‘ARMAN’ archaea depend on association with euryarchaeal host in culture and in situ

Olga V. Golyshina1, Stepan V. Toshchakov2, Kira S. Makarova3, Sergey N. Gavrilov4, Aleksei A. Korzhenkov2, Violetta La Cono5, Erika Arcadi5, Taras Y. Nechitaylo6, Manuel Ferrer7, Ilya V. Kublanov2,4, Yuri I. Wolf3, Michail M. Yakimov2,5 & Peter N. Golyshin1

Intriguing, yet uncultured ‘ARMAN’-like archaea are metabolically dependent on other members of the microbial community. It remains uncertain though which hosts they rely upon, and, because of the lack of complete genomes, to what extent. Here, we report the co-culturing of ARMAN-2-related organism, Mia14, with Cuniculiplasma divulgatum PM4 during the isolation of this strain from acidic streamer in Parys Mountain (Isle of Anglesey, UK). Mia14 is highly enriched in the binary culture (ca. 10% genomic reads) and its ungapped 0.95 Mbp genome points at severe voids in central metabolic pathways, indicating dependence on the host, C. divulgatum PM4. Analysis of C. divulgatum isolates from different sites and shotgun sequence data of Parys Mountain samples suggests an extensive genetic exchange between Mia14 and hosts in situ. Within the subset of organisms with high-quality genomic assemblies representing the ‘DPANN’ superphylum, the Mia14 lineage has had the largest gene flux, with dozens of genes gained that are implicated in the host interaction.
Deep metagenomic analysis of environmental samples from acidic environments across our planet has demonstrated the existence of previously neglected uncultured archaea that are only very distantly related to recognised phyla. Initially detected at Iron Mountain (California, USA), these archaeal lineages were subsequently confirmed to occur in various acid mine drainage (AMD) systems. This enigmatic group of archaea (the so-called ‘Archaeal Richmond Mine Acidophilic Nano-organisms’, or ‘ARMAN’) was initially found in the fraction of cells filtered through 0.22 μm membrane filters. Metagenomic assemblies suggested average genome sizes of these organisms to be relatively small for free-living organisms (approximately 1 Mbp). An interesting observation documented by electron microscopy was that some cells of a small size (<500 nm) interact through pili-like structures with larger cells that lacked cell walls. Comolli and colleagues suggested the ‘ARMAN’ organisms were the ‘small’ cells, whereas cell wall-deficient larger cells were attributed to some members of the order Thermoplasmatales, a group of organisms known to be widely represented in AMD systems. Emerging findings from metagenomic data sets of ARMAN-like archaea and especially their ubiquity, suggest that this group plays important roles in the environment, although the exact roles have yet to be established. The phylogenomic placement of archaea from this group still represents a matter for discussion.

The known example of small-sized cultured archaea is represented by Nanoarchaeum equitans, currently the only validly described member of the phylum Nanoarchaeota. Cells are about 500 nm (or smaller) in diameter and exhibit atypical archaeal ultrastructure. Nanoarchaeum equitans exists only in association with the host, Ignicoccus hospitalis, which supplies certain organic compounds (lipids and amino acids), growth factors and likely ATP to N. equitans. Other nanoarchaeota-related examples include an Nst1 archaeon forming an association with its host, the Sulfolobales-related organism, and ‘Candidatus Nanopusillus acidilobi’, thriving in a partnership with Acidilobus spp. These nanoarchaeota are hyperthermophilic marine and terrestrial organisms with extremely compact genomes that likely are not of an ancestral nature, but rather probably resulted from massive gene loss. Nanoarchaeota-related organisms (including those known only by metagenomics-resolved genomes) are phylogenetically clustered within the ‘DPANN’ candidate superphylum (abbreviated after candidate divisions ‘Diaphetotrites’, ‘Parvarchaeta’, ‘Aenigmarchaeota’, ‘Nanohalararchaeota’ and the only validly described phylum Nanoarchaeota). Recently, a number of uncultured ‘DPANN’ archaea with almost complete genomes were predicted by Castelle and co-authors to be symbiotic and/or have a lifestyle based on fermentation. To summarise, all experimentally validated examples of interactions between co-cultured small (or ‘nano-sized’) archaea and their partners are limited to Crenarchaea being the hosts. All of them (except Ignicoccus sp.) are acidophiles, while so far no associations have been co-cultured or characterised for Eurarchaeota, except those from the recent report on a four-member consortium containing a fungus, two strains of Thermoplasmatales and ARMAN-1-related organism with, due to the complexity of this enrichment culture, only a partially sequenced genome.

Here, we report the co-cultivation and analysis of the ungapped genome of an ARMAN-like organism, the ‘Candidatus Mancarchaeum acidophilum’ Mia14, which was enriched in the laboratory binary culture with Cuniculiplasma divulgatum PM4, a recently described representative of the family Cuniculiplasmataceae within Thermoplasmata. After additional sampling campaigns and de novo metagenome sequencing of the microbial community of the acidic streamer of Mynydd Parys/Mynydd, we revealed possible in situ interactions of these organisms with other microbial community members. Furthermore, we analysed the voids in its metabolic pathways (and thus dependencies on potential hosts) and mapped its phylogenetic position. Finally, using data on arCOGs gains and losses, we reconstructed its evolutionary trajectory starting from the last archaeal common ancestor (LACA), which pointed at Mia14 having the greatest known extent of gene fluxes within the ‘DPANN’ superphylum.

Results
Coexistence of Mia14 with Cuniculiplasma divulgatum PM4. We have previously isolated and described two strains of a new archaeal family, genus and species, named Cuniculiplasma divulgatum (order Thermoplasmatales), from acidic streamers at Parys Mountain (UK) and Cantareras mine (Spain). Both strains S5 and PM4 were characterised as acidophilic organoheterotrophes with mesophilic optima for growth and as facultative anaerobes. The genomes of these isolates were remarkably similar to one another (>98% average nucleotide identity) and to that of the genomic assembly ‘G-plasma’ from Iron Mountain (USA)11. During the isolation, C. divulgatum strain PM4 was co-cultured for 2 years with another archaeon designated Mia14 with a proportion of genomic reads, PM4: Mia14 of approx. 10:1. The initially poor growth of the PM4 component was significantly improved by the addition of complex organic compounds, such as beef extract and tryptone (0.1% w/vol). However, the enhanced growth of C. divulgatum and an increased frequency of re-inoculations had a dramatic effect on the growth of Mia14, which was eliminated from the culture and, after approximately 2.5 years of regular (every 20–22 days) passages into the fresh medium, was not detectable by PCR with specific primers. Another possible explanation is that the faster growth of C. divulgatum strain PM4 was the result of the elimination of Mia14, which may have negatively affected the growth of PM4 in earlier cultivation stages. Whatever the case, we could not maintain Mia14 for longer than 2.5 years. However, as Mia14 was highly enriched in the initial enrichment cultures with C. divulgatum strain PM4, we obtained enough coverage of its genomic reads (approximately 40-fold) to assemble a single chromosome. After the loss of Mia14 from the enrichment culture, we performed additional sampling of the acidic streamer (from the same site in Parys Mountain where the isolate PM4 was derived from), and detected Mia14 initially by PCR using specific primers, then by the de novo sequencing of environmental DNA, and ultimately, by catalysed reporter deposition fluorescence in situ hybridisation (CARD-FISH).

Analysis of metagenomic contigs showed that the most abundant group (up to 57%) was Thermoplasmatales–related archaea. Small-genome archaeal lineages (‘Candidatus Parvarchaeta’ and ‘Ca. Micarchaeota’) were also detected at 0.31% and 3.84%, respectively (Fig. 1).

Interestingly, both median contig coverage and coverage-based abundance calculation indicate that the amount of Cuniculiplasma cells in the Parys Mountain acidic streamer is nearly equal to the amount of ‘Candidatus Mancarchaeum’ cells (Figs. 1 and 2). Also, analysis of read coverage vs. GC content of metagenomic contigs reveals that ‘Ca. Mancarchaeum’ and C. divulgatum-related contigs form a very compact cluster similar both in coverage and GC content (Fig. 2). Notably, we also observed another Thermoplasmatales, ‘Ca. Micarchaeota’ contig cluster, in the Parys Mountain metagenome, suggesting that there are several stable two-member microbial associates in the community at this site.
Fluorescence microscopy shows interaction of Mia14 and the host. Microbial cells from enrichment cultures set-up with the environmental sample from 2014 were either hybridised with probe EUB338(I-III) mix or probe ARCH915 to target Bacteria or Archaea, respectively. The following CARD-FISH analysis revealed dense populations of Archaea and the almost complete absence of bacterial cells. Pleomorphic morphologies of cells of various size, typical for Cuniculiplasma/Thermoplasmatales16, were confirmed with hybridisations with Cuniculiplasma-specific probe Clpm-1100R. Besides bright signals, the CARD-FISH microphotography retrieved numerous debris-like structural forms, which likely could be referred to as either dying or metabolically dormant cells. This observation is typical for both natural samples and initial enrichments, where the cells of different metabolic states coexist. Noteworthy, parallel hybridisations with ‘Ca. Mancarchaeum’-specific probe ARM-MIA1469R and Thermoplasmata-specific probe Thpmt680R showed quite similar images (Fig. 3), suggesting that the organisms live in a tight association. Cross-hybridisation of ARM-MIA1469R probe with pure Cuniculiplasma culture was controlled at specific hybridisation conditions and no positive signals were retrieved. Side-by-side comparisons of ‘Ca. Mancarchaeum’ vs. Cuniculiplasma cells revealed that the former are slightly smaller in size and only a minor fraction of cells do not overlap in each frame. Detailed view of some double-hybridised cell formations revealed single coccoid-shaped Cuniculiplasma cells were surrounded by ARM-MIA1469R probe-labelled organisms (Fig. 3f).

Phylogenetic position of Mia14 and related organisms. Based on 16S ribosomal RNA (rRNA) gene sequence, Mia14 was found to be only distantly related to organisms with established taxonomic status. Less than 75 % SSU rRNA gene sequence identities with the thaumarchaeon Nitrospira viennensis and euryarchaeon Methanosaeta consilii are observed. Among candidate status holders (Supplementary Fig. 1), the nearest relative was inferred to be ARMAN-2 (‘Candidatus Micarchaeum acidiphilum’ excluding Ca. Mancarchaeum acidiphilum* excluding Ca. Mancarchaeum acidiphilum’ original detected in acidic environments and sharing 92% 16S rRNA sequence identity with Mia14. Other similar sequences (92% sequence identity) belong to PCR-amplified and cloned SSU rRNA genes from fumarolic thermal and acidic green biofilms, Mexico, Michoacan, Los Azufres (KJ907762). Interestingly, both above sequences and the sequence of Mia14 possess introns in their 16S rRNA genes. In addition, sequences with a lower sequence identity and coverage (91%, 58%) were detected in a PCR-amplified SSU rRNA clone from Rio Tinto (FN865418)18, acidic hot springs (JF280243; 91%, 58%)19 and a number of other AMD and volcanic environments. Furthermore, similar sequences have been retrieved from southern Appalachian peatlands (PF82012)20 and wetlands in Finland (AM905392, AM905420)21. The two latter sites were oligotrophic, with temperatures in the range from 0 to 15 °C and slightly acidic pH (4–5.6 and 3.9–4.3, respectively). Along with wetland clones, similar signatures (BioProject PRJNA279923) have been found in metagenomic data from another oligotrophic environment, the pH-neutral groundwater from Fennoscandian terrestrial deep biosphere22. All these records suggest a wide distribution of organisms similar to Mia14 and ARMAN-2 in natural settings with various pH characteristics, not necessarily tied to acidic environments.

The placement of Mia14 on the phylogenetic tree constructed with concatenated ribosomal proteins is presented in Fig. 4. In agreement with previous observations12, the position of Mia14 within the ‘DPANN’ superphylum is strongly supported.

Genome statistics. The genome of Mia14 is a single, circular chromosome with 952,257 bp, with the molar G+C% of 39.36% (Fig. 5). The coding density in the genome is of 1.032 genes per
kbp (968 bases per gene). About ~ 150–200 hypothetical proteins were present. The genome encodes 45 transfer RNAs. Three introns were detected across the chromosome. All these traits are typical for small archaeal genomes, e.g., in *Nanoarchaeum equitans* (491 kbp) \(^{23}\), *Candidatus Nanobsidianus stetterii*, Nst1 belonging to the phylum Nanoarchaeota (592 kbp) \(^{10}\), ARMAN-2 (~ 1 Mbp) \(^{5}\) and other host-associated or symbiotic microorganisms.

**Lateral gene transfer between Thermoplasmatales and Mia14.** Comparative analysis of in silico proteomes of Mia14 and strains S5 and PM4 with ProteinOrtho\(^{24}\) revealed several clusters of orthologous genes shared between Mia14 and *C. divulgatum* S5, but absent in *C. divulgatum* PM4 (Supplementary Data 1, Supplementary Fig. 2). These genes encode several membrane-associated proteins (MIA14\(_{0876, 0886, 0893, 0478}\)), two SAM-dependent methyltransferases (MIA14\(_{0883, 0885}\)), sulfocyanin (MIA14\(_{0884}\)) and peroxiredoxin (MIA14\(_{0479}\)). It should be noted that the majority of these proteins have homologues in PM4, but are more distant to those from both Mia14 and S5 and have different gene context. Few Mia14 genes from these clusters have no homologues in PM4.

Analysis with IslandViewer\(^3\)\(^{25}\) showed that altogether five genomic islands (GIs) are present; the largest GI contains 41 genes and spans 36.5 kbp (Fig. 5). A closer inspection of this island reveals that the integration occurred in the gene for zinc-binding pyruvate-formate lyase-activating enzyme (MIA14\(_{0850}\)), splitting it in two parts: MIA14\(_{0850}\) and MIA14\(_{0891}\), with the latter located in the immediate vicinity of 23S rRNA gene. About 50% of genes within this GI could not be assigned to known arCOGs and represent small proteins that often contain transmembrane segments, which is typical for archaeal ‘dark matter’. In turn, the genes assigned to arCOGs (i.e., MIA14\(_{0898}\), the DNA invertase Pin homolog, MIA14\(_{0894}\) similar to those from other Thermoplasmatales), ParA family chromosome partitioning ATPase and MIA14\(_{0890}\), integrase of XerD family) were shown to be strongly associated with ‘dark matter’ islands in archaeal genomes and could be specifically attributed to integrated mobile elements\(^{26}\).

Among GI-associated genes, we also found cation transport ATPase/copper-transporting P-type ATPase (MIA14\(_{0877}\)), which may have significance for the fitness of this organism in the harsh conditions of Parys Mountain AMD. Phylogenetic analysis of this ATPase showed that its close homologues are widely distributed among acidophilic Thermoplasmatales. At the same time, the copper-transporting ATPase of ARMAN-2 seems only quite distantly related to MIA14\(_{0877}\) (Fig. 6, Supplementary Table 1). Gene neighbourhood of MIA14\(_{0877}\) included...
an Lrp-AsnC family transcriptional regulator and a copper chaperone, resembling functional copper fitness islands described for *Ferroplasma acidarmanus* [27]. This gene cluster was found to be conserved in *Cuniculiplasma*-related archaea. Furthermore, *C. divulgatum* S5 genome possessed two copies of this copper fitness island (Fig. 6). Interestingly, one of the *C. divulgatum* S5 copper fitness islands was adjacent to genetic loci for SHOCT family and DUF 302 family proteins as in the Mia14 copper gene cluster, while another *C. divulgatum* S5 copper gene island had a high level of gene synteny with *C. divulgatum* PM4 (Figs. 5 and 6). The above observation supports the lateral gene transfer from ancestral *Cuniculiplasma*-related lineage(s) to Mia14. In that case, it is more likely that LCA of *Cuniculiplasma* had two copies of this gene cluster, one of which was lost during the evolution of *C. divulgatum* PM4 and ‘G-plasma’.

Detailed analysis of de novo metagenome sequencing data from Parys Mountain samples shows that gene clusters similar to the abovementioned copper fitness island of Mia14 are widely present in different metagenomic contigs (Supplementary Data 2). It indicates that this highly mobile gene set is important for heavy metal resistance in microbial communities inhabiting acidic environments with high concentrations of dissolved metal ions.

Other smaller GIs of Mia14 (Fig. 5) contain defence systems (toxin/antitoxin and type III restriction-modification proteins), 2-oxoacid dehydrogenase multienzyme complexes, 2-oxoacid decarboxylase (E1) component subunits α and β, glycosyltransferases and numerous hypothetical proteins. Interestingly, the laminin G-encoding gene locus is also situated on the GI (see the section 'Secretion systems').

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**Fig. 4** Phylogenetic position of Mia14 within Archaea. An approximate Maximum Likelihood tree based on the concatenated alignment of 56 ribosomal proteins universally conserved in Archaea. In total, 285 genomes were analysed. Taxa are named according to the NCBI taxonomy. Candidate phyla are shown in quotation marks. Lineages with cultured/co-cultured representatives are highlighted in blue. NCBI Genome Assembly IDs are shown for individual genomes. Scale bar reflects 0.1 substitutions per amino-acid position.
Carbohydrate metabolism. The Mia14 genome has no genes for central carbohydrate metabolism pathways such as glycolysis and gluconeogenesis, pentose phosphate pathway or tricarboxylic acid (TCA) cycle. A detailed manual inspection suggested that the genome encodes a complete set of enzymes for glucose oxidation via the non-phosphorylating Entner-Doudoroff (ED) pathway28: a glucose dehydrogenase (MIA14_0575), d-gluconate dehydratase (MIA14_0298), 2-dehydro-3-phosphogluconate aldolase (MIA14_0299) and NAD-dependent d-glyceraldehyde dehydrogenase (MIA14_0297). Surprisingly, no enzymes for further conversion of glyceral, e.g., to glyceral-2-phosphate or glyceral-3-phosphate were found. The pyruvate released during the action of 2-dehydro-3-phosphogluconate aldolase, could be carboxylated to malate and oxaloacetate in the reaction catalysed by NAD-dependent malic enzyme (MIA14_0243/EC 1.1.1.38). Characterised homolog of scfA from *E. coli* (or *maeA*29) was reversible despite the carboxylation reaction being 28 times slower than the forward reaction. Other enzymes found to catalyse pyruvate conversions are phosphoenolpyruvate synthase/pyruvate phosphate dikinase (MIA14_0437, EC 2.7.9.2 and MIA14_0462) and pyruvate kinase (MIA14_0326, EC 2.7.1.40).

It is worth mentioning that ARMAN-2, one of the most closely related organisms to Mia14 among those with partially sequenced genomes, exhibits a relatively scarce repertoire of genes in comparison to sibling lineages ARMAN-4 and -5. Only a few genes for glycolysis in ARMAN-2 and a near-complete set of genes in ARMAN-4 and -5 were predicted. TCA, which is dysfunctional in Mia14, was reported to be complete or almost complete in all ARMAN cluster organisms mentioned above5. Furthermore, central metabolic pathways in Mia14 starkly contrast with AR10 assembly representing ‘Ca. Diapherotrites’, but to some extent resemble those predicted for a more phylogenetically distant AR20 (‘Ca. Woesarchaeota’)13. Furthermore, the inspection of amino-acid biosynthetic pathways in Mia14 found them to be either incomplete or entirely missing. However, the total number of proteins in this functional category is higher in comparison to *N. equitans* and Nst110, 23.

Cofactors, vitamins, prosthetic groups and pigments. No genes for coenzyme A, folate, lipoic acid, NAD and NADP cofactor, pyridoxin (Vitamin B6), heme and siroheme, thiamin biosynthesis and riboflavin, FMN and FAD metabolism were present in the entire genome of Mia14. The lack of functional pathways for cofactors and amino acids is quite characteristic for organisms with reduced genomes10, 23.

Protein metabolism. Protein processing and modification-related genes (G3E family of P-loop GTPases, peptide methionine sulfoxide reductase and Rio family of protein kinases, amino-
carboxy-terminal intein-mediated trans-splice, and ribonucleotide reductase of class III (anaerobic), large subunit (EC 1.17.4.2) were missing in Mia14. Altogether, we have identified 35 large- and 26 small-subunits of ribosomal proteins; L37e (arCOG04126), S17e (arCOG01885) and S27e (arCOG04108) were absent. All three and, correspondingly, two former proteins were found in N. equitans and Nst1 (Supplementary Data 3).

Secretion systems. We have identified a number of genes affiliated with secretion processes in the genome of Mia14 (Supplementary Table 2). Two distinct type IV pili systems are present in the genome: one belongs to Methanococci/Methanothermobacteria/Thermococci group (MIA14_0170-_0177) and another is more genome: one belongs to Methanococci/Methanothermobacteria/Thermococci group (MIA14_0170-_0177) and another is more similar to a euryarchaeal group (MIA14_0252-_0260) 30. No Thermococci-related genes were found, in agreement with the loss of motility in most of ’DPANN’ species. Only one FlaK-like prepilin peptidase (MIA14_0570) was found. Key components of both systems are shared by many ‘Ca. Micarchaeas’ species. Additionally, the genome encodes Sec translocon genes for pre-protein translocase subunits SecYE (MIA14_0832, _0132) and Sec61Beta (MIA14_0736), SecDF (MIA14_0121 and _0122), signal peptide peptidase and signal recognition particle subunits and receptors. The presence of Sec-independent Tat pathway genes, suggests this system is operational for secretion of folded proteins.

The Mia14 surface layer deserves special attention. Besides a protection function in archaea, this compartment often regulates both cell adhesion and cell–cell interaction. We identified at least eight different proteins that eventually account for the architecture of the surface layer. It contains strain-specific secreted proteins with polycystic kidney disease (PKD) superfamily fold and β-propeller repeat domains fused to CARD (cell adhesion related domain found in bacteria)-like adhesion module. Proteins of the β-propeller fold are ubiquitous in nature and widely used as structural scaffolds for ligand binding and enzymatic activity. This fold comprises between four and twelve four-stranded β-meaders, the so-called blades that are arranged circularly around a central funnel-shaped pore.

Another observation is the expansion of genes encoding jellyroll fold LamG-like proteins in the Mia14 genome, which are only distantly similar to other LamG proteins from archaea with Candidatus status and from bacteria, but generally abundant in ’DPANN’ superphylum species. This finding suggests an association of these proteins with laminin (glycoprotein)-containing extracellular matrix and their key role in host cell interactions. Some of them are linked to aforementioned type IV pili loci and are localised in GI (Fig. 5).

Respiration. The Mia14 genome encodes all typical subunits K, E, C, F, A, B, D, G, H and I of V/A Na+- and H+-transporting type ATP synthases, in this particular order (MIA14_0355-0364), whereas the genome of N. equitans encodes only five subunits of ATP synthase23. The analysis of conserved motifs supported H+-translocating V-type ATPase31.

All genes coding for cytochrome bd quinol oxidase were identified. Subunits I and II are encoded by MIA14_0653-0654. This type of oxidoreductase could generate proton motive force (PMF) by transmembrane charge separation, but do so without being a ‘pump proton’. The main electron acceptor for them is oxygen, but cytochrome bd oxidoreductases are usually induced in response to low oxygen concentrations and serve for oxygen detoxification32. The role of the cytochrome bd oxidoreductase in Mia14 is puzzling, as the organism completely lacks any genes for biosynthesis of isoprenoid quinones, which are the only electron donors for this electrogenic enzyme complex. Moreover, no genes were identified coding for electron donating type I NADH dehydrogenase or succinate dehydrogenase or other known respiratory complexes (III and IV).

Transporters. ABC transporters, amino-acid permeases, Major Facilitator Superfamily and others have been predicted in Mia14 (Supplementary Table 3) to notably outnumber those in nanoarchaeal genomes31.

Evolutionary patterns. Overall, compared to other ’DPANN’ group members, the Mia14 genome experienced an unusually high level of gene flux (Fig. 7). In addition to the 226 genes that do not belong to known arCOGs (a large fraction of such genes was probably also acquired at the terminal branches of the ’DPANN’ tree), Mia14 has lost determinants for over 400 arCOG families from the genome of its common ancestor with ’Ca. Iainarchaeum’/AR10 lineage (46% of the ancestral set), but also gained over 130 arCOGs (21% of its current arCOG complement). Gains and losses of comparable scale exist within the ’DPANN’ group tree (e.g., the loss of 49% of the ancestral genome in the lineage of AR17 or acquisition of 18% of the gene complement in the lineage of G17-L22-OTU1), but not on the same tree branch. Gene gains and losses seem to affect all functional groups equally with a notable exception of the ’Cell motility’ group where more gains than losses were predicted (Fig. 7). This functional group includes components of secretion systems, which might play a key role in interaction of Mia14 with its host. Moreover, as mentioned before many of the unique genes in Mia14 belong to GI, many of which encode membrane proteins and are associated with potential conjugative elements which might be involved in the extensive gene exchange between Mia14 and its host.

Analysing the trajectory of evolution of Mia14 from LAC through the prism of losses and gains of functional genes, a few interesting facts became apparent. Genes for the majority of enzymes of the TCA cycle were already lost during the transition from LACA to the ’DPANN’ ancestor, together with many genes involved with amino acid, vitamin and cofactor biosynthesis along with the CRISPR-Cas system. Glycolysis and gluconeogenesis were present in all ascendants of Mia14 (i.e., in LACA, ’DPANN’- and Mia14-’Ca. Nanohaloarchaeota’-ancestors). However, many genes of these pathways were lost en route to the extant Mia14 species. Pathways for pyrimidine and purine biosynthesis and salvage were also lost at the very last (and long) step of evolution from Mia14/Ca. Iainarchaeum’ ancestor to the modern Mia14, with 414 genes lost and only 131 gained (Fig. 7a and Supplementary Data 4).

Analysis of the taxonomic affiliations of the ’DPANN’ group species (Supplementary Data 5 and 6) shows that, in contrast with the other group members, the genome of Mia14 was and continues to be involved in extensive gene exchange with the Thermoplasmata lineages. Unsurprisingly, the most common source of the acquired genes is identified as Cuniculiplasma divulgatum, the Mia14 host.

Etymology. ’Candidatus Mancarchaeum acidiphilum’

Mancarchaeum (Manc. archaeum) M.L. mancus (adj.) crippled, maimed, referred to the absence of many pathways in the genome; N.L. neut. n. archaeum (from Gr. adj. archaios -ê- on, ancient), ancient one, archaeon; N.L. neut. n. Mancarchaeum, an archaeon with absence of many pathways in the genome.

a. c.i.d. ’phi.lum.’ M.L. neut. n. acidum from an acid; Gr. adj. philos from loving; M.L. neut. adj. acidiphilum means acid-loving.
Fig. 7 Gains and losses of arCOG families in the evolution of ‘DPANN’ group. a, Reconstruction of gene loss and gain along the ‘DPANN’ subtree. Triplets of numbers indicate the estimates for the arCOG complement, arCOG gains and arCOG losses, respectively, for the selected extant or ancestral genomes and adjacent tree branches. Estimates for the terminal branches are shown next to the extant genome names. The number at the base of the tree indicates the estimates for the arCOG complement, arCOG gains and arCOG losses, respectively, for the selected extant or ancestral genomes and associated tree branches. b, Number of arCOGs predicted to be gained or lost in the course of evolution of Mia14 lineage with respective COUNT probability >50% by arCOG functional categories. The functional classification of the arCOGs is shown for two major groups: C-Q—metabolic genes; J—informational genes; V—defence genes; R-S poorly characterised or uncharacterised genes (for details see ftp://ftp.ncbi.nih.gov/pub/wolf/COGs/arCOG/funclass.tab).

Discussion
In the present work, the enrichment culture from Parys Mountain AMD system was set-up to grow acidophilic members of the order Thermoplasmatales. The culture was eventually highly enriched in archaea from the genus Cuniculiplasma, and incidentally, with the significant (ca. 10% genomic reads or 20% of total population) community component belonging to yet uncultured archaea distantly related with ARMAN-2. Due to its high numbers in the enrichment, we were able to produce the fully assembled genome of the ‘ARMAN’-related organism. Based on the genome annotation and experimental data (co-existence in an enrichment culture and fluorescence microscopy), we inferred that the metabolic needs of this sentinel of Cuniculiplasma spp. termed ‘Ca. Mancarchaeum acidiphilum’ resemble to some
extent those of other archaea co-occupying the environment (e.g., reliance on external proteinaceous compounds and amino acids). However, the incompleteness or absence of the central metabolic pathways (e.g., TCA, glycolysis, quinone biosynthesis, etc.) and reduced genome size support an obligate partner-dependent (or ‘ectoparasitic’) lifestyle. Our data (Fig. 3) further suggest that sizes of Mia14 cells (and likely other ARMAN-related archaea) have a broad range, usually larger than the diameter of membrane filter pores (0.22 μm) used to enrich for these organisms. The penetration of cells through the 0.22 μm pores of membrane filters observed previously may also be explained by the lack of rigid cell walls in these organisms. For example, the majority of 1–2 μm, cell wall-deficient Thermoplasmatales may squeeze through pores of this diameter.

The occurrence of laterally transferred genes and GIs from *Cuniculiplasma* spp. in Mia14 highlights the relative connection between these organisms co-existing in one environment. It is, furthermore, likely that extracellular structures such as pili or pili-like organelles might be present in Mia14. One may also speculate on massive exchange of DNA through surface pores or by using the Type IV pili system and numerous membrane proteins encoded within GIs, the likely conjugative elements. Under our experimental conditions, the preferred partner of Mia14 was *Cuniculiplasma divulgatum* (previously known as ‘G-plasma’), which is an abundant inhabitant in AMD. However, the distribution of archaea related to Mia14 (or to ARMAN-2 cluster) is diverse, sometimes non-acidic environments, emphasises their higher plasticity and ability to adapt to the broader range of environmental conditions. This broader distribution of ‘ARMAN’-related organisms in other environments also suggests that *Cuniculiplasma* spp. may not necessarily be the exclusive partner (host) for ARMAN-2-like organisms.

Mia14 is characterised by a very rudimentary metabolic capability. It is even devoid of minimal sets of enzymes required for biosynthesis of both types of nucleotides (purine and pyrimidine) and of 12 out of 20 amino acids (lysine, methionine, arginine, asparagine, alanine, aspartate, leucine, isoleucine, threonine, phenylalanine, tyrosine and tryptophan). Biosynthetic pathways for vitamins and cofactors (B1, B2, Coenzyme A, Coenzyme PQQ, B6, B12, heme, methanopterin, and ubiquinone/ menaquinone) are incomplete.

In Mia14, all glycolytic enzymes are missing. The majority of enzymes for the pentose-phosphate pathway and the entire TCA cycle are also absent. On the other hand, the non-phosphorylating ED pathway of glucose oxidation is present. Additionally, fatty acid metabolism and beta-oxidation, folate cycle, phospholipid biosynthesis, aminosugar metabolism, glycine and serine catabolism pathway, urea cycle, and YVTN domain found in bacteria)-like adhesion module. Six of these surface layer proteins (MIA14_0152, _0793 and_0946) require almost 2.5% of the whole genome. We identified two domain types in surface layer proteins displaying the PKD superfamily fold and beta-propeller Kelch and YVTN β-repeat domains fused to CARDB (cell adhesion related domain found in bacteria)-like adhesion module. Six of these surface layer proteins are predicted to be gained from various methanogenic and acidiphilic euryarchaea and the members of ‘TACK’ superphylum. As previously hypothesised, the expansion of proteins containing PKD and YVTN domains indicates their function in cell–cell interactions. Thus, we propose that the very rudimentary metabolic capability of Mia14 indicates a *Cuniculiplasma*-associated lifestyle and that numerous systems such as Type IV pili, surface proteins and membrane channels provide an interface for the exchange of metabolites, energy, macromolecules including DNA between Mia14 and its host.

**Methods**

**Sample proceedings.** Samples from sediments and water of acidic streamer were taken for the establishment of enrichment cultures in March of 2011 from copper-containing sulfidic ores, Parys Mountain, Anglesey, North Wales, UK (53°23′13.6″N 4°20′58.6″W). The enrichment cultures were supplemented with yeast extract and glucose each at concentrations of 0.1% (w/vol) and grown at pH 1.12 and 37°C in AR medium.

DNA was extracted by G’NOME DNA Kit (MP Biomedicals). For the metagenomic study and second series of enrichment cultures set-up for...
CARD-FISH experiments, sediments and water were collected in July, 2014 from the same sampling spot as in March, 2011. The metagenomic DNA was isolated with DNA Power Isolation Kit for Soil (MoBio).

DNA concentrations in all cases were measured using Varian Cary Eclipse fluorescence spectrophotometer using Quant-iT DNA Assay Broad Range Kit (Life Technologies).

**Genome sequencing and annotation.** The genomes were sequenced, assembled and annotated at Fidelity Systems, Inc. (Gaithersburg, MD), as previously reported. Final assemblies provided ca. 364 and 561-fold coverages for strain S5 and PM4, respectively, while Mia14 genome was covered 42-fold. GIs were inspected using Island Viewer 3 using two different algorithms: IslandPath-DIMOB, based on the analysis of mobile element-related genes and dinucleotide distribution biases, and SIGI-HMM, exploiting biases of codon usage implementation of a hidden Markov model approach. In most cases, after manual inspection of taxonomic affiliation of best blast hits of predicted horizontally transferred proteins, both predictions were considered as GIs. Analysis of proteins shared between Mia14 and C. divulgatum S5, but absent in C. divulgatum PM4, was performed by ProteinOrthoV5 using default parameters (10−8 blast e-value, 50% minimal query coverage and 25% minimal percent identity).

Based on the genomic data, the specific primers for the detection of Mia14-related organisms in enrichment cultures were: 5′-3′ F Mic (GCTTGGCAATTAAATGCTGGG) and R Mic (ATCTTGCCAGCCTATC CCCAG).

**Metagenome sequencing of Parys Mountain community.** For sequencing of the Parys Mountain acidic streamer metagenome, both pair-end and mate-paired DNA libraries were used. Paired end library was prepared from 400 ng of environmental DNA with NEBNext Ultra DNA library preparation kit (New England Biolabs, Ipswich, USA) according to the manufacturer’s instructions to obtain 12–14 million reads. Mate-paired libraries were prepared using Nextera Mate Pair Library Prep Kit (Illumina Inc., San Diego, CA, USA) using gel-free protocol supplied by manufacturer. Both libraries were sequenced with 2 x 250 bp reads with MiSeq Personal Sequencing System (Illumina Inc., San Diego, CA, USA). After sequencing, all reads were subjected to stringent quality filtering with CLC Genomics Workbench 8.5 (Qiagen, Germany). After filtering, overlapping paired-end library reads were merged with SeqPrep tool (https://github.com/jstjohn/SeqPrep) resulting in 4,110,617 single reads and 7,539,176 read pairs. Mate-paired reads were treated with NextClip tool, resulting in 663,171 read pairs with mean insert size of 2170 bp. Reads were assembled with metaSPADES, resulting in metagenomic assembly of about 200 Mb of total length consisting of 93,342 contigs with N50 of 3295.

For the binning for metagenomic contigs they were aligned against NCBI non-redundant protein database using DIAMOND in ‘blastx’ mode with e-value of 10−3. Results of the alignment were imported to MEGAN 6.4.2248 with default settings adjusted as follows: min score—80, top percent—10 min support—20. Binning by MEGAN was performed using default settings. After the initial automatic binning step, additional manual inspection was performed. In particular, contigs with ambiguous taxonomic affiliation, characterised by mixed blast hits were either reassigned to a bin of a higher taxonomic level or moved to the 'Unassigned' bin. Contigs related to Mia14-related lineages were identified manually using blasting their genome sequences with blast against the local Parys Mountain metagenomic contigs nucleotide blast database.

For the calculation of the taxon abundance, all metagenomic reads were mapped to the contigs with Bowtie 2249. Total length of all sequencing reads mapped to every particular bin was calculated with samtools25. Abundance was calculated as a ratio between total lengths of all sequencing reads to the average genome size of the corresponding taxon (based on NCBI genomes database). Relative abundance value which was used for the Fig. 1 was calculated as ratio of bin abundance to the sum of bin abundances.

**CARD-FISH.** Samples were fixed for 1 h at room temperature with pre-filtered formaldehyde (final concentration 2% vol/vol). Sample (diluted from 10−3 to 10−3, according to cell concentrations) was filtered through 0.22 μm (0.25 mm) polycarbonate membranes (New Technologies Group Srl, NTG). Cell permeabilisation was performed by incubation for 1 h with lysozyme (10 mg ml−1 in TE buffer pH 8.0) followed by incubation with a chomopeptidase for 30 min (5 mg ml−1), both at 37°C. Filters were cut into sections and cells were hybridised with universal horseradish peroxidase (HRP)-labelled oligonucleotide probes for Eubacteria (UB383 1Ⅱ, 1Ⅲ probe mix)26,27 to check for bacterial presence and for Archaea (Arch91528). Absence of unspecific hybridisation was controlled by implication of the non-specific probe NON338. The CARD-FISH probes specific for membranes and pigments (order Thaumarchaeota (Thaum888R), of family Cuniculiplasmae (Cpnm-1100R) and of Ca. Manarchaeas acidiphilum Mia14 (ARM-MIA1469R) were designed through this study. Detailed information about the probes is given in Supplementary Table 3. Intracellular peroxidase was inhibited by treatment with 1% H2O2 at room temperature for 20 min. For signal amplification, tyramide Alexa488 and Alexa594 were used26,28. The filter sections were counter-stained with 4,6-diamidino-2-phenylindole (DAPI) (2 μg ml−1) in a four-to-one ratio of Citifluor (Citifluor Ltd, Leicester, UK): Vectashield (Linaris GmbH, Wertheim-Bettingen, Germany). At least 200 DAPI-stained and Alexa-positive cells were counted in a minimum of 10 fields under an AXIOPLAN 2 Imaging microscope (Zeiss, Germany).

**Sequence analysis and evolutionary reconstructions.** Protein coding genes of Mia14 were assigned to archaeal Clusters of Orthologous Groups (arCOGs) as follows: PSSMs derived from arCOG alignments were used as PSI-BLAST queries in a search against a database of archaeal proteins with e-value cutoff of 10−5. Proteins (fragments) and Alexa338 were assigned to arCOGs with the highest-scoring hit.55 Also, sequences of the 56 ribosomal proteins universally conserved in archaea66 from 285 organisms with completely or almost completely sequenced genomes were aligned using the MUSCLE program. Alignments were concatenated; the phylogenetic tree was reconstructed using the FastTree program8 with WAG evolutionary model and gamma-distributed site rates.

Manual curation of automatic functional predictions was performed according to the recent protocol.56 In particular, the proteins of central carbohydrate pathways (Embden-Meyerhoff and Gluconeogenesis, ED, pentose-phosphate, TCA) including currently known archaeal modifications56 were searched by BLAST (with a consciously low e-value cut-off = 1.0 to avoid loss of distantly related sequences) of sets, including several amino-acid sequences of biochemically characterised (mainly, that with ‘Evidence at protein level’ in Swissprot database) archaeal and bacterial proteins against the genome (BLASTn) or in silico translated proteome (BLASTp) of Mia14. If no hits with all queries were found the protein was regarded as absent. If all/many of proteins of the pathway were absent the pathway was regarded as absent. If any BLAST hits were obtained, these sequences were BLASTed against Uniprot and Swissprot (e-value threshold = 0.01) and resulted hits analysed. The co-localisation of genes for a particular pathway was also taken into account.

arCOG phylogetic patterns of the 15 ‘DPANN’ group genomes were analysed using the COUNT program as described previously57. A matrix with the numbers of orthologs in the given arCOG in the given organism and the tree of the corresponding genomes were used to estimate the parameters of a phylogenetic birth and death model with gamma-distributed gain, loss and duplication rates.54 The model produces posterior probabilities for the presence or absence of a gene in ancestral genomes as well as the probabilities of gene gains and losses on all tree branches, providing a comprehensive picture of these events in the evolutionary history of the ‘DPANN’ group. The reconstructed ‘DPANN’ group ancestor was compared to the previously reconstructed last common ancestor of all Archaea63. To identify actual arCOGs in three groups (likely present, lost, gained) probability of each event more or equal 50% has been chosen for each lineage of interest.

Taxonomic affiliations for proteins, encoded in the 10 out of 15 ‘DPANN’ genomes (to the exclusion of the three genomes in the Nanoarchaeota archaean SCG CAA0011-G17 lineage and two genomes in the ‘Candidatus Halorovovirus’ lineage that have close relatives within the ‘DPANN’ group) was assessed by running protein BLAST search other archaeal genomes. The database contained 704,591 proteins from 286 complete and nearly complete archaeal genomes, available in GenBank and the protein set encoded by Mia14. The top BLAST hit (e-value threshold of 10−5) outside of the self genome was recorded as an approximate indication of the taxonomic affiliation of the protein.

**Data availability.** Sequence data determined in this study are available at NCBI under BioProject Accession PRJNA335339. Genome sequencing and assembly are deposited in the GenBank under Accession CP019964. Metagenomic reads and contigs were submitted to MG-RAST and can be provided from the corresponding author upon request.

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Author contributions

O.V.G. and P.N.G. conceived the research. O.V.G., S.V.T., K.S.M., Y.I.W., S.N.G., A.A.K., I.V.K., T.Y.N., M.F., M.M.Y. and P.N.G. did the genome analysis. V.L.C. and E.A. did the CARD-FISH experiment. O.V.G. drafted the manuscript. S.V.T., K.S.M., Y.I.W., S.N.G., A.A.K., I.V.K., M.M.Y. and P.N.G. contributed to the manuscript writing.

Additional information

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