Phages actively challenge niche communities in the Antarcticsoils

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16	Running Head: host-phage interactions in Antarctica
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24 Abstract

25 By modulating the structure, diversity and trophic outputs of microbial communities, 26 phages play crucial roles in many biomes. In oligotrophic polar deserts, the effects of 27 katabatic winds, constrained nutrients and low water availability are known to limit 28 microbial activity. Although phages may substantially govern trophic interactions in 29 cold deserts, relatively little is known regarding the precise ecological mechanisms. 30 Here, we provide the first evidence of widespread antiphage innate immunity in 31 environments using metagenomic sequence data from Antarctic hypolith 32 communities as model systems. In particular, immunity systems such as DISARM 33 and BREX are shown to be dominant systems in these communities. Additionally, we 34 show a direct correlation between the CRISPR-cas adaptive immunity and the 35 metavirome of hypolith communities, suggesting the existence of dynamic host-36 phage interactions. In addition to providing the first exploration of immune systems in 37 cold deserts, our results suggest that phages actively challenge niche communities 38 in Antarctic polar deserts. We provide evidence suggesting that the regulatory role 39 played by phages in this system is an important determinant of bacterial host 40 interactions in this environment.

42 Importance

43 In Antarctic environments, the combination of both abiotic and biotic stressors results in simple trophic levels dominated by microbiomes. Although the past two decades 44 45 have revealed substantial insights regarding the diversity and structure of microbiomes, we lack mechanistic insights regarding community interactions and 46 47 how phages may affect these. By providing the first evidence of widespread 48 antiphage innate immunity, we shed light on phage-host dynamics in Antarctic niche 49 communities. Our analyses reveal several antiphage defense systems including 50 DISARM and BREX, which appear to dominate in cold desert niche communities. In 51 contrast, our analyses revealed that genes, which encode antiphage adaptive 52 immunity were under-represented in these communities suggesting lower infection 53 frequencies in cold edaphic environments. We propose that by actively challenging 54 niche communities, phages play crucial roles in the diversification of Antarctic 55 communities.

56

58 Introduction

59 Antarctic terrestrial environments including open soils, permafrost and the surface and interior of rocks, are typically oligotrophic and dominated by 60 61 psychrophilic and psychrotolerant microbial communities (1-4). It has been 62 suggested that the extreme abiotic pressures of the environment such as 63 temperature, desiccation stress and UV radiation are dominant drivers of both the 64 diversity and function of cold-adapted bacterial communities in terrestrial polar 65 deserts (5-7). Similarly, biotic interactions such as competition, symbioses, horizontal 66 gene transfer (HGT) and predation have also been shown to play a role in the 67 distribution and diversity of microbial communities in these soil ecosystems (8-10). 68 The presence of viruses, including bacteriophages, in these cold hyper-arid desert 69 soils potentially adds an additional layer of complexity to the microbial system, but 70 the extent to which phage-host interactions play a role in shaping community 71 compositions and processes in cold desert soil niches remains a matter of 72 speculation (11, 12).

Antarctic desert hypolithic communities, in particular, have been shown to contain substantial viral populations, dominated by tailed bacteriophages of the order *Caudovirales* (11, 13-15). Micro-array analysis of lithic niches identified an even greater phage diversity, including signatures of RNA bacteriophages of the family *Leviviridae*, ssDNA phage of the family *Microviridae* and non-tailed dsDNA tectiviruses (16). Together, these observations suggest that phages may play an important role of in microbial community structures and functions.

The presence of active bacteriophages in a microbial community inevitably leads to the evolution of specialized bacterial defensive measures (17), and a diverse range of bacterial defense mechanisms against parasitic phages have been

83 identified (18, 19). These include adaptive immunity elements, such as the CRISPR-84 Cas systems, and innate immunity mechanisms, such as restriction-modification (RM) and toxin-antitoxin abortive infection (Abi) systems (18). Recent pangenomics 85 86 studies have also identified novel defense systems that are widely distributed across 87 bacterial taxa and are thought to play a role in anti-phage resistance (20-23). These 88 include the bacteriophage exclusion (BREX) system, coded by a 4-8 gene cluster, 89 that provides resistance to Siphoviridae and Myoviridae tailed phages by inhibition of 90 phage DNA replication (21), and other less well characterized systems such as the 91 Thoeris, Shedu and Gabija elements that increase bacterial host resistance to 92 specific groups of phages (22).

93 Combining the valuable evidence on phage diversity and prevalence in polar 94 desert soils, we hypothesize that phage-host interactions play an important role in 95 shaping the structure of edaphic microbial communities in these environments. To 96 test our hypothesis, we assess the known bacterial defense systems in 97 metagenomic sequence data derived from niche Antarctic hypolith community. We 98 were able to link some of these data to specific phage genomes and propose that 99 phages play an active role in shaping the immunity of Antarctic soil microbial 100 communities.

101

102 **Results**

The distribution of anti-phage defense mechanisms shows an abundance of
 innate immunity genes

105 The distribution of antiphage defense systems in the metagenome was 106 determined by mapping defense genes against the taxonomically assigned contigs. 107 In total, 24,941 defense genes were detected, compromising 1.2% of the entire

108 metagenome gene count. Approximately 40% of these were found in contigs 109 attributed to unknown phyla. The general distribution of defense genes across known 110 phyla was consistent with the relative abundance of each phylum in the metagenome 111 (Figure 1A, Table S4). Proteobacteria harbored the highest number of anti-phage 112 genes (5289 genes, 1.1% of total gene count for this phylum), followed by 113 Actinobacteria (3808, 0.9% total gene count) and Bacteroidetes (2128, 1.08% of total 114 contig count). RM, DISARM and BREX systems were the most abundant systems in 115 the metagenome, contributing 67.6% of the total gene hits for anti-phage defense 116 systems. On the other side of the spectrum, the defense systems Shedu, Hachiman 117 and CRISPR-type 2 were present at relatively low abundances, and therefore had 118 little apparent contribution to the global defense system distribution. The average 119 contribution of defense genes to the total gene count per phyla was 1.8%, with 120 Deferribacteres and Candidatus Tectomicrobia as outliers. However, it is important 121 to note that these phyla represent a very small portion of the metagenome, and 122 therefore the possibility that the high percentage of defense genes is biased toward 123 the low gene count for these phyla cannot be disregarded.

124 Analysis of the relative contribution of each defense system within each 125 phylum also showed that genes belonging to the RM, DISARM, and BREX systems 126 were the main contributors across the majority of phyla (Figure 1B). The recently 127 discovered Zorva system was predominantly represented in the phyla 128 Gemmatimonadetes, Bacteroidetes, Planctomycetes, Proteobacteria and 129 Verrumicrobia, while CRISPR systems showed the highest contribution in 130 Cyanobacteria and Euryarchaeota. Interestingly, non-canonical anti-phage systems 131 represented more than 50% of the defense systems identified for all phyla aside from Euryarchaeota, with Verrucomicrobia, Planctomycetes and Acidobacteria possessingthe highest distribution of non-canonical defense genes.

134

135 Innate immunity is dominated by BREX and DISARM genes

As highlighted above, anti-phage systems across phyla in the hypolith metagenome were dominated by non-canonical innate systems. Further analysis of the distribution of defense genes revealed that anti-phage systems in the majority of phyla were dominated by BREX and DISARM genes. The two systems together accounted for 33.4% of defense genes, compared to 31.7% genes belonging to canonical RM systems.

142 A total of 3758 genes for the DISARM system were identified. These included 143 the Class I marker gene drmD (449 counts, 11.9% of DISARM genes), which 144 encodes the SNF2-like helicase (23), as well as the Class II marker gene drmA 145 (1020, 17.1% of DISARM genes), which encodes a protein with a putative helicase 146 domain (23). Similarly, a total of 4598 genes representing all BREX types were 147 identified in the metagenome. Interestingly, the most abundant gene from this 148 system found in the metagenome, pqW (2640, 57.4% of BREX genes), which codes 149 for a serine/threonine kinase, is specific to the type 2 BREX system, also called the 150 Pgl system (21). By comparison, of the 7908 RM genes found in the metagenome, 151 the most abundant is hsdM (1423, 18% of RM genes), a type I DNA methylase 152 responsible for the protection of host DNA (24). In fact, more than 50% of RM 153 defense genes were attributed to type I RM systems.

The third non-canonical system representing more than 10% of the anti-phage defense systems in a subset of the phyla, the Zorya system, included a total of 2411 genes in the metagenome. The majority of these were homologous to the two genes

that make up a proton channel, *zor*A and *zor*B. This is a common feature in all types
of Zorya system and is thought to cause depolarization of the membrane upon
infection (22).

160

Type I CRISPR-Cas genes comprise the bulk of anti-phage adaptive immunity genes

163 In total, 2234 CRISPR-cas genes were identified in 1601 contigs by searching 164 for shared sequence similarities against the CDD database. A substantial proportion 165 of all classified CRISPR-cas loci (71.4%) belonged to type I CRISPR-Cas systems, 166 followed by type III (18.5%) and type II (10.2%) (Table S5). While the abundance of 167 Cas I-B loci sequences in the public databases suggests that the Cas-I mechanism 168 is the most common in both bacteria and archaea (20 and 30% of total CRISPR loci 169 (25), less than 3% of these loci were present in our composite metagenome (Table 170 S5, Figure 2). Surprisingly, CRISPR-cas loci linked to Types I-C and I-E were the 171 most prevalent, at 24.1% and 12.9% of classified CRISPR-cas loci, respectively. 172 Another subtype identified at higher relative abundances than previously reported 173 (25) was I-U, at 10.76% of classified cas loci. This subtype is characterized by the 174 marker GSU0054 domain, which was the fourth most abundant cas CDD overall 175 (108 occurrences) after cas4, cas1, and cas2.

176 **Phage presence in the niche community is correlated with the CRISPR arrays**

177 CRISPR arrays represent the history of infection by invading DNA (e.g. 178 phages, plasmids (26, 27), and a study of their composition and frequencies 179 provides insights into phage-host interactions in an ecological context (28). A total of 180 878 CRISPR arrays harboring 10,292 spacers were identified in the metagenome, 181 with an average length of 36 protospacers per array (Figure S1A). CRISPR array sizes ranged from 2 to 249, with the majority (83.5% of total arrays) falling between 2
and 18 protospacers per array (Figure S1B).

184 The distribution of CRISPR array sizes in the metagenome was compared to 185 data collected from a ground-water microbiome (29), to compare the array size 186 distributions from environments with potentially different phage-host dynamics (11). 187 The results show that CRISPR arrays in the hypolith metagenome exhibited a 188 smaller and narrower size range, compared to the ground-water community 189 metagenome (Figure 3). This suggests the existence of distinct phage infection 190 frequencies between the different environments; i.e., lower infection frequencies in 191 the cold edaphic community.

192 In addition to using the CRISPR array as a tool for understanding infection 193 history, the viral population in the Antarctic soil community was also assessed by 194 assembly of the metavirome. A total of 793 contigs was assembled from the 195 metagenomic sequence data using VirSorter (30). Taxonomic annotation of these 196 contigs, using a database of viral reference genomes (31), unambiguously assigned 197 645 of these as viral, 560 of which were further assigned to the order of tailed 198 phages *Caudovirales*. Within this order, the majority of contigs were assigned to 199 Siphoviridae (52%), followed by unclassified Caudovirales (14%), and viruses with 200 no assigned family (13%) (Figure S2). To access the correlation between the viral 201 contigs and the CRISPR arrays, spacers from the metagenome were matched to 202 both the VirSorter contigs and a set of contigs from environmental datasets (IMG/VR 203 (32), which allowed for the taxonomic assignment of 394 (3.8% of total number of 204 spacers) CRISPR-cas spacers (Figure S3). The resulting similarity network (Figure 205 4) showed that all 73 VirSorter phage contigs included in the network (red nodes) 206 matched to CRISPR-cas spacers (grey nodes), suggesting that a substantial fraction (11.3%) of the identified viral population had a history of infection *in situ* in the host
population, and may therefore be actively involved in shaping the adaptive immunity
of the microbial community. In addition, several distinct clusters showed matches
between a single VirSorter contig and several spacers, suggesting these viral contigs
are common infection agents.

212 Functional analysis using eggNOG showed the presence of genes that 213 facilitate infection such as genes that code for chitinases, which are involved in the 214 degradation of the protective biofilm (33), as well as a AntA/AntB antirepressor 215 gene, thought to be involved in phage anti-immunity (34) (Figure 5). In addition, the 216 eggNOG functional analysis of the 645 VirSorter viral contigs also revealed the 217 presence of genes contributing to phage virulence (Table S2), the most abundant of 218 which encode for methyltransferases, which are actively involved in the evasion of 219 the R.M systems (35). This result suggests the possibility of an evolutionary pressure 220 for the phages to develop evasion mechanisms against their hosts, which further 221 hints at active phage-host dynamics in these long-enduring Antarctic hypoliths.

222

223 Discussion

224 Due to the relatively simple trophic structures in cold desert systems, 225 including Antarctic soils, cryptic microbial communities are considered to be 226 important drivers of local ecosystem services (36). However, the extent to which 227 these communities are influenced by phages remains largely unexplored. Such 228 interactions may shape the diversification and community interactions in cold desert 229 systems. Qualitative surveys of Antarctic metaviromics have reported a high diversity 230 of viruses associated with microbial communities of open soils, and cryptic niches 231 (12, 13). Evidence, albeit limited, that Antarctic soil phages exist predominantly in a

lysogenic rather than lytic lifestyle (14), has led to suggestions that the functional role
of phages in this spatially restricted, water-constrained desert soil niche may be
limited (11).

235 The results presented in this study provide the first evidence of interaction 236 between phage and hosts in this psychrophilic edaphic environment. This is most 237 evident in the correlation between the metavirome of the hypolith community and the 238 CRISPR-arrays, which suggest the active evolution of the adaptive immune system 239 against local viral threats. This idea of community adaption to local phage threat is 240 further implied by the positive correlation between the CRISPR arrays and viruses 241 extracted from local soils. In fact, a previous study (37) has already suggested that 242 recruitment from surrounding soils plays an important role in the development of 243 hypoliths, and this might also be extended to the recruitment of phages from the 244 surrounding ecosystem. Another indication of active interaction between phage and 245 host is suggested by the presence of several methyltransferases in the 246 metagenome-assembled viral contigs, which are a hallmark of viral evasion against 247 native host RM systems (31, 35). Other genes found in this virome contigs include 248 genes specifically involved in the degradation of biofilm matrices and evasion against 249 RM systems, further suggesting that there is a complex network of interactions at 250 play between phages and their hosts in the hypolithic environment.

251

While the metagenomics data analysed in this study does not give a direct indication of the temporal scale of the phage-host interactions occurring in the hypolith, the short sizes of CRISPR-array sizes in the hypolith metagenome suggest a low frequency of infection. This low frequency is further hinted at when comparing the hypolith CRISPR-array sizes with those of a more fluid and homogenous

257 environment, where viral-host interactions are assumed to be a frequent occurrence 258 (29). Together, these results imply a model for viral-host interactions in hypoliths 259 that follows the 'static-step-static' development model suggested by Pointing et al. 260 (38), driven by the stochastic and intermittent nature of rain events in such water-261 limited ecosystems. A surprising result from this study is the prevalence of non-262 canonical innate immunity systems, the most prominent of which are the BREX and 263 DISARM systems. While these two systems have been shown to be widespread in 264 bacteria using a pan-genomic dataset (21, 23), the present study represent the first 265 evidence for the prevalence of these systems in ecological samples. As such, this 266 result implies that non-canonical innate immunity is more important for anti-phage 267 microbial community defense than previously thought and should therefore be the 268 focus for future studies into innate immunity in the ecological context. There are also 269 indications from the hypolith metagenome that the prevalence of non-canonical 270 innate immunity over traditional RM and Abi system for defense against phages is 271 related to the adaptation of the hypolith communities to specific local viral 272 populations. For instance, the Zorya system, the third most prevalent non-canonical 273 immunity system in the metagenome, is hypothesized to operate similarly to the Abi 274 system (22). In turn, Zorya systems provide resistance against a limited range of 275 phages, including the ssDNA family *Microviridae* (22), which has been shown to be 276 prevalent in Antarctic aquatic and soil niches (39).

277

278 Conclusion

Together, these results are not consistent with the suggestion that the constraints of the environment, such as low temperatures, low a_w and resulting very limited capacity for inter-particle diffusion, lead to extremely localized phage-host

282 interactions (11). Rather, the data are suggestive of a dynamic and continual 283 interaction between host and phage. Nevertheless, inter-particle communication and 284 exchange may be limited to brief periods when bulk liquid water is present, after 285 snow melt, for example. Furthermore, the low metabolic rates (the inevitable 286 consequence of Arrhenius effects (temperature dependence of reaction rates) in cold 287 environments) should also limit the rates at which phages can replicate and 288 propagate, further limiting the frequency of interactions with their hosts (40). We 289 suggest that the localized nature of host-phage interactions in the hypolith niche and 290 the limited inter-particle communication, where bacterial hosts are not frequently 291 challenged by novel phage threats, leads to a reliance of microbial communities on 292 innate immunity as the primary defense against phage infection. The smaller sizes of 293 CRISPR arrays in the Antarctic soil metagenome sequences compared to those from 294 a temperate aquatic environment, and the under-representation of CRISPR systems, 295 give further credence to the temporally sporadic interaction between phages and 296 their hosts. Nevertheless, the correlation between the metavirome and the CRIPR-297 cas arrays, together with the presence of bacteriophage evasion genes in the 298 metavirome, suggest that phage-host interactions within the hypolith community are 299 a dynamic process that leads to co-evolution of both phages and hosts. We therefore 300 suggest that phages play a hitherto underestimated role in driving the evolution of 301 Antarctic soil microbial communities by shaping their collective immunity.

302

303 Materials and Methods

Sample collection, DNA extraction and metagenomic sequencing

305 The sample collection, DNA extraction and metagenomic sequencing 306 protocols used in this study have been described previously (41). Briefly, a total of 50 307 samples were collected from hypolithic niches in the Antarctica Miers Valley (GPS 308 78°09'36.0"S 164°06'00.0"E) and stored in sterile Whirl-Pak bags (Nasco 309 International, Fort Atkinson, WI, USA) at -20 °C. Metagenomic DNA was extracted 310 from each sample using a PowerSoil DNA isolation kit (MO BIO, Carlsbad, CA, 311 USA), and the purified DNA was pooled before further processing. Purified DNA was 312 sheared into fragments of approximately 300 bps and further purified from 1% 313 agarose gels. Subsequent sequencing was performed using Illumina HiSeq-2000 314 paired-end technology (2 x 101 bp), and the resulting reads were trimmed and 315 assembled as described below.

316

317 Metagenome assembly and taxonomical annotation

Metagenomic DNA sequence data were quality-filtered by trimmomatic version 0.36 using a phred cut-off > 30 (42). The assembly of high-quality reads from the metagenome sequence dataset was conducted using the IDBA-UD tool (43) and contig lengths were extended (scaffolded) using SSPACE Basic (43). The statistics for the assembly of the metagenome are presented in Table S1. Contigs were taxonomically assigned using the MEGAN v6 pipeline (44) with the NCBI taxonomy database for taxon ID assignment.

325

326 **Detection of the innate and adaptive defense systems**

Metagenomic contigs were used for functional gene predictions using prodigal v2.50, with the –meta parameter implementation (45). Predicted genes were subsequently screened for domain similarity with known defense systems against 330 the conserved domain database (CDD) of clusters of orthologous groups (COGs) 331 and protein families (Pfams) using rps-blast (E value < 1e-02) (33). These results 332 were manually filtered for the identification of phage-specific defense systems, which 333 include restriction-modification (R.M), bacteriophage exclusion (BREX), abortive 334 infection (Abi), defense island system associated with restriction-modification 335 (DISARM), and other recently identified systems using a refined list of COG and 336 Pfam position-specific score matrices (PSSMs) for marker genes in these systems 337 (21-23, 46). A list of the marker genes used in this study can be found in Table S2. 338 Additionally, defense genes that could not be clustered into a specific system were 339 classified as ambiguous as were not considered for subsequent analysis (Table S3).

ORFs predicted using prodigal v2.50 were queried against the CDD database for the presence of putative CRISPR-cas genes (47), using delta-blast at a cutoff E value of 1e-03. Multi-gene cas modules were identified as those having multiple cas annotated genes with \leq 5 ORF spacings. Type and subtype classifications were assigned following the updated classification set by Makarova et al. (25).

345

346 **Phage genome identification and CRISPR spacer matching**

347 Antarctic hypolith phage genomes were identified from the assembled 348 metagenome using VirSorter (30) on the iVirus platform hosted by Cyverse (48), 349 using the virome database and the microbial decontamination option. Only 350 predictions of categories 1, 2, 4 and 5 were used (phages and prophages identified 351 with the "pretty sure" and "quite sure" qualification). Additional phage environmental 352 phage contigs were downloaded from the IMG/VR database version 2018-07-01 4 353 (32) and used for the network construction. Taxonomic assignment of assembled 354 contigs was performed by using the DIAMOND blastx function with a viral database downloaded from the NCBI Viral Genomes Resource and e-value set to 1e-5. ORFs of VirSorter contigs were predicted using Prodigal v2.50 (31, 49) with the virus genomes setting and annotated using eggNOG-mapper v1 (50) with the DIAMOND option and the EggNOG v4.5.1 database (51). Annotation were visualized with the ApE v2.0.55 plasmid editor (http://jorgensen.biology.utah.edu/wayned/ape/).

360 The CRISPR recognition tool (CRT) v1.2 was used with the default settings to 361 search for CRISPR arrays in the hypolith metagenome (52). The identified spacers in 362 the arrays were matched with the VirSorter phage database and the IMG/VR 363 database using blastn of the BLAST+ suite with the following parameters: -364 gcov hsp perc 80 -task blastn -dust no -soft masking false (53). Spacer matches of 365 > 90% sequence identity for the VirSorter genomes and >95% identity for the 366 IMG/VR genomes were exported and visualized as a network in Cytoscape (54), 367 where the nodes are spacers (grey) or genomes (blue = IMG/VR; red = VirSorter) 368 and the edges blastn matches.

369

371 Acknowledgements

We are grateful to the National Research Foundation (NRF) (Grant ID 118981, the South African National Antarctic Programme (SANAP 110717), and the University of Pretoria for funding. TPM also wishes to acknowledge the Fulbright Visiting Scholar Program for providing sabbatical funding. EMA gratefully acknowledges the support of the Biotechnology and Biological Sciences Research Council (BBSRC); this research was funded by the BBSRC Institute Strategic Programme Gut Microbes and Health BB/R012490/1 and its constituent project(s)

379 BBS/E/F/000PR10353 and BBS/E/F/000PR10356

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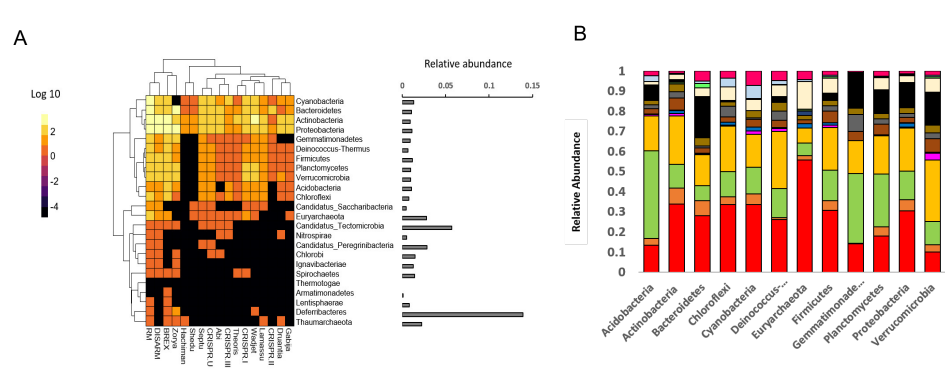
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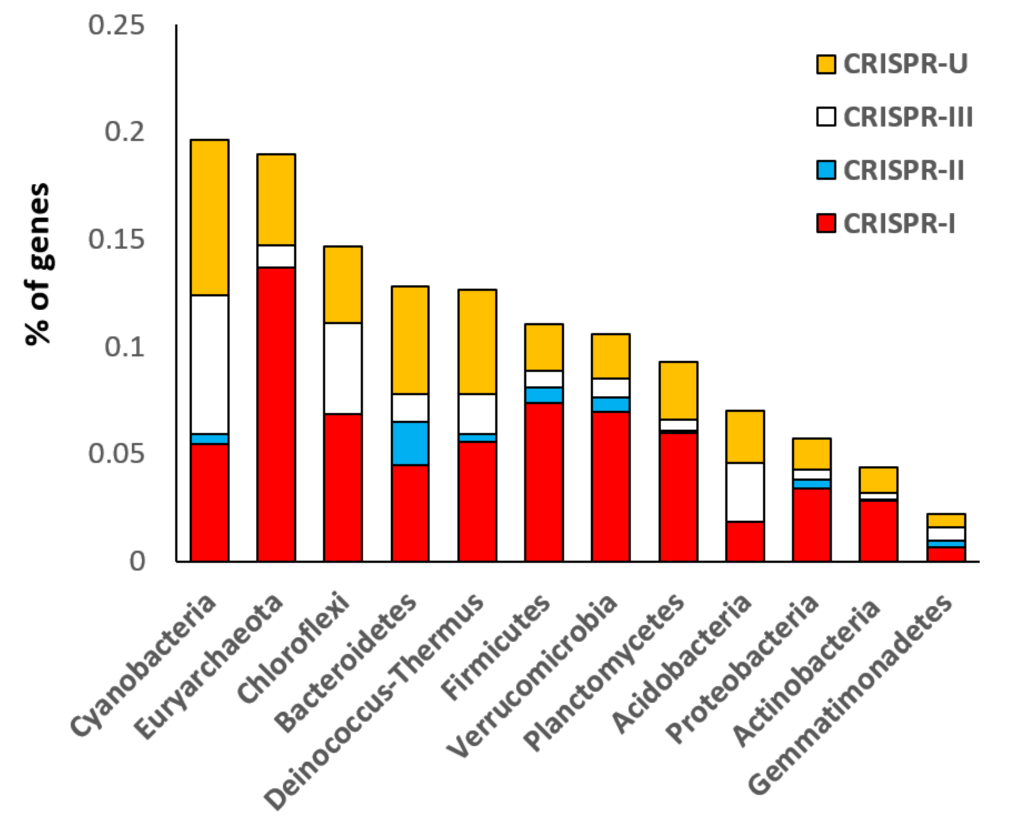
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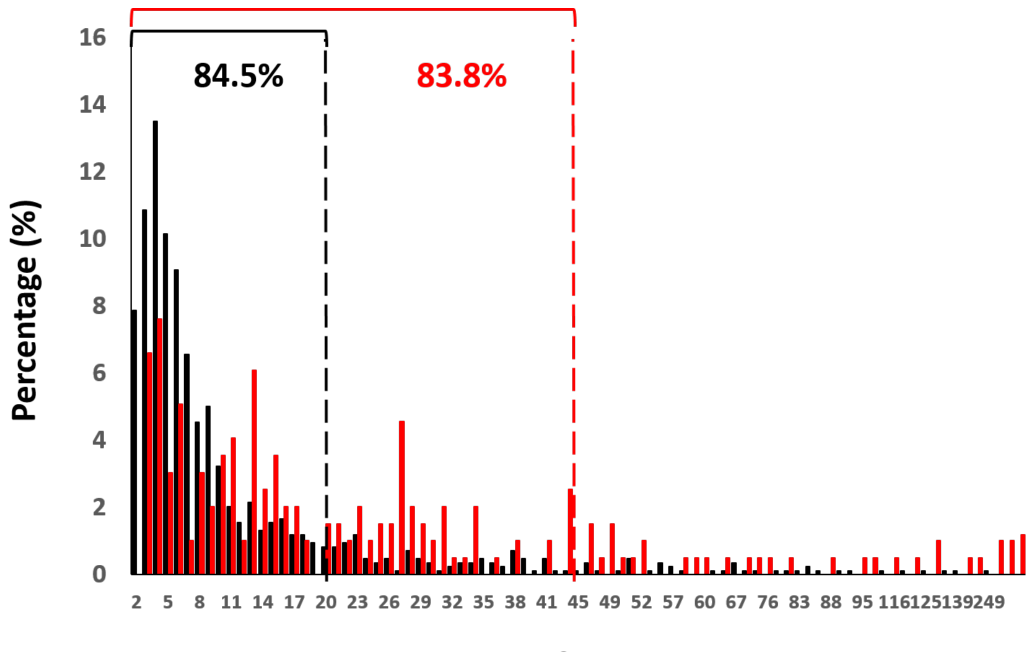
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CRISPR-U CRISPR-III CRISPR-II CRISPR-I Zorya Theoris Shedu Septu Lamassu Wadjet Hachiman 🗖 Gabija Druantia DISARM BREX 🗖 Abi RM





Size of array

