

Biology of archaea from a novel family Cuniculiplasmataceae (Thermoplasmata) ubiquitous in hyperacidic environments

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1	Biology of archaea from a novel family Cuniculiplasmataceae				
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- 2
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- 30

31 ABSTRACT

The order *Thermoplasmatales* (*Euryarchaeota*) is represented by the most acidophilic organisms known so far that are poorly amendable to cultivation. Earlier culture-independent studies in Iron Mountain (California) pointed at an abundant archaeal group, dubbed 'G-plasma'. We examined the genomes and physiology of two cultured representatives of a Family *Cuniculiplasmataceae*, recently isolated from acidic (pH 1-1.5) sites in Spain and UK that are 16S rRNA gene sequenceidentical with 'G-plasma'.

39 Organisms had largest genomes among Thermoplasmatales (1.87-1.94 Mbp), that 40 shared 98.7-98.8% average nucleotide identities between themselves and 'G-41 plasma' and exhibited a high genome conservation even within their genomic 42 islands, despite their remote geographical localisations. Facultatively anaerobic 43 heterotrophs, they possess an ancestral form of A-type terminal oxygen reductase 44 from a distinct parental clade. The lack of complete pathways for biosynthesis of 45 histidine, valine, leucine, isoleucine, lysine and proline pre-determines the reliance 46 on external sources of amino acids and hence the lifestyle of these organisms as 47 scavengers of proteinaceous compounds from surrounding microbial community 48 members. In contrast to earlier metagenomics-based assumptions, isolates were 49 S-layer-deficient, non-motile, non-methylotrophic and devoid of iron-oxidation 50 despite the abundance of methylotrophy substrates and ferrous iron *in situ*, which 51 underlines the essentiality of experimental validation of bioinformatic predictions.

53 Introduction

3

54

55 Acidic environments are widely distributed across the globe and are represented 56 by natural (e.g. volcanic or geothermally heated), or man-made (mines or acid 57 mine drainage (AMD)) sites, with a constantly low pH¹. Microbial communities 58 inhabiting such niches were considered to be of a relatively low complexity², 59 however, recent OMICS studies pointed at a greater variety of yet uncultured 60 prokaryotes¹. Due to the low numbers of cultured microorganisms that may serve 61 as a functional reference, their physiological features and hence the roles in the 62 environment largely remain at the level of *in silico* predictions from metagenomic data. In that context, while a certain success has been achieved in isolation of new 63 64 bacterial taxa from these specific environments³, only a handful of cultured, 65 taxonomically described and physiologically studied archaeal representatives have 66 been obtained³. Recent data based on genomes assembled from metagenomes documented a number of archaeal clades mostly affiliated with the order 67 *Thermoplasmatales*, phylum *Euryarchaeota*⁴. Among archaeal populations from 68 69 the above order, cultured members of Ferroplasmaceae together with yet 70 uncultured archaea from the so-called 'alphabet plasmas' were the most abundant 71 and hence suggested to play important roles in carbon cycling in the environment⁵. 72 Initially identified in 16S rRNA gene clone libraries from Iron Mountain⁶, these 73 archaea have later been found in a number of acidic environments of different temperature regimes¹. Their presence in iron-rich environments have guite logically 74 promoted discussions on their iron oxidation potential. Apart from the iron oxidation 75 76 experimentally confirmed only in cultured members of *Ferroplasmaceae*^{7,8}, other 77 *Thermoplasmatales* were described as facultatively anaerobic heterotrophs⁹. Their

78 appearance in biofilms alongside with chemolithoautotrophs suggests that the 79 metabolism of this group may be depended on organic compounds (sugar 80 polymers/oligomers, peptides, lipids or carbohydrate monomers) derived from primary producing organisms¹⁰. "Alphabet plasmas" were furthermore predicted to 81 oxidise carbon monoxide and utilise methylated compounds⁴. However, the dearth 82 83 of experimental evidence has largely limited our entire current understanding on metabolism, physiology, and environmental roles of these archaeal lineages. 84 85 One of important members of *Thermoplasmatales* in AMD systems from Iron 86 Mountain (California) was a group of organisms dubbed 'G-plasma', which was so 87 abundant, that the genomes of some of the representatives were almost fully 88 assembled^{2,4}. These organisms were the third-abundant community members 89 (following Leptospirillum spp. "Group II and III") and contributed up to 22 % of total 90 community proteome¹¹. Elsewhere, 'G-plasma' contributed approx. 15 % of total metagenomic reads in this environment¹². However, despite their abundance and 91 92 ubiquity these organisms escaped the cultivation until very recently, when first 93 representatives were isolated from Cantereras AMD site (Spain) and Parys Mountain/Mynydd Parys (Anglesey, UK) and described as representatives of a 94 95 novel family, Cuniculiplasmataceae, new genus and species Cuniculiplasma divulgatum within the order Thermoplasmatales¹³. The 16S rRNA gene sequences 96 of isolates PM4 and S5 were identical to those from 'G-plasma' cluster from 97 98 metagenomic data analysed at Richmond mine at Iron Mountain, USA^{2,4}, terrestrial acidic springs in Japan¹⁴, high-temperature fumarole and acidic biofilm from 99 100 Mexico (GenBank Acc Nrs. JX997948, AB6000334 and KJ907759), Frasassi hydrogen sulphide-rich cave system, Italy¹² and from AMD system, Los Rueldos, 101 Spain^{1,10} and other low pH systems. Altogether, these documentations point at the 102

- 5
- 103 ubiquity of *Cuniculiplasmataceae*-related organisms in acidic systems and volcanic
- areas (Fig. 1, Supplementary Table S1).
- 105 New isolates provided a great opportunity to perform for a first time the
- 106 comparative genomic analysis of very closely related members of *Thermoplasmata*
- 107 from very distant geographic locations, to analyse their physiology and functions
- related to the environment in the context of the earlier genomic predictions, and,
- 109 finally, to analyse their evolutionary relationships with other clades within the class
- 110 *Thermoplasmata*, which harbours organisms known as acidophilic 'champions'^{9,15}.
- 111

112 **Results**

Physiological traits: *in silico* predictions in 'G-plasma' vs experimental data *lron oxidation.*

- 115 Despite earlier suggestions of iron-oxidising capabilities based on the occurrence
- 116 of rusticyanin/sulfocyanin-encoding gene homologs⁴, no iron oxidation was
- 117 confirmed with ferrous sulfate and pyrite in either *C. divulgatum* isolate.
- 118 Noteworthy, the presence of genes for rusticyanin/sulfocyanin homologs might not
- 119 necessarily be connected with the iron oxidation in archaea of the order
- 120 Thermoplasmatales, e.g. Picrophilus torridus does not oxidise ferrous iron despite
- 121 the presence of sulfocyanin. It was suggested¹⁶ that this respiratory complex in *P*.
- 122 torridus is situated on a genomic island, which seems also to be the case for
- 123 rusticyanin/sulfocyanin genes acquired by a lateral transfer in *C. divulgatum* S5 (s.
- 124 below) and *F. acidiphilum* (Golyshina *et al.*, in preparation).
- 125
- 126 Archaeal flagella and pili

127 In 'G-plasma', the full operon encoding FlaBCDEFGHIJ with individual proteins being homologous to those from Methanococcus voltae and Halobacterium 128 salinarum has been reported earlier⁴, however, corresponding loci have not been 129 130 detected in either genome of Cuniculiplasma divulgatum. Our analysis suggested that *M. voltae* and *H. salinarum* flagellar proteins do not have significant (e-values 131 132 <0.01 and guery coverage > 50%) BLAST hits in 'G-plasma' in silico translated proteome. Electron microscopy of C. divulgatum grown under optimal conditions did 133 not provide evidence for an archaellum, but occasionally showed the presence of 134 distinct pili¹³, (s. also Fig. 2c). This feature is also reflected in the genomic data of 135 136 both isolates of Cuniculiplasma, as discussed further (s. subsection 'Secretion 137 system and motility').

138

Regarding the S-layer prediction in 'G-plasma', corresponding genes to be linked 139 with S-layer formation⁴ were also found in both *Cuniculiplasma* genomes. These 140 genes annotated in 'G-plasma' to code for "S-layer protein P. torridus"⁴ (and 141 142 annotated as a surface protein in *P. torridus* itself), are affiliated with COG3391, arCOG0652 and arCOG2560 that have five homologs in each Cuniculiplasma 143 144 divulgatum genome). Additionally, genes encoding oligosaccharyltransferase AgIB in 'G-plasma' are present in both genomes of *Cuniculiplasma* strains, as well. 145 146 However, as revealed by electron microscopy, the cells of strains S5 and PM4 147 were only surrounded by cytoplasmic membranes and lacked distinct (predicted in 'G-plasma') S-layers (Fig. 2 a, b). An S-layer should provide a certain rigidity to 148 149 cells and its absence is consistent with the characteristic pleomorphism in C. 150 divulgatum, as exemplified in Fig. 2 c,d. Apparently, within the order Thermoplasmatales, the cell wall-deficient members clearly outnumber S-layer-151

exhibiting organisms, which are represented only by *Picrophius* spp.⁹. This feature
is also reflected in the genomic data of two strains, as discussed further.

154

155 *Methylotrophy*

In the growth experiments performed with both strains of C. divulgatum with a 156 range of methylated compounds¹³ we were not able to confirm the methylotrophy 157 earlier predicted in 'G-plasma'⁴. In this regard, the genes predicted to be present in 158 159 'G-plasma', namely methenyl tetrahydrofolate cyclohydrolase and formyl-160 tetrahydrofolate synthetase have also been found in both Cuniculiplasma 161 genomes. However, the gene encoding 'methanol dehydrogenase' in 'G-plasma' 162 has not been confirmed in Cuniculiplasma. Furthermore, the protein referred as 163 such in 'G-plasma' itself had a low amino acid sequence identity (>26%) to alcohol dehydrogeneases of unknown substrate specificity and was equally (dis)similar 164 165 with maleylacetate reductases. The very homolog was found in the S5 genome, 166 but not in PM4. Among tested substrates, e.g. methylamines, could not be utilised 167 by Cuniculiplasma isolates since no genes for methylamine dehydrogenase or dimethylamine and trimethylamine dehydrogenase were found. Whatever the case, 168 169 methylotrophy was not experimentally confirmed in any *Thermoplasmatales*, even though the methanol is a common product of organic matter degradation and may 170 171 be available in studied environments.

172

173 Genome analysis of *Cuniculiplasmataceae*

174 Genome statistics

175 The genomes of *C. divulgatum* strains (Table 1) are larger as compared to the

176 relatives from *Thermoplasma* spp (1.58 Mbp for *T. volcanium* and 1.56 Mbp for *T.*

- 8
- 177 *acidophilum*) and *Picrophilus torridus* (1.55 Mbp), being within the common range
- to archaea of the family *Ferroplasmaceae* (1.94 Mbp for *"Ferroplasma*

acidarmanus", 1.75-1.78 Mb for Acidiplasma aeolicum and 1.74 Mbp for A.

180 *cupricumulans*). Low G+C contents of genomic DNA of strains S5 and PM4 are

- 181 rather typical for *Thermoplasmatales*¹⁷.
- 182

183 Genome comparisons

184 The three genomes exhibited a high average nucleotide identity (ANI)¹⁸ and

average amino acid identity (AAI)¹⁹, which also supports similar physiological

patterns in both isolates: strains S5 and PM4 had 98.8 % ANI, while ANI of both

isolates with 'G-plasma' genome were about 98.7 and 98.4 %, respectively,

188 pointing at their similar evolutionary trajectories despite transcontinental

189 localisation of their niches and highly complementary microbial structure and gene

190 pools in AMD settings¹ (also s. Fig. S1 for the AAI data).

191 The core *in silico* proteome of *C. divulgatum* strains and 'G-plasma' is represented

192 by 1174 protein groups. 111 protein clusters were identified as exclusively

distributed among PM4 and S5 strains, 13 among PM4 and 'G-plasma' and 27

among S5 and 'G-plasma' (Fig. 3, 4 and Fig. S1, Supplementary Table S2). 79, 52

and 114 unique single-copy genes and 1, 1 and 10 strain-specific paralogue

196 clusters were identified for S5, PM4 and 'G-plasma' respectively. Analysis of their

197 distribution across the chromosomes revealed that most of them are highly

198 clustered (Fig. 4), supporting the hypothesis that LGT (lateral gene transfer) is an

¹⁹⁹ important driving force in evolution of AMD-related microorganisms²⁰, however with

200 very similar patterns of foreign DNA integration in the genomes of recipients.

202 Lateral gene transfer (LGT), genomic islands (GIs) and defence systems 203 Analysis of arCOG distribution within variable and core parts of Cuniculiplasma-204 related in silico proteomes revealed a significant enrichment in "Defense 205 mechanisms" group in PM4 strain. This observation together with the fact that PM4 possesses 92 non-redundant CRISPR spacers as opposed to 52 in S5 strain and 206 207 only 10 in 'G-plasma' give an opportunity to speculate that Parys Mountain/Mynydd Parys mine is characterised by much higher viral load than other investigated acid 208 mine habitats²¹. In turn, unique and accessory part of 'G-plasma' genome 209 210 characterised by the lowest proportion of 'defence mechanisms' is highly enriched 211 with 'replication, recombination and repair' proteins including integrases, 212 transposases and recombinases pointing on higher level of genome mobility in 'G-213 plasma' (Fig. 3). Another point related to arCOG distribution is the prevailing comparative number of unique strain-specific proteins in S5 for categories energy 214 215 production and conversion, cell cycle control, transcription, inorganic ions transport 216 and metabolism (Fig. 3). 217 Lateral gene transfer (LGT), genomic islands (GIs) and defence systems. 218 The strain S5 harboured ten GIs in its genome, whereas its counterpart from 219 Parys Mt/Mynydd Parys only four (Fig. 4 (a, b) and Supplementary Table S2). As expected, numerous insertion sequences elements (IS), integrases and 220 221 transposases from different families (IS3, IS5, IS6, IS66, IS256, IS200/605, IS110, 222 IS1634) were associated with the GIs, as well as tRNAs reflecting the commonality of tRNA co-occurrence in genomic islands²². The G+C molar content in predicted 223 GIs varied within the range 37.7 - 43.2 %, i.e. marginally higher than average 224 values in PM4 and S5 genomes (Supplementary Table S3), which may be a result 225 of old integration events and consequent DNA amelioration in GIs making GC 226

227 similar to that in the core genomes. Notably, a slight difference between G+Ccontent in genomes of S5 and PM4 strains (37.16% in PM4 vs 37.30 in S5) is 228 229 determined by the presence of six additional GIs in the former isolate. Analysis of 230 taxonomic affiliation of GIs revealed that almost all lateral transfers originated from other acidophilic euryarchaea. This observation implies the existence of a highly 231 232 mobile gene pool in acidophilic Archaea, which determines rapid adaptations of Thermoplasmatales members to toxic concentration of heavy metals and to a high 233 234 viral load.

Thus, some GIs could clearly be attributed to 'defence' islands, e.g. GI3, GI7 and GI9-10 in S5 and GI4 in PM4 due to the localisation therein of genes for restrictionmodification and toxin-antitoxin systems. Others (e.g. GI 4, GI 5 and GI 8 from S5) were transport-, efflux-, metal- and oxidative stress response-related). GI1 from the strain S5, which is absent in PM4, harboured an array of genes for site-specific recombinases, metal-transporting ATPases, multipass membrane proteins,

241 metallochaperones, cupredoxin COX2 family proteins, heavy metal reductases,

and rustycyanin/sulfocyanin homolog.

We have identified several toxin-antitoxin systems (TAS) -encoding genes, mostly 243 associated with GIs in both isolates. The most abundant ones were represented by 244 245 vapBC of the type II system: six clusters of corresponding ORFs in PM4, and 246 seven in S5 and, besides three vapB toxin genes were found across chromosomes 247 in both *Cuniculiplasma* isolates. In addition, three clusters of genes were found in PM4 and two such loci in S5 with corresponding MazE and MazF family proteins 248 249 affiliated with COG2336/arCOG03943 and COG2337, respectively. 250 Furthermore, three and two re/EF loci were identified in PM4 and S5 genomes,

correspondingly. Commonly, TAS are known to be stress response-connected and

lateral gene transfer-related^{23,24}, which is confirmed by the GI analysis. Notably, no
 TAS were previously reported in *Thermoplasmatales*²⁵.

All genomes of *Cuniculiplasma* spp. showed the presence of Clustered Regularly
 Interspaced Short Palindromic Repeats (CRISPR)-Cas defence systems: in S5, we

have identified the cluster of genes for Cas3, Csx17, Cas7, Cas5, Cas4/Cas1 and

257 Cas2 with an adjacent CRISPR repeat region with 57 spacers. Interestingly, all

258 proteins exhibited 100% polypeptide identity with counterparts from 'G-plasma'

259 (apart from Cas4 and 1 which had psi-blast hits of about 54% identity with

acidobacterial polypeptides).

ATDV01000019 contig of 'G-plasma' exhibited a remarkable similarity in gene

arrangement (ADMU5_GPLC00019G0101-G0107) with the corresponding region

in S5 (CIP_1636-1642) albeit with only 10 spacers of repeats found on the

terminus of the contig ATDV01000011. According to²⁶ systems from 'G-plasma' and

265 S5 can be classified as Type I-C. The strain PM4, in contrast to the above, coded,

in this order, for Cas6 endoribonuclease, Cas8b, Cas7, Cas5, Cas3, Cas4, Cas1

and Cas2, flanked by a repeats-spacers array of 92 spacers, suggesting its

affiliation with the Type I-B system²⁶. Interestingly, all sequences of Cas proteins

were equally distant (28-57% sequence identity (Supplementary Table S4) with the

270 proteins from 'F. acidarmanus' and other archaea and, to the same extent, with

271 polypeptides from representatives of *Bacteria*, e.g. ∂-Proteobacteria or

272 Acidobacteria (pointing at an unclear origin of corresponding gene clusters).

273 Remarkably, very similar pseudogenes CPM_1008 and CSP5_0996 for Cas1 were

detected in both isolates, in similar locations on chromosomes, within the same

275 genomic context in the region severely affected by transposon integration and

pseudogenisation. Analysis of CRISPR repeats in S5, PM4 and 'G-plasma' by

277 blastn-short algorithm revealed no cross-matches of spacers between these three genomes. Nevertheless, eight of 92 PM4 repeats and four of 57 S5 spacers 278 279 showed high (90-100%) identity with sequences of Richmond mine microbial and viral communities^{21,27}, suggesting the existence of viruses common for these 280 extreme acidic ecosystems. Interestingly, CRISPR array of PM4 contains two 281 282 spacers with the significant level of similarity (83 and 96%) to marine metagenomic sequences (Supplementary Table S5). Despite a significantly high probability of 283 284 false positive hits (e-values are 0.015 and 0.14, respectively), this finding might be 285 speculated as relic genomic signatures of an ancient hydrothermal ecosystem which existed 480-360 my BP in the place of contemporary Parys Mountain site²⁸. 286 287 From the analysis of GIs in *Cuniculiplasma* spp. two important facts become 288 apparent. First, the co-occurrence of GIs and the majority of 'unique' genes (numbers in the outermost segments in Fig. S1 and green lines in Fig. 3). Most 289 'unique' genes had likely been acquired from organisms other than 290 291 Thermoplasmata and had no hits above the e-value cut-off (0.005) either with 292 'alphabet plasmas' or with isolates from cultured/genome-sequenced 293 Thermoplasmata, suggesting a high probability of lateral gene transfer also in the 294 vicinity of GIs. Second, a remarkable similarity in gene arrangements was observed within some GIs in both strains and their positioning in both 295 296 chromosomes, i.e. in 'defence islands' GI9-10 of S5 and GI4 of PM4 (homologous 297 to 'G-plasma' contig ADMU5 GPLC00019G0004-G0013) (Supplementary Fig. S2 298 (b) and Supplementary Table S6) and GI2 of S5 and GI1 and 2 from PM4 that were 299 mostly composed by ORFs for hypothetical proteins conserved in both organisms 300 (Supplementary Fig. S2 (a)). Such conservation in gene arrangements in GIs is indicative for an important role these genes may play in metabolism in iron-rich 301

environments and that they can relatively easily be transferred between organisms
 and remain in genomes due to the selective pressure, providing a competitive
 advantage, much like 'catabolic transposons' for xenobiotics or hydrocarbon
 metabolism²⁹. This was the case in, e.g., three transposases-adjacent operons in
 strain S5 encoding metallochaperone and metalloreductases that showed high
 similarities with counterparts in all *Thermoplasmatales* type strains.

308

309 Secretion systems and motility

310 In the genomes of both strains PM4 and S5 no operon essential for archaella biogenesis (flaCDFGB)³⁰ was found, and consistently, no archaella and no motility 311 312 were observed by microscopy, despite earlier suggestions^{4,13}. The strain PM4 exhibited filaments or pili-like cell surface structures¹³, according with the presence 313 in genomes of genes for proteins of type IV pili biosynthesis. In accordance with 314 the recent census of archaeal clusters of orthologous groups of proteins (arCOG) 315 related with pili formation³¹, we have identified principal components in both 316 317 genomes as follows. In S5, CSP5 0712 and CSP5 0715 encoded Type II secretion system ATPase subunits (Flal, arCOG01817) forming a gene cluster with 318 319 genes for CSP5 0713-14, encoding homologs of flagellar assembly proteins J2 and J1 (TadC, arCOG01808) and major pilins (FlaB/FlaF/PilA family, arCOG02423) 320 coded by clustered CSP5 1254-1255 and stand-alone CSP5 0804 and 321 322 CSP5 0881. The arrangement of two gene clusters harbouring six former gene loci resembled that in both genomes of "Aciduliprofundum" spp.³¹. In the strain PM4, 323 corresponding loci were CPM 0710 and 0713 (secretion ATPases), CPM 0711-324 0712 (TadC-like proteins) and CPM_1256-57, 0800 and 0878 (major pilins), with 325 the very same arrangement of gene clusters across the chromosome, as in the 326

327 strain S5. Function of these surface formations could be various: surface adhesion, intercellular connection, DNA exchange or probably attachment to the substratum 328 rather than the motility³². Both strains encode type IV secretion components: 329 TraG/TraD/VirD4 family ATPases (arCOG04816) by CSP5 0791 and CPM 0795; 330 membrane protein (arCOG05340), VirB4 component (arCOG04034), multipass 331 332 protein (arCOG05369) and membrane protein (conserved in Thermoplasmatales only) with four latter encoded by gene clusters CSP5 1185-1189 and CPM 1190-333 93. Furthermore, both genomes encode Sec translocon components, preprotein 334 335 translocase subunits SecYE and Sec61beta, signal peptide peptidase and signal 336 recognition particle subunits and receptors. Another feature to be addressed here 337 is the presence of Sec-independent Tat pathway genes for folded proteins' 338 secretion. Twin-arginine translocase subunits A and C are presented in PM4 and 339 S5 genomes, which may be functional in an analogy with a Gram-positive bacterial Tat system, known to work without additional TatB protein³³. 340

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342 Peptidases, peptide/amino acids transporters

Consistently with the substrate preferences for proteinaceous compounds, each 343 344 genome contained more than 50 various peptidases. Among those, eight were 345 predicted to be secreted due to the presence of signal peptides. Five peptidases 346 were most probably responsible for extracellular hydrolysis of proteins and 347 peptides: three serine peptidases of S53 family and two thermopsins, aspartic peptidases of A5 family³⁴. S53 family peptidases have 3D structures similar with 348 349 other representatives of SB clan, their distant homologs, subtilases of S8 family, 350 but differ in acidic pH optima for activity. Since all Thermoplasmatales are 351 extremely acidophilic microorganisms, it is quite logic that S8 peptidases-coding

352 genes were not found in their genomes, and were 'replaced' by S53 peptidases. A thermopsin, also characterised as an acidic endopeptidase³⁵ is another reflection 353 354 of adaptation of *Cuniculiplasma* spp. to extremely acidic conditions. The genomes 355 of the strains S5 and PM4 encoded two almost identical thermopsins, however, one of S5 thermopsins lacked 130 amino acid on its N-terminus and hence lacking 356 357 secretion system motifs. Genomic context analysis revealed the presence of various transporters in close vicinity of A5 peptidases of both strains and almost no 358 359 transporters in S53 neighbourhood. Among transporters, surrounding thermopsins, 360 the most probable amino acid and peptides importers were among the members of 361 Major Facilitator Superfamily (MFS, 2.A.1), according to TCDB database³⁶.

362

363 TCA

364 All genes, coding for TCA proteins were clearly identified in *Cuniculiplasma*

365 genomes except 2-oxoglutarate dehydrogenase (EC 1.2.4.2) and fumarate

366 reductase (EC 1.3.5.1). A 2-ketoacid dehydrogenase complex was found

367 (CSP5_0253-0256 and CPM_0219-0222), however it was related to rather 2-

368 oxoisocaproate dehydrogenase (EC 1.2.4.4) than to 2-oxoglutarate

dehydrogenase (EC 1.2.4.2) or pyruvate dehydrogenase (1.2.4.1). Still, the

370 conversion of 2-oxoglutarate to succinyl-CoA could be catalyzed by 2-oxoglutarate

371 synthases (CSP5_0284-0285 and CPM_0255-0253) and CSP5_1378-

372 1379/CPM_1377-1378). These ferredoxin-dependent enzymes are known to be

373 highly sensitive to oxygen, thus, presumably being active during anaerobic growth

of *C. divulgatum* or being highly stable to oxygen as it was shown for a homolog

from *Mycobacterium tuberculosis*³⁷. CSP5_1895 and CPM_1834 (COG1027) are

homologous to several characterised class II aerobic fumarases (EC 4.2.1.2),

377 however the phylogenetic analysis shows (Supplementary Fig. S3) their marginally closer relatedness with aspartases (EC 4.3.1.1) than with fumarases (yet with high 378 379 AA identity/similarity values (38/57%) with the Class II fumarase from Sulfolobus 380 sp.). Whatever the case, a possible absence of fumarase would imply incompleteness of the TCA cycle, however it would still be able to generate the 381 382 proton motive force via the Complex II (succinate dehydrogenase CSP5 0486-0489 and CPM 0468-0451). As expected, glyoxylate bypass seems to be 383 384 inoperative: isocitrate lyase was found, but not the malate synthase. 385 In the course of growth of *C. divulgatum* on peptides, the lack of recirculation of TCA metabolites due to its incompleteness can be compensated by their synthesis 386 387 from amino acids. During potential sugars-driven growth, PEP can be converted to 388 oxaloacetate in a reversible reaction (which is not favourable, but possible), catalysed by GTP-dependent phosphoenolpyruvate carboxykinase (CSP5 1337 389 390 and CPM 1336) while malate or oxaloacetate can be synthesized from pyruvate 391 by a reverse reaction catalysed by malic enzyme (CSP5 0838, CPM 0835). 392 Despite the generation of the proton motive force at aerobic growth (complex II) on peptides or sugars (the latter was not confirmed experimentally in the current 393 394 experimental setup) TCA cycle enzymes play a crucial role in anabolism during 395 growth on peptides at both aerobic and anaerobic conditions. For example, the 396 mentioned above GTP-dependent phosphoenolpyruvate carboxykinase and malic 397 enzyme uses its metabolites for the first stages of gluconeogenesis: phosphoenolpyruvate and pyruvate synthesis, respectively. 398 399

400 NADH dehydrogenase

401 Both *Cuniculiplasma* genomes contain genes for four major respiratory complexes, with some unusual details, as specified below. A set of genes of the proton-402 translocating type I NADH-dehydrogenase (complex I) nuoABCDHIJJKLMN is 403 encoded by CSP5 1737-1726 in the strain S5 and CPM 1708-1687 in the strain 404 PM4, in the same order. Both genomes encode neither NuoG subunit, nor subunits 405 406 NuoE or NuoF homologs essential to provide the catalytic site for NADH oxidation, which raises doubts in NADH-oxidizing activity of the complex I and its involvement 407 in respiratory electron transfer chain in C. divulgatum. Alternative pathway of 408 electron inflow into the respiratory chain could be provided by succinate 409 410 dehydrogenase/fumarate reductase (Complex II), encoded by CSP5 0486-0489 in 411 S5 and CPM 0458-0461 in PM4 genomes. It should be mentioned that none of 412 Thermoplasmatales genomes available to date contain genes of NuoEF subunits, indicating possibly inherent feature of respiratory complex I in the organisms of this 413 414 deep phylogenetic lineage and possible existence of other yet unknown alternative 415 mechanisms of electron flow from NADH oxidation - in analogy to those proposed 416 in aerobically respiring Helicobacter pylori, also lacking NuoEF subunits of the complex I³⁸. 417

418

419 Quinol oxidising complex III and oxygen respiration

420 Quinol oxidising complex III in both *C. divulgatum* genomes is represented by 421 clustered genes of Rieske Fe-S protein and cytochrome *b* subunit of a typical 422 cytochrome *bc*₁ complex encoded by CSP5_1460-1459 in the S5 and CPM_1454-423 1453 in the PM4 strains. These clusters are located remotely with the genes of 424 terminal respiratory reductases in both genomes. No genes of an alternative 425 complex III have been detected in *C. divulgatum* genomes.

426 Terminal oxygen reductases are represented in both C. divulgatum strains by a typical cytochrome bd quinol oxidase (CSP5 0552-0553 and CPM 0524-0525) 427 and a heme copper oxygen reductase (HCO) encoded in the clusters CSP5 1313-428 429 1312 and CPM 1312-1311. The first enzyme complex possesses a high affinity to oxygen and is usually involved in oxygen detoxification or respiration under 430 microaerophilic conditions providing relatively low energy yield to the cell³⁹. The 431 heme-copper oxygen reductase (complex IV) is a typical terminal enzyme of 432 433 aerobic respiratory electron transfer chain, coupling oxygen reduction to proton 434 translocation at aerobic or microaerophilic conditions. Sequence analysis of 435 catalytic subunits I of the heme-copper oxidases of C. divulgatum strains with a 436 web-based classifying tool (http://evocell.org)⁴⁰ clearly showed that both of them 437 belong to the type A1 oxygen reductases possessing two proton translocating 438 channels, and consequently, the highest proton pumping stoichiometry of 2H⁺ per one electron^{41,42,43}. Our phylogenetic reconstruction of full-size CoxI available so 439 440 far (Fig. 5) generally reproduced the recently reported topology of HCO 441 phylogenetic trees and revealed that the A1-type heme-copper oxidases of 442 Cuniculiplasma species form a distinct clade, most closely branching to B-type 443 oxygen reductases and to the root of all the other A-type reductases. Interestingly, heme-copper oxygen reductases from other Thermoplasmatales (from Acidiplasma 444 and Picrophilus species) are located on a distinct clade of A-type oxidases (Fig. 5). 445 Furthermore, both C. divulgatum strains lack genes for membrane-integral oxygen 446 447 reductase subunits III and IV (either separately encoded or fused to the C-terminus 448 of the catalytic subunit I), while those were found in Acidiplasma and Picrophilus 449 genomes being fused with coxl genes. The subunits III and IV are regarded to be 450 distinguishing features of A-type (SoxM) heme-copper oxygen reductases acquired

during their evolution from less energetically effective and more ancient B-type enzymes⁴². The lack of these subunits in *C. divulgatum* together with the phylogenetic position of its CoxI proteins allows assuming that this organism possesses an ancestral form of all known A-type terminal oxygen reductases.

455

456 A crucial point for the activity of the heme-copper oxygen reductase is the pathway of electron transfer from the quinone pool or complex III. In C. divulgatum genomes, 457 458 there are no genes of type I monoheme *c*-type cytochromes, providing the electron 459 transfer from respiratory complexes III to terminal oxidases. Alternative pathway could be driven by blue-copper redox proteins (cupredoxins), as described in several 460 461 acidophiles⁴⁴. A homolog of such cupredoxins has been found to be involved in the respiratory chain of *Ferroplasma acidiphilum*⁴⁵. As mentioned above, the gene 462 encoding a cupredoxin rusticyanin was identified only in C. divulgatum strain S5 463 (CSP5 0076). The absence of both genes of type I cytotchrome c and 464 465 rusticyanin/sulfocyanin does not allow predicting the electron transfer pathway between respiratory complexes III and IV in the strain PM4. The possibility still exists 466 that the complex IV in strain PM4 possesses guinol oxidizing activity and, similarly 467 to some other heme-copper oxidases, could directly accept electrons transferred 468 from the complexes I and II via the guinone pool. In such a case, the complex III in 469 470 strain PM4 would serve as an additional proton-translocating site, which is not 471 directly involved in oxygen respiration and could transfer electrons to an extrinsic, yet unidentified acceptor. However, this assumption needs experimental evaluation. 472 All analysed genomes code for subunits K, E, C, F, A, B, D, H and I of V/A type H⁺-473 474 transporting ATP synthases, in this particular order (CSP5 0034-0042 and 475 CPM 0034-0042).

476 Further central metabolic and protein folding pathways detailed in SI suggest
477 *Cuniculiplasma* spp. largely share these with other *Thermoplasmata*.

478

479 **Comparison with other Thermoplasmatales**

480 Phylogenomic analysis of *Thermoplasmata* based on concatenated amino acid

481 sequences of 11 conservative ribosomal proteins of each representative of the

482 phylum with a sequenced genome (Fig. 6 a), indicates a slightly different tree

483 topology than that suggested by 16S rRNA gene phylogenetic analysis¹³, likely due

to this selection of particular molecular markers for phylogenetic reconstruction. On

485 the other hand, IMG COG-based hierarchical clustering placed

486 *Cuniculplasmataceae* representatives close to the root of the order

487 *Thermoplasmatales* (Fig. 6 b). This might be an indication that *Cuniculiplasma* spp.

488 share more parental properties than other cultivated members of

489 *Thermoplasmatales* and thus could be a good model for analysis of yet

490 uncultivated members of the class *Thermoplasmata*.

491 In contrast to other *Thermoplasmatales*, the genome of *C. divulgatum* strain PM4

492 (but not S5) had no restriction-modification system Type I. Pyrimidine and purine

493 conversion and utilization pathways, RNA processing and modification processes

494 showed their incompleteness in *C. divulgatum*, in comparison to the rest of

495 *Thermoplasmatales.* We also infer that amino acid biosynthesis category for other

496 Thermoplasmatales (T. acidophilum, P. torridus, and "F. acidarmanus") showed

497 some discrepancies to C. divulgatum. Thus, P. torridus has been proposed to

498 possess all pathways for the amino acid synthesis¹⁶. *"F. acidarmanus"* occurred to

499 encode incomplete histidine, valine, leucine and isoleucine synthesis pathways⁴.

500 The genomes inspection of *C. divulgatum* suggested, in addition to the above,

incomplete pathways for lysine and proline, pointing at the organisms' dependenceon external peptides and hence suggesting their role in the environment as

503 'scavengers'.

504 Incidentally, C. divulgatum and "F. acidarmanus" genomes, but not T. acidophilum

505 or *P. torridus* encode proteins for capsular heptose metabolism and

506 polyhydroxybutirate metabolism (with an exception of the gene encoding for

507 acetoacetyl-CoA synthetase (EC 6.2.1.16 in "F. acidarmanus").

508 The organisms have a weak potential for synthesis of polymeric storage

509 compounds: both genomes for a similar folylpoly(gamma)glutamate synthase

510 (CPM_0655 and CPM_1446). The PM4 also contains an inorganic

511 polyphosphate/ATP-NAD kinase (CPM_0378) putatively active in energy gaining

512 from environmental polyphosphate deposits. However, no cell inclusions were

513 observed.

514 Formate dehydrogenase complex, involved into catabolism of C1 compounds, which is a common trait for T. acidophilum and "F. acidarmanus" has not been verified in 515 either *Cuniciliplasma* genome. The gene coding for aguaporin Z (MIP superfamily) 516 was found in both genomes of C. divulgatum, potentially contributing to the osmotic 517 518 stress response and adaptive fitness, but absent in the genomes of other members of Thermoplasmatales. Another distinctive feature is the lack of molybdenum 519 520 cofactor and coenzyme M biosynthesis in C. divulgatum genomes in contrary to 521 other Thermoplasmatales. Finally, the lack of ATP-dependent DNA ligases in C. divulgatum genomes has been observed. The global analysis of distribution patterns 522 of arCOGs in Thermoplasmatales is further detailed in SI and suggests 523 524 *Cuniculiplasma* is a common member of the order.

526 **Discussion**

527 Isolation of previously uncultured microorganisms from the environment remains one of the bottlenecks in microbiology hindering physiological and biochemical 528 529 studies and demanding a resolution. It is especially important for archaea, the relatively recently discovered Domain, and which embraces a majority of difficult-530 531 to-culture organisms. The cultured diversity of archaea is dramatically low: according to the Euzeby LSPN online resource (http://www.bacterio.net/), only 532 533 some 116 genera and 451 species with validly published names of archaea (of 534 which 55-60% are haloarchaea-related organisms) vs some 2277 genera и 11940 535 species of cultured and described bacteria are known to-date. The acidophiles of 536 the order *Thermoplasmatales* are a good example of this status of things, 537 accounting for only six cultured genera published since 1970, despite numerous documentations on the presence of highly diverse Thermoplasmatales-like 538 539 organisms in low-pH habitats worldwide. The present genomic analysis of new successfully cultured *Thermoplasmatales* members¹³ brought us closer to the 540 541 understanding of functional diversity within this archaeal group. Interestingly, these archaea represent a unique case for Thermoplasmatales, when organisms from 542 543 the same species and almost identical genomes from different geographic 544 locations became cultured. Metabolically, Cuniculiplasmataceae resemble other 545 Thermoplasmatales members, however certain discrepancies suggest some 546 variety of their evolutionary trajectories. *Cuniculiplasma* spp. genomes encode the A1-type heme-copper oxidases forming a distinct clade at the root of A-type 547 548 reductases and closely branching to the B-type oxygen reductases and are 549 deficient in membrane-integral oxygen reductase subunits III and IV, suggesting 550 that, in contrast with other *Thermoplasmatales*, they have a more ancient and less

551 energetically efficient B-type enzymes. Cuniculiplasma spp. exhibit largest genomes among *Thermoplasmatales* seemingly at the expences of genetic loci for 552 553 heavy metal resistance and defense systems. Scavenger type of nutrition was 554 confirmed as a characteristic trait for *Cunicuiplasma* spp., which is reflected in their genomic blueprints and physiology, suggesting these organisms feed in situ on 555 556 proteinaceous compounds derived from primary producing organisms. Based on the reconstructions of metagenomic data, the archaea related to this species 557 558 previously supposed to be uncultured and associated to 'G-plasma cluster' are found in many acidic environments^{1,6}. Certain features predicted from the 559 metagenomic assembly "G-plasma" have not been confirmed highlighting the 560 561 essentiality of cultivation efforts and experimental functional validation of genomic 562 predictions. Almost identical genomes of the two European isolates and their North American sibling and strong conservation within their genomic islands, suggest a 563 massive stabilizing selective pressure in similar acidic environments and/or 564 565 significant fidelity of DNA repair systems assure their genome stability. Isolation of reference strains and experimental validation of genomic predictions for 566 567 this archaeal group should be considered in the future as tasks of a highest priority. 568

569 Methods

570 Sampling, and culturing, DNA isolation and sequencing.

Samples from acidic streamers for isolation were taken in March-April of 2011 from
Cantareras (Spain) and Parys Mountain/Mynydd Parys (UK) copper-containing
sulfidic ores. Both cultures were grown in AB Medium, pH 1-1.2, as described
previously¹³. DNA was isolated by GNOME DNA Isolation Kit (MP Biomedicals).

576 Genome sequencing and analysis protocol

The genomes were sequenced at Fidelity Systems, Inc. (Gaithersburg, MD) using 577 Illumina HiSeq 2000 platform, combining short paired-end libraries of 400 bp and 578 579 long mate-paired 3,600 bp inserts with an average read length of 100 bases using manufacturer protocols with the only modification that for the PCR amplification of 580 the genome library the TopoTag DNA polymerase was used⁴⁶. Initially, Velvet v. 581 1.2.10 was used to assemble the contigs⁴⁷. Scaffolding, filling the gaps, and repeat 582 resolution were performed using the Phred, Phrap, Consed software package⁴⁸ 583 584 and in-house software of Fidelity Systems. The error rate quality score of the 585 completed genome sequences was of Phred 50. The final assemblies provided 586 564- and 561-fold coverages for strain S5 and PM4, correspondingly. The genome 587 annotation was done at Fidelity Systems Ltd. using FgenesB 2.0 (SoftBerry, Inc., NY) followed by manual curation. The Rfam 11.0 database 588 (http://rfam.sanger.ac.uk)⁴⁹ and Infernal 1.0.2 (http://infernal.janelia.org)⁵⁰ were 589 used for annotation of RNA genes. 590 591 For analysis of shared and unique proteins all in silico translated genes were filtered by a length of 150 amino acids to exclude false predictions from the 592 593 analysis. Resulting proteins were subjected to 'all vs all' alignment with blastp algorithm⁵¹ and e-value cut-off of 10⁻⁵. Resulting blast table was used as an input 594 595 for OrthoMCL analysis with grain value of 2.5.

596 Assignment of predicted CDS to the archaeal clusters of orthologous groups

597 (arCOGs) was made using blastp against the latest version of arCOG database⁵²

⁵⁹⁸ with maximal e-value of 10⁻⁵. blastp hits were filtered to have minimum alignment

599 length more than 50% of query and subject sequences length. arCOG was

assigned to a protein if the hits to at least 3 different genera were registered.

Phylogenetic analyses were performed in MEGA 6⁵³ using Maximum likelihood
method and bootstrap confidence test. Sequence alignments were performed in
MAFFT v. 7⁵⁴.

Metagenome data search was performed through the following databases: MG-604 RAST⁵⁵, IMG-M-ER databases⁵⁶ and SRA archive⁵⁷. Metagenome sequencing 605 projects related to acidic environment were identified using keywords "acid" "mine" 606 "drainage" "copper" and its combination. Cuniculiplasma related sequences were 607 detected using blastn algorithm. Sequences with identity > 95% were considered 608 609 positive hits for MG-RAST and IMG-M-ER, while for NCBI SRA sequences identity cut-off was set to 99 %. CRISPR repeat sequences were analysed locally using 610 611 blastn tool of NCBI blast 2.4.0+ package against NCBI nt/nr, env nt and htgs 612 databases, PM4, S5 and 'G-plasma' genomes and against metagenomes acidic 613 environments found in IMG-M database (Gp0051182, Gp0097388, Gp0097859, Gp0097858, Gp0053344 and Gp0053343). Parameters were as follows: word size: 614 615 7, match score: 1, mismatch penalty: -1, gap open penalty: 10, gap extension penalty: 2, percentage of query covered: 90, percentage identical bases: 90. 616 Genomic islands (GIs) in C. divulgatum were inspected using Island Viewer 3⁵⁸. 617 618

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626 **Conflict of interest**

627 The authors declare no conflict of interest.

628

629 Authors contributions

630 OVG and PNG convened the research. OVG, IVK, TH, AAK, TYN, SNG, SVT and

631 PNG did genome analysis. OVG, HL, IVK, SNG, SVT and PNG wrote the

632 manuscript.

- 635 Supplementary Information is available at the Journal website.

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802

803 Table 1. Overview of general genomic features of *Cuniculiplasma divulgatum*,

804 strains PM4 and S5 and 'G-plasma'

	Strain PM4	Strain S5	G-plasma*
Number of bases	1878916	1938699	1827255
Number of chromosome contigs	1	1	22
introns	1	5	ND
GC mol %	37.16	37.3	38.9
Coding density, %	87.1	87.4	88.5
Genes	1948	2016	1923
tRNA	46	46	48

* "G-plasma" genome stats may be affected by the application of a different annotation
pipeline and the fact that it was assembled from metagenomic reads⁴ as opposed to the
genomic assembly of pure cultures of strains S5 and PM4. ND, not determined.

812 Figure legends

813 **Figure 1.** Worldwide distribution of *Cuniculiplasma*-related archaea. Map was

814 created using Plotly online package (<u>https://plot.ly/</u>) using geographical

815 coordinates, retrieved from metadata of database entries.

816

Figure 2. Electron micrographs of *Cuniculiplasma divulgatum* showing monolayer

818 membranes and absence of the S-layer (a & b), pilus (c, arrow) and pleomorphism

of cells. Scale bars: 500 nm (a), 200 nm (b), 1 μm (c, d). Ultrathin sections (a, b)

and Pt-C shadow castings (c, d). Figure shows cells of the strain PM4 (b, c and d)

and S5 (a). Arrowheads in c and d indicate the direction of shadow cast, arrows in
a and b point to the cytoplasmic membrane.

823

Figure 3. Distribution of arCOG Functional Categories within core, accessory and

strain-specific (unique) proteomes of *C. divulgatum* S5 and PM4 and 'G-plasma'.

826 Circle area is proportional to the fraction of corresponding arCOG FC to the total

number of arCOG hits in every group of proteins. Core group corresponds to

828 proteins found and all three genomes, accessory group includes proteins found in

829 at least two genomes and unique group consists of strain-specific proteins.

830

Figure 4. Genomic islands (GIs) in *C. divulgatum* strains PM4 and S5. Rings from
outside to inside: genomic coordinates (grey colour); plus-strand CDS (blue) and
RNA (red);); minus-strand CDS (blue) and RNA (red); strain-specific CDS (red)
and genomis islands (green); blastn hits with e-value cutoff 10⁻⁵ vs other *C*.

835 *divulgatum* isolate; blastn hits with e-value cutoff 10⁻⁵ vs 'G-plasma'. Function of

Gls is marked by small figures: 'defense' islands – squares, 'transporter' islands –
triangles, islands of non-specific function – circles.

838

Figure 5. Maximum Likelihood phylogenetic tree of PF00115 polypeptides (COX1 839 family). Totally 112 sequences were used in analysis after 50% sequence identity 840 841 filtering. The tree with the highest log likelihood (-68482.6023) is shown. The percentages of trees in which the associated taxa clustered together (bootstrap 842 843 values, 1000 replicates) are shown next to the branches. The tree is drawn to 844 scale, with branch lengths measured in the number of substitutions per site. All positions with less than 95% site coverage were eliminated. Unclassified group 845 846 was first mentioned⁴². Nitric oxide reductases (NOR) were placed as an outgroup. 847

848 **Figure 6.** Phylogentic position of *Cuniculiplasma* spp. within *Thermoplasmata*.

849 A. Maximum Likelihood phylogenetic tree, based on concatenated sequences of 11

850 conservative ribosomal proteins of two Cuniculiplasma strains, nine

851 Thermoplasmata representatives with the genomes, publically available in IMG and

852 Methanopyrus kandleri AV19 as an outgroup (not shown on the tree). The proteins,

involved into analysis were: COG0048, ribosomal protein S12; COG0049,

ribosomal protein S7; COG0081, ribosomal protein L1; COG0197, ribosomal

protein L16/L10AE; COG0200, ribosomal protein L15; COG0244, ribosomal protein

L10; COG1631, ribosomal protein L44E; COG1890, ribosomal protein S3AE;

857 COG2004, ribosomal protein S24E; COG2051, ribosomal protein S27E; COG2125,

ribosomal protein S6E (S10). The tree with the highest log likelihood (-24738.5123)

is shown. The percentages of trees in which the associated taxa clustered together

860 (bootstrap values, 1000 replicates) are shown next at branching points. All

- 36
- positions with less than 95% site coverage were eliminated. There were a total of
- ⁸⁶² 1607 positions in the final dataset. The tree was constructed in MEGA6⁵³.
- 863 **B.** IMG COG-based hierarchical clustering. The analysis was performed using IMG
- genomic annotations of two *Cuniculiplasma* strains and nine publically available
- 865 *Thermoplasmata* representatives. Bars indicate the number of substitutions per
- 866 site.
- 867