microbes (see methods), there were a total of 241 unique genera in the broader homes dataset (Dunn et al. 2013). Our extreme samples contained 135 of the remaining broader homes genera, but ~44% of the genera found in the broader homes were absent from our extreme home samples (Supp. Table 4), the absence of which might simply be

due to the larger number of samples in Dunn et al. (2013). More interestingly, we found 20 318 genera present among our samples that were absent from the broader homes dataset. Nine of 319 these genera were found in all three categories of extreme environments (Table 1); one genus 320 (Solibacter) was absent from habitats with extreme pH, but occurred in both extreme chemical 321 and temperature environments. Solibacter is a common and abundant soil microbe, especially in 322 323 tropical regions (Guan et al. 2013, Wang et al. 2015). There was also one genus (Brevundimonas) that was absent from extreme chemical environments, but present in both 324 extreme temperature and extreme pH environments; *Brevundimonas* is one of the only genera 325 thought to be able to survive the low temperatures and ionizing radiation on Mars (Dartnell et al. 326 2010). There were three genera (Azobacteroides, Elizabethkingia, and Xiphinematobacter) that 327 occurred in both extreme pH and chemical environments that were absent in extreme 328 temperature environments. Both Azobacteroides and Xiphinematobacter are gut symbionts of 329 invertebrates; Azobacteroides is commonly found inside the protozoan symbionts of termites 330 331 (Noda et al. 2007), and *Xiphinematobacer* is an endosymbiont of nematodes (Vandekerckhove et al. 2000). In invertebrate guts these microbes likely experience extreme chemical and pH 332 environments frequently, while being relatively protected from temperature stress. 333 334 *Elizabethkingia* is a cosmopolitan genus, with species that are endosymbionts of mosquitoes (Kämpfer et al. 2011), and others that are pathogens of both humans (Ceyhan and Celik 2011) 335 336 and frogs (Xie et al. 2009). There was one genus that was only found in extreme chemical 337 environments (*Helcococcus*). Interestingly, members of the genus *Helcococcus* possess the 338 ability to degrade detergents. In fact, the detergent Tween-80 can be added to media to enrich Helcococcus (Collins et al. 1993, Chagla et al. 1998). Finally, we found 5 genera (Brochothrix, 339 340 Buchnera, Polynucleobacter, Ralstonia, and Thermicanus) unique to extreme temperature

environments. *Brochothrix* is a common spoilage bacterium in meat (Rattanasomboom et al. 341 1999). Buchnera is a widespread aphid endosymbiont (Shigenobu et al. 2000). Recently, a 342 survey of homes in Raleigh, NC demonstrated that aphids could be quite common in human 343 homes (Bertone et al. 2016), which could explain how this genus arrived in the homes in our 344 study (via aphids in the home). The genus *Polynucleobacter* includes both free-living species and 345 346 species that are endosymbionts of nematodes (Vannini et al. 2007). Ralstonia metallidurans is a bacterium specifically adapted to toxic metal environments (Mergeay et al. 2003). Other species 347 of *Ralstonia* have been shown to be effectively controlled using high temperature treatments in 348 commercial crops (Kongkiattikajorn and Thepa 2007). In our study, Ralstonia were collected in 349 both high and low temperature environments. Finally, *Thermicanus* is, as its name suggest, a 350 thermophilic bacterial genus (Wrighton et al. 2008). 351

352 Conclusions:

This study has provided a glimpse into the microbial diversity that lives in habitats of human 353 homes similar in their extreme temperature, pH and chemical conditions to some of the most 354 extreme habitats on Earth. We discovered that these conditions have lower diversity than the 355 surrounding home environment; yet tens of bacterial lineages can be found in these extreme 356 habitats of the human home, including many taxa with known associations with extreme 357 conditions. Habitats with extreme temperatures alone appear to be able to support a greater 358 diversity of microbes than habitats with extreme pH or extreme chemical environments alone. 359 Microbial diversity is significantly lowest when habitats have both extreme temperature and one 360 of these other extremes. Interestingly, environments in homes often alternate between periods of 361 362 extreme and non-extreme conditions. For example, dishwashers are only likely to have extremely high temperatures while cleaning and drying dishes. This variability could lead to temporal shifts 363

in microbial composition, similar to those found for human vaginal microbes (Gajer et al. 2012). 364 This variability may also explain the presence of human-associated generalist species in our 365 samples. Future work, with samples taken before and after appliances (like many of those used in 366 our study) are operated, could elucidate the importance of episodic extreme conditions for 367 microbial communities in homes. Additionally, a key next step is understanding which of the 368 369 relatively few species that are found in these poly-extreme environments in the home are metabolically active there and both whether these polyextreme taxa pose health threats (as was 370 recently suggested by Gümral et al. 2015) and/or might be useful industrially. 371

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539 Figure Captions:

540 <u>Figure 1: OTU accumulation curves for each extreme environment</u>, expressed as number of

541 OTUs by number of reads from sequencing. Each curve was constructed using 1000 iterations,

and the dotted lines represent 95% confidence intervals. Non-overlapping confidence intervals

543 indicate that the accumulation curves are significantly different. Thus, habitats with extreme

544 temperatures had significantly more accumulated species than habitats with either extreme pH or

545 extreme chemical environments. However, the accumulated species in habitats pH and chemical

546 extremes did not differ significantly.

547 Figure 2: Comparison of rarefaction curves between extreme and non-extreme habitats. (A)

548 extreme vs. intermediate temperatures, (B) extreme vs. neutral pH environments, and (C)

549 extreme chemicals present *vs.* absent. Rarefaction curves are expressed as number of OTU by

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number of reads from sequencing. Each curve was constructed using 1000 iterations, and the
dotted lines represent 95% confidence intervals. Significance tests were as described for Figure
1.

553 Figure 3: NMDS ordinations OTU occurrence by (A-B) Temperature & pH and (C-D)

- 554 <u>Temperature & chemical environments in the home.</u> Large symbols represent centroids ± 1 SE
- 555 (A, C), and small symbols represent each sample (B, D). The interaction between temperature
- and pH was significant (PERMANOVA: (pseudo)- $F_{1,85} = 2.53$, P(perm) = 0.0001), as was the
- 557 interaction between extreme temperature and chemical conditions (PERMANOVA: (pseudo)- $F_{1,}$
- 558 $_{85}$ = 3.16, P(perm) = 0.0001). The ordination was constructed with Bray-Curtis distances and 100
- restarts; 2-D stress was 0.21. PERMANOVA analyses were conducted using type III sums of
- squares and 9,999 iterations.

561 Figure 4: Average distances between samples and centroids (β -diversity) across home

- 562 <u>environments</u> that differ with respect to extreme temperatures and (A) extreme pH conditions &
- 563 (B) extreme chemical conditions. Data were assessed using PermDisp; dispersion was
- significantly different across extreme temperatures and extreme pH conditions ($F_{3, 82} = 4.08$,
- 565 *P*=0.024) and across extreme temperatures and extreme chemical conditions ($F_{3, 82} = 6.99$,
- 566 *P*=0.0017). Post-hoc pairwise tests: * *P*<0.025, ** P<0.01, *** P<0.001.

567 **Table Captions:**

568 <u>Table 1:</u> Summary of occurrences of microbes that were present in samples from extreme home 569 environments, but absent from the broader home samples. Numbers indicate the number of reads 570 of each genus by extreme environment. The first group includes genera that were only present in 571 one extreme environment, the second group includes genera that were present in two extreme

572 environments, and the last group includes genera that were present in all three extreme home

573 environments.

574 Supplementary Tables and Figures:

575 Raw Data: Output file from QIIME at the genus level (L6), rarefied to 1000 reads.

576 Supp. Table 1: Description of sample locations. Standardized locations were sampled in all 6

houses, while special locations were only sampled in a subset of the houses (due to availabilityof samples across houses)

579

- Supp. Table 2: Classifications of sampled extreme home environments based upon temperature,pH and chemical conditions.
- 582 Supp. Table 3: Results from PERMANOVAs testing (a) the effects of house ID on microbial
- composition; and (b) the effects of temperature class, pH class, and house ID on microbialcomposition.
- Supp. Table 4: List of non-human associated microbes in extreme and non-extreme (Dunn et al.
 2013) home habitats

587

- 588 Supp. Figure 1: Map of houses that were sampled for the study
- 589 Supp. Figure 2: NMDS plot with houses and sampling locations labeled.
- 590 Supp. Figure 3: Histograms depicting the % of reads from bacterial and archaeal classes in (a)
- habitats with extreme temperatures, (b) habitats with extreme pH, and (c) habitats with extremechemicals.
- 593
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Table 1(on next page)

Table 1

Summary of occurrences of microbes that were present in samples from extreme home environments, but absent from the broader home samples. Each X indicates that the genus was found in a given extreme environment. theh6\0e>

1

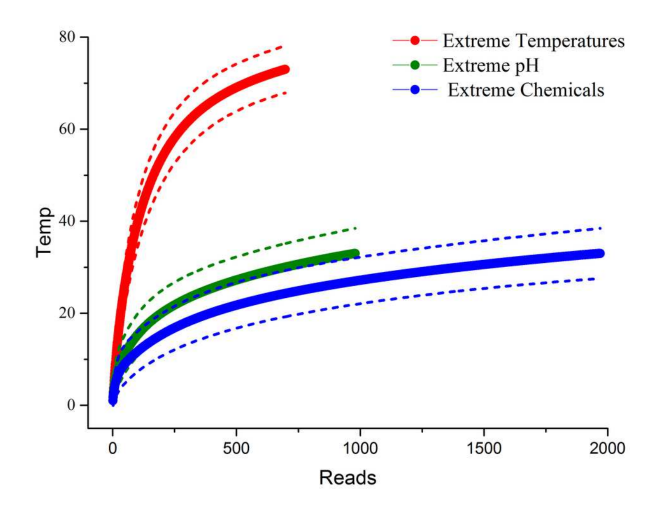
- 2 <u>Table 1</u>: Summary of occurrences of microbes that were present in samples from extreme home
- 3 environments, but absent from the broader home samples. Numbers indicate the number of reads of each
- 4 genus by extreme environment. The first group includes genera that were only present in one extreme
- 5 environment, the second group includes genera that were present in two extreme environments, and the
- 6 last group includes genera that were present in all three extreme home environments.

7

8	Genus	Extreme Temperatures	Extreme pH	Extreme Chemical
9	Brochothrix	265	0	0
10	Buchnera	22	0	0
11	Polynucleobacter	33	0	0
12	Ralstonia	21	0	0
	Thermicanus	34	0	0
	Helcococcus	0	0	22
			<u> </u>	1
	Solibacter	86	0	30
	Brevundimonas	184	189	0
	Azobacteroides	0	33	33
	Elizabethkingia	0	25	24
	Xiphinematobacter	0	19	21
				I
	Azospira	139	33	44
	Brachybacterium	101	52	69
	Enhydrobacter	452	387	408
	Gluconobacter	23	21	22
	Oligella	40	74	77
	Parascardovia	141	46	107
	Photobacterium	71	65	93
	Propionibacterium	73	31	40
	Salinibacterium	108	334	355

OTU accumulation curves for each extreme environment

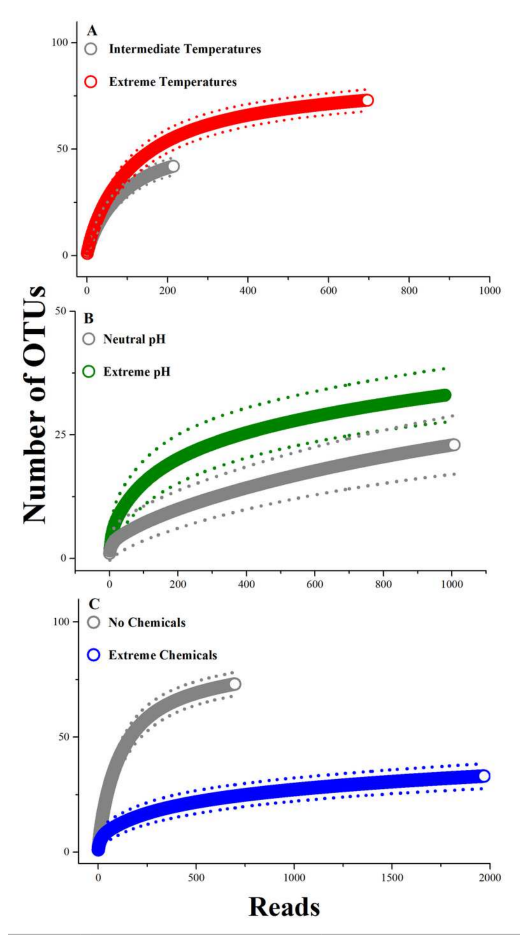
OTU accumulation curves for each extreme environment , expressed as the number of OTUs by the number of reads from sequencing. Each curve was constructed using 1000 iterations, and the dotted lines represent 95% confidence intervals. Non-overlapping confidence intervals indicate that the accumulation curves are significantly different. Thus, habitats with extreme temperatures had significantly more accumulated species than habitats with either extreme pH or extreme chemical environments. However, the accumulated species in habitats pH and chemical extremes did not differ significantly.



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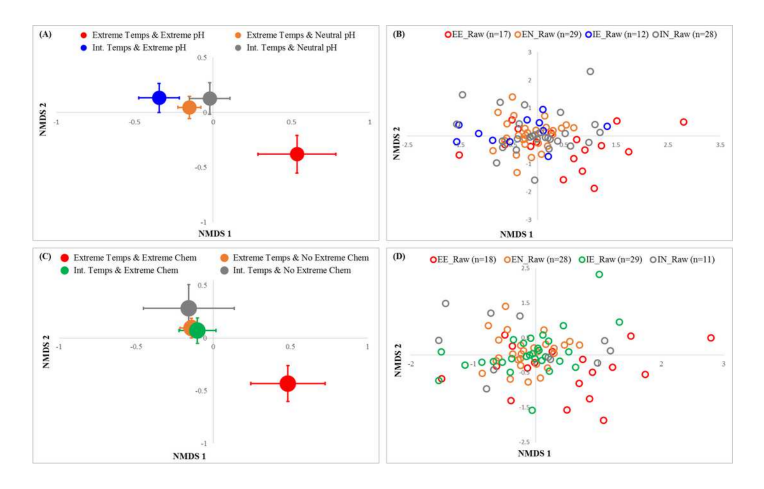
Comparison of rarefaction curves between extreme and non-extreme habitats

Comparison of rarefaction curves between extreme and non-extreme habitats . (A) extreme *vs.* intermediate temperatures, (B) extreme *vs.* neutral pH environments, and (C) extreme chemicals present *vs.* absent. Rarefaction curves are expressed as number of OTU by number of reads from sequencing. Each curve was constructed using 1000 iterations, and the dotted lines represent 95% confidence intervals. Significance tests were as described for Figure 1.



NMDS ordinations OTU occurrence by (A-B) Temperature & pH and (C-D) Temperature & chemical environments in the home.

NMDS ordinations OTU occurrence by (A-B) Temperature & pH and (C-D) Temperature & chemical environments in the home. Large symbols represent centroids ± 1 SE (A, C), and small symbols represent each sample (B, D). The interaction between temperature and pH was significant (PERMANOVA: (pseudo)-F_{1,85} = 2.53, *P*(perm) = 0.0001), as was the interaction between extreme temperature and chemical conditions (PERMANOVA: (pseudo)-F_{1,85} = 3.16, *P*(perm) = 0.0001). The ordination was constructed with Bray-Curtis distances and 100 restarts; 2-D stress was 0.21. PERMANOVA analyses were conducted using type III sums of squares and 9,999 iterations.



Average distances between samples and centroids (β -diversity) across home environments

Average distances between samples and centroids (β -diversity) across home environments that differ with respect to extreme temperatures and (A) extreme pH conditions & (B) extreme chemical conditions. Data were assessed using PermDisp; dispersion was significantly different across extreme temperatures and extreme pH conditions ($F_{3, 82} = 4.08$, P=0.024) and across extreme temperatures and extreme chemical conditions ($F_{3, 82} = 6.99$, P=0.0017). Post-hoc pairwise tests: * P<0.025, ** P<0.01, *** P<0.001.

