The importance of biofilm formation for cultivation of a Micrarchaeon and its interactions with its Thermoplasmatales host

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Micrarchaeota is a distinctive lineage assigned to the DPANN archaea, which includes poorly characterised microorganisms with reduced genomes that likely depend on interactions with hosts for growth and survival. Here, we report the enrichment of a stable co-culture of a member of the Micrarchaeota (Ca. Micrarchaeum harzensis) together with its Thermoplasmatales host (Ca. Scheffleriplasma hospitalis), as well as the isolation of the latter. We show that symbiont-host interactions depend on biofilm formation as evidenced by growth experiments, comparative transcriptomic analyses and electron microscopy. In addition, genomic, metabolomic, extracellular polymeric substances and lipid content analyses indicate that the Micrarchaeon symbiont relies on the acquisition of metabolites from its host. Our study of the cell biology and physiology of a Micrarchaeon and its host adds to our limited knowledge of archaeal symbioses.

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n 2002, Huber and colleagues described a novel nano-sized archaeon, *Nanoarchaeum equitans*. Later, metagenomic data of environmental samples revealed that the Nanoarchaeota are part of a tentative superphylum of nano-sized archaea referred to as DPANN – an acronym of its first members’ lineages, the Diphtheriatores, Parvarchaeota, *Aemigrarchaeota*, Nanoarchaeota, and Nanohaloarchaeota. Most DPANN representatives have reduced genomes and are thought to comprise a diversity of potential archaean symbionts. Besides the name-giving phyla, the DPANN also include the Woese- and Pacearchaeota, Huberarchaeota, Micrarchaeota, Altiarchaeota, Undinarchaeota and Mamarchaeota as well as several so far undefined phyla. Nano-sized archaea are globally distributed and can comprise non-negligible proportions of microbial communities. Yet only a few representatives have been enriched under laboratory conditions. Current genomic data suggest that most DPANN archaea have reduced genomes, limited metabolic capabilities and various auxotrophies and might depend on interactions with other community members. The extent of genome reduction varies within the DPANN members. For instance, marine Nanoarchaeota were characterised by highly reduced genomes of about 0.5 Mbp and seem to represent ectoparasites that are strongly dependent on their host. On the other hand, the first members of the Nanohaloarchaeota and Micrarchaeota have larger genome sizes and seem metabolically more flexible. Yet cultivated Nanohaloarchaeota representatives are nevertheless host-dependent. While recent work has provided more insights into symbiotic interactions characterising certain representatives of the DPANN, additional model systems remain to be established. We have recently succeeded in enriching a member of the Micrarchaeota in a community of four different microorganisms. Here, we report the isolation of a stable co-culture of this Micrarchaeon together with its host, a previously unknown member of the *Thermoplasmatales*, as well as the isolation of the latter. This allows us to conduct experiments aiming at understanding the interaction of the two organisms and the response of the *Thermoplasmatales* member to growth in coculture with the Micrarchaeon.

**Results and discussion**

**Isolation of the *Ca. Micrarchaeum harzensis*-Ca. *Scheffleriplasma hospitalis* co-culture and pure culture of *Ca. Scheffleriplasma hospitalis*.** We have previously enriched for a putative DPANN symbiont belonging to the Micrarchaeota, i.e., *A. DKE*, in a culture, which besides the inferred host *Thermoplasmatales* archaea *B. DKE*, also contained a *Cuniculiplasma*-related archaeon referred to as *C. DKE* and a fungus (with *Acidothrix acidophila* as its closest related organism). In order to obtain a better understanding of the interactions between the symbiont and its hosts, a co-culture of the putative symbiont and host organisms was generated by eliminating all other organisms. *C. DKE* represented the minority of the archaea in the enrichment culture and is closely related to an organism reported to have a low pH optimum of 1.0–1.2. Hence, it was possible to eliminate *C. DKE* by transferring the culture in media with a pH of 2.5, which exceeds its optimal pH range, while still supporting the growth of *A. DKE* and *B. DKE*. Secondly, the fungus, which was isolated and via sequence analysis revealed to be most closely related to an isolate from acidic soils in the Czech Republic, was successfully eliminated by incubating the enrichment cultures at 37 °C over three consecutive culture transfers. Microscopic analysis, as well as 16S rRNA and ITS region gene analyses confirmed the absence of those contaminants. From the previous microscopic analysis, it was known that *A. DKE* thrives together with *B. DKE* in biofilm-like structures. We discovered that it was possible to enhance the biofilm formation of the host by lowering the pH-value of the medium from 2.5 to 2.0, which led to robust biofilm formation of the co-culture (Supplementary Figure 1). At the same time, we were able to isolate the host *B. DKE* by enriching at pH 2.5 for planktonic organisms. The composition of cultures was verified via PCR with organism-specific primers and CARD-FISH, periodically.

Based on the isolation of the *Thermoplasmatales* host and the reconstruction of complete genome sequences of both organisms, we propose the names *Candidatus Micrarchaeum harzensis* sp. nov. (N.L. masc./fem. adj. *harzensis*, pertaining to the German region of the Harz Mountains, where the organism was isolated) and *Candidatus Scheffleriplasma hospitalis* gen. nov. sp. nov. (Scheffler.i.plas’ma. N.L. gen. masc. n. *scheffleri* of Scheffler, named in honour of the geologist Dr. Horst Scheffler and in recognition of his work on mine geology and commitment to our work; Gr. neut. n. *plasma* something shaped or moulded; hos.pi.ta’lis. L. masc. adj. *hospitalis* relating to a guest, hospitable, referring to its ability to serve as a host for *Candidatus Micrarchaeum harzensis*).

Previous attempts to purify mesophilic members of the Micrarchaeota with their respective hosts yielded in relatively diverse enrichments or led to the disappearance of the symbiont after some time of incubation. Here, we used physiology-informed strategies for deselecting against additional community members and show that selection for biofilm formation of the *Thermoplasmatales* host population was the critical factor for obtaining a stable co-culture.

**Characterisation of *Ca. Micrarchaeum harzensis*-Ca. *Scheffleriplasma hospitalis* co-culture.** The pure culture of *Ca. Scheffleriplasma hospitalis* and the *Ca. Micrarchaeum harzensis*-Ca. *Scheffleriplasma hospitalis* co-culture reduced ferric iron (Fe(III)) to ferrous iron (Fe(II)) during growth indicating that *Ca. Scheffleriplasma hospitalis* is a dissimilatory ferric iron reducing organism. During growth of the co-culture *Ca. Micrarchaeum harzensis* was able to reach higher cell densities compared to *Ca. Scheffleriplasma hospitalis* and maintained a higher cell number for several weeks, before dropping under the *Ca. Scheffleriplasma hospitalis* cell number at the end of the growth curve (Fig. 1). Furthermore, *Ca. Scheffleriplasma hospitalis* showed a slower growth rate compared to *Ca. Micrarchaeum harzensis*. Source data are provided as a Source Data file.

![Fig. 1 Growth curve of a *Ca. Micrarchaeum harzensis*-Ca. *Scheffleriplasma hospitalis* co-culture.](https://example.com/growth_curve.png)
increase and decrease in cell number compared to the Micrarchaeon.

To examine the impact of the co-cultivation with the Micrarchaeon on \textit{Ca. Scheffleri}plasma hospitalis its growth characteristics were compared between pure and co-culture under otherwise identical conditions (growth curve of \textit{Ca. Scheffleri}plasma hospitalis pure culture in Supplementary Fig. 2). Cells of the \textit{Thermoplasmatales} member showed a similar growth and ferric iron reduction in pure culture in comparison with the co-culture. Also, the doubling time of \textit{Ca. Scheffleri}plasma hospitalis in co-culture (7.49 ± 1.45 days, \(n = 3\)) was not significantly different (unpaired, two-sample \(t\)-test, two-sided, significance level 0.05) to the doubling time of the organism cultivated without its interaction partner (7.19 ± 3.80 days, \(n = 3\)). Hence, growth of the host seems to be little affected by the Micrarchaeon under the tested conditions.

Genomic potential of \textit{Ca. Micrarchaeum} harzensis and \textit{Ca. Scheffleri}plasma hospitalis. DNA of the co-culture containing \textit{Ca. Micrarchaeum} harzensis and \textit{Ca. Scheffleri}plasma hospitalis was sequenced using a combination of PacBio and Illumina sequencing. For comparison, Illumina sequencing was also performed on the \textit{Ca. Scheffleri}plasma hospitalis pure culture. The two organisms have circular chromosomes of 1,959,588 base pairs (bp) (\textit{Ca. Scheffleri}plasma hospitalis) and 989,838 bp (\textit{Ca. Micrarchaeum} harzensis), and GC contents of 44.4% and 45.8%, respectively. The genomes of the pure \textit{Ca. Scheffleri}plasma hospitalis isolate and the strain within the co-culture were 100% identical which was important for the later comparative transcriptomic analysis. An analysis of clusters of orthologous groups (COGs) revealed that \textit{Ca. Micrarchaeum} harzensis contains more proteins with unknown function (29%; 300 putative proteins without an arCOG-assignment) relative to the overall number of genes compared to \textit{Ca. Scheffleri}plasma hospitalis (20%; 419 putative proteins without an arCOG-assignment).

The complete genome of \textit{Ca. Micrarchaeum} harzensis confirmed earlier findings\textsuperscript{15,18,22}, such as an extremely limited set of genes coding for proteins involved in central carbon metabolism. We could only detect one gene encoding a putative enzyme of the pentose phosphate pathway and two genes for putative enzymes of a glycolysis or gluconeogenesis pathway (Supplementary Data 1). However, we did identify a putative set of genes coding for enzymes for the conversion of glucose to glycerate, which together comprise four of the seven reactions of the non-phosphorylative Entner-Doudoroff pathway (Supplementary Data 1). The \textit{Ca. Micrarchaeum} harzensis genome also contains genes encoding enzymes for the conversion of pyruvate to acetyl-CoA, though we could not identify candidate proteins for reactions leading from glycerate to pyruvate. In agreement with our previous study, enzymes for almost all steps of the tricarboxylic acid cycle (TCA) were detected in \textit{Ca. Micrarchaeum} harzensis. Earlier publications proposed that the missing succinyl-CoA synthetase in Micrarchaeota genomes could be replaced by a methylisocitrate lyase, which generates succinate from methylisocitate\textsuperscript{10}. A corresponding gene, encoding methylisocitrate lyase, was detected in the \textit{Ca. Micrarchaeum} harzensis genome, as well (Supplementary Data 1). The thereby potentially completed TCA cycle, may through the production of NADH, fuel an electron transport chain and the generation of a proton gradient for ATP production. In particular, we detected gene clusters encoding a full NADH dehydrogenase and a membrane-bound \(A_{1}/A_{0}\) ATP synthase complex\textsuperscript{23} in the \textit{Ca. Micrarchaeum} harzensis genome. Moreover, we identified genes encoding one subunit of the cytochrome bc1 complex and two subunits of the cytochrome c oxidase (Supplementary Data 1). Although the organism might have the ability to conserve energy and produce reducing equivalents, it will be dependent on building blocks acquired either from the environment (or the culture medium) or from the partner organism \textit{Ca. Scheffleri}plasma hospitalis. For instance, \textit{Ca. Micrarchaeum} harzensis has major gaps in various biosynthesis pathways including for the production of amino acids; the few catalytic steps encoded by \textit{Ca. Micrarchaeum} harzensis comprise the synthesis of aspartate from oxaloacetate, glutamate from \(\alpha\)-ketoglutarate and phenylalanine from phenylpyruvate. Phenylpyruvate could be produced from tyrosine, which was taken up from the medium by the co-culture (see below). Other amino acid biosynthesis pathways could not be detected. Furthermore, genes encoding known amino acid transporters seem to be absent\textsuperscript{13,19} and DNA, RNA, and lipid biosynthesis pathways are incomplete (see below). Consequently, \textit{Ca. Micrarchaeum} harzensis may acquire certain metabolites or building blocks directly from its partner \textit{Ca. Scheffleri}plasma hospitalis through cell-cell interactions as seen in the \textit{Ignococcus hospitalis-Nanoarchaeum equitans} system\textsuperscript{24,25}. In turn, the dependency of \textit{Ca. Micrarchaeum} harzensis on growth in a biofilm (see above) may be due to the need to establish cellular contact with its host. Possible central metabolic pathways were summarised in a cell cartoon in Fig. 2.

Biofilm composition of pure and co-cultures. As the isolation experiments and our previous results point towards the importance of extracellular polymeric substances (EPS) for successful cultivation of \textit{Ca. Micrarchaeum} harzensis, we next investigated the composition of the EPS matrix in the co-culture as compared to the pure culture of \textit{Ca. Scheffleri}plasma hospitalis. To this end, the glycoconjugates were analysed with fluorescently labelled lectins, and the signals were correlated to the individual cell type by CARD-FISH analysis (Fig. 3). Lectins are complex proteins, which bind specifically to carbohydrate structures\textsuperscript{30}. In this study, 70 different fluorescently labelled lectins, which represent all commercially available lectins (Supplementary Data 2), were used to analyse the EPS in pure and co-cultures.

Among the tested lectins, only those specific to galactose- and mannose-related conjugates bound the extracellular matrix of the co-culture and of \textit{Ca. Scheffleri}plasma hospitalis cells in pure culture (see Fig. 3 and Table 1). Notably, lectins, IAA, HHA, and PTA seemed to discriminate between the extracellular matrix of \textit{Ca. Scheffleri}plasma hospitalis in the presence or absence of \textit{Ca. Micrarchaeum} harzensis, suggesting a potential influence of \textit{Ca. Micrarchaeum} harzensis on growth in a biofilm of \textit{Ca. Scheffleri}plasma hospitalis. This may reflect the inability of \textit{Ca. Micrarchaeum} harzensis to build carbohydrate polymers or the production of less common polymers for which we did not have a lectin. It suggests that the detected signals on \textit{Ca. Micrarchaeum} harzensis are likely due to growth within the biofilm matrix of \textit{Ca. Scheffleri}plasma hospitalis.

Overall, the results indicate that \textit{Ca. Scheffleri}plasma hospitalis displays galactose and mannose on its cell surface and that these carbohydrates are also components of the co-culture EPS matrix. This is corroborated by the presence of transcriptionally expressed genes for metabolic pathways leading to UDP-glucose, UDP-galactose, GDP-mannose, UDP-N-acetylgalactosamine, and UDP-N-acetylglucosamine in the genome of \textit{Ca. Scheffleri}plasma hospitalis (Supplementary Data 3 and Supplementary Fig. 3).

Membrane lipids of \textit{Ca. Micrarchaeum} harzensis and \textit{Ca. Scheffleri}plasma hospitalis and lipid biosynthetic pathways. An analysis of the intact polar lipids (IPLs) of co-cultures of \textit{Ca. Micrarchaeum} harzensis and \textit{Ca. Scheffleri}plasma hospitalis revealed the archaeal isoprenoidal glycerol dibiphytanyl glycerol (GDGT) with zero cyclopentane rings (i. e., GDGT-0, also known as
calderarchaeol) as the main lipid, making up to 97% of the total intact polar lipids, together with a minor amount of archaeol (2,3-di-O-phytanyl glycerol diether) (Supplementary Data 4). Comparison of the results to the pure Ca. Scheffleriplasma hospitalias-culture revealed no differences in the relative abundance of the archaeal IPLs, suggesting that the Micrarchaeon Ca. Micrarchaeum harzensis has an identical membrane lipid composition as its host Ca. Scheffleriplasma hospitalis. Archaeal membrane lipids are formed by isoprenoid side chains linked through ether bonds to glycerol-1-phosphate (G1P, synthesised by the G1P-dehydrogenase) either as a bilayer of diethers (archaeoles) or a monolayer of tetraethers (i.e., GDGTS). The isoprenoid building blocks are synthesised by one of the four variants of the archaeal mevalonate (MVA) pathway, which differ with regard to the enzymes mediating the last three enzymatic steps (see ref. 28 for a review). Figure 4 shows an overview of the different MVA pathways known. The isoprenoid C20 units are linked to the G1P backbone through ether bonds by the geranylgeranyltransferase (GGGP) synthase. Also, a putative G1PDH (i.e., glycerol-1-phosphate dehydrogenase) could be identified. Genes encoding enzymes for the ether bond formation (i.e., GGGP and DGGGP synthase) are also encoded in the Ca. Scheffleriplasma hospitalis genome (see Supplementary Data 6). In contrast, Ca. Micrarchaeum harzensis has an incomplete mevalonate and archaeal lipid pathway (Supplementary Data 4 and 5). In particular, while an ancestor of Ca. Micrarchaeum harzensis and some other Micrarchaeota are likely to have acquired three key genes of the variant-III mevalonate pathway from Thermoplasmatales archaea (i.e., mevalonate-3-kinase (acCOG02937), mevalonate-3-phosphate-5-kinase (COG02074; M3P5K), and mevalonate-3,5-bisphosphate-decarboxylase (arCOG02937; MBD), characterising this pathway, while it lacks genes for a canonical mevalonate kinase (Supplementary Data 5 and 6) similar to other members of the acidothrophic Thermoplasmatales. Prenyltransferases found in the genome are farnesyl diphosphate synthase and geranylgeranyl diphosphate synthase. Also, a putative G1PDH (i.e., glycerol-1-phosphate dehydrogenase) could be identified. Genes encoding enzymes for the ether bond formation (i.e., GGGP and DGGGP synthase) are also encoded in the Ca. Scheffleriplasma hospitalis genome (see Supplementary Data 6). In contrast, Ca. Micrarchaeum harzensis has an incomplete mevalonate and archaeal lipid pathway (Supplementary Data 4 and 5). In particular, while an ancestor of Ca. Micrarchaeum harzensis and some other Micrarchaeota are likely to have acquired three key genes of the variant-III mevalonate pathway from Thermoplasmatales archaea (i.e., mevalonate-3-kinase (arCOG02937), mevalonate-3-phosphate-5-kinase (COG02074), and mevalonate-3,5-bisphosphate-decarboxylase (arCOG02937)) (Fig. 5). Ca. Micrarchaeum harzensis lacks a homologue of the hydroxymethylglutaryl-CoA reductase (Supplementary Data 5). Note, its genome does not provide any evidence for the presence of another variant of the mevalonate pathway (Supplementary Data 5). Furthermore, Ca. Micrarchaeum harzensis does not appear
to encode a G1PDH. One of the encoded geranylgeranyl reductase homologues of \textit{Ca. Micrarchaeum harzensis}, likely involved in lipid biosynthesis, also seems to be acquired by horizontal gene transfer (HGT) from \textit{Thermoplasmatales} (arCOG00570) (Fig. 5). The result is in line with a recent study which revealed that 16.3\% of the examined Micrarchaeota genomes contained a horizontally acquired mevalonate-3,5-bisphosphate-decarboxylase\textsuperscript{22}.

Together with our experimental data, the presence of an incomplete variant-III mevalonate and lipid biosynthesis pathways in \textit{Ca. Micrarchaeum harzensis}, indicates that this organism depends on lipids or precursors thereof from its host, similar to what has been previously described in the DPANN archaeon \textit{N. equitans}\textsuperscript{31} and likely other DPANN members such as \textit{Nanohaloarchaeum antarcticus}\textsuperscript{16}.

**Impact of growth in co-culture on the \textit{Ca. Schefflerioplasm}a hospitalis transcriptome.** To further elucidate the effect of the symbiont on its host, we compared gene expression levels of \textit{Ca. Schefflerioplasm}a hospitalis with and without co-cultivation with \textit{Ca. Micrarchaeum harzensis} under otherwise identical growth conditions. In particular, we compared three pure cultures with four co-cultures and analysed differentially expressed genes with p-values lower than 0.01, a false discovery rate of 0.01 and log2fold changes higher or lower than 2 or −2. This revealed 17 genes that were differentially expressed based on these criteria (Table 2).

Of the 17 differentially expressed genes, 16 genes were downregulated in the host \textit{Ca. Schefflerioplasm}a hospitalis in the co-culture compared to the pure culture of \textit{Ca. Schefflerioplasm}a hospitalis.

**Table 1** Results of lectin staining of co-culture of \textit{Ca. Micrarchaeum harzensis} and \textit{Ca. Schefflerioplasm}a hospitalis and \textit{Ca. Schefflerioplasm}a hospitalis pure culture. The table shows binding lectins, their abbreviation, carbohydrate-binding specificity and the strength of the signal in pure and co-culture (with + as weak binding, ++ as binding, and − as no binding). The experiment was repeated at least three times with equivalent results.

| Lectin | Single-sugar binding specificity | \textit{S. hospitalis} pure culture | Co-culture |
|--------|---------------------------------|-----------------------------------|------------|
|        |                                 | \textit{S. hospitalis} | \textit{M. harzensis} | \textit{S. hospitalis} | \textit{M. harzensis} |
| AAL    | β-fucose                        | +                        | −          |
| CA     | lactose>N-acetylgalactosamine>galactose and related sugars | ++                      | +          |
| GNA    | mannose                         | +                        | ++         |
| GS-I   | galactose, N-acetylgalactosamine | +                        | +          |
| HHA    | mannose                         | ++                       | −          |
| HPA    | N-acetylgalactosamine           | ++                       | +          |
| IAA    | not determined                  | ++                       | −          |
| PTA    | galactose, N-acetylgalactosamine | +                        | −          |
| RCA    | β-galactose, lactose            | +                        | +          |
| RPA    | N-acetylgalactosamine           | ++                       | −          |
| SSA    | β-N-acetylgalactosamine         | +                        | ++         |
| TKA    | galactose                       | −                        | +          |

**Fig. 3** Results of lectin staining of co-culture of \textit{Ca. Micrarchaeum harzensis} and \textit{Ca. Schefflerioplasm}a hospitalis and \textit{Ca. Schefflerioplasm}a hospitalis pure culture. Shown are example images of the microscopic analysis of the pure culture (left) and co-culture (right). The images show the results for staining with lectin CA (co-culture) and HHA (pure culture). Colour allocation: \textit{Ca. Schefflerioplasm}a hospitalis was stained with the general archaea probe Arch915 (blue) which does not stain \textit{Ca. Micrarchaeum harzensis}. \textit{Ca. Micrarchaeum harzensis} was stained using the Micrarchaeota-specific ARMAN980 probe (magenta). Lectin staining is shown in green. Scale bars represent 10 \(\mu\)m. The experiment was repeated at least three times with equivalent results.
Fig. 4 Schematic overview of mevalonate pathways and lipid metabolism. The different pathways are indicated with dark blue (MVA pathway I), light blue (MVA pathway II), violet (MVA pathway III) and magenta arrows (MVA pathway IV), respectively. Names of enzymes are boxed. Enzymes expressed in Ca. Micrarchaeum harzensis and Ca. Scheffleriplasma hospitalis are indicated with black and grey circles. Abbreviations are AMPD anhydromevalonate phosphate decarboxylase, DGGGPS 2,3-bis-O-geranylgeranyl glycerol-1-phosphate synthase, DMD diphosphomevalonate decarboxylase, GGPS 3-O-geranylgeranyl glycerol-1-phosphate synthase, GGPPS geranylgeranyl diphosphate synthase, G1PDH glycerol-1-phosphate dehydrogenase, IDI isopentenyl-diphosphate-delta-isomerase, IPK isopentenyl phosphate kinase, M3K mevalonate 3-kinase, M3P5K mevalonate 3-phosphate 5-kinase, MBD mevalonate-3,5-bisphosphate-decarboxylase, MVK mevalonate kinase, PMK phosphomevalonate kinase.
hospitals. Five of these genes are part of or are associated with the Ca. ScheffleriPlasma hospitalis archaellum complex. This suggests a decreased cellular motility of the host in the co-culture, which is in line with the observed tendency of Ca. ScheffleriPlasma to form a biofilm in the presence of Ca. Micrarchaeum harzensis. The closest relative of Ca. ScheffleriPlasma hospitalis, Cuniculiplasma divulgatum, does not contain any archaeellin-related genes, whereas the related ‘G-Plasmas’ are described to contain the full arl-operon (arBCDEFGHII). Furthermore, the gene for the hexuronic acid methyltransferase AglP, a component of the protein glycosylation machinery was downregulated. This may indicate an alteration of the glycosylation pattern of Ca. ScheffleriPlasma hospitalis, which could influence cell-cell-interaction in the presence of Ca. Micrarchaeum harzensis. It may also change the exopolysaccharide (EPS) matrix of Ca. ScheffleriPlasma hospitalis, which would explain the observed binding differences of some lectins (see above). Three other downregulated genes code for transport proteins that might be involved in the uptake of carbohydrate molecules. While the effect of the decreased expression level of these transporters is unclear, it may be speculated that it could lead to higher availability of certain metabolites in the medium and support growth of Ca. Micrarchaeum harzensis. Other downregulated genes encode hypothetical proteins, a putative aminopeptidase, a transposase-associated protein and an iron-sulfur-protein. Thus far, their potential impact on the interaction of Ca. ScheffleriPlasma hospitalis with Ca. Micrarchaeum harzensis remains unclear. Only the gene for a putative membrane protein was upregulated in the co-cultures compared to the pure culture. The encoded protein shows similarities to a protein domain of unknown function (DUF1648), which is part of a receptor protein in Bacillus subtilis. Membrane localisation was verified using the TMHMM algorithm which detected eight transmembrane helices in the protein.

Metabolomic analysis in the presence and absence of Ca. Micrarchaeum harzensis. Next, we performed a metabolomic analysis to compare metabolites produced in the co-culture to the pure host culture and determine whether the presence of Ca. Micrarchaeum harzensis changes the pattern of depleted and produced organic carbon compounds. Growth of both cultures was estimated based on the change in ferrous iron concentration over time. Ca. ScheffleriPlasma hospitalis pure culture showed in these experiments a shorter lag-phase compared to the co-culture (Supplementary Fig. 4). Therefore, we compared samples with equal ferrous iron concentration (3, 4, and 5 weeks of growth for the pure culture; 5, 6, and 7 weeks of growth for the co-culture) as they indicate similar growth phases. Figure 6 shows a heatmap representing all significantly different metabolites (p < 0.05 false discovery rate, FDR corrected) present in the samples at the various time points. While the majority of the detected compounds changed simultaneously in the pure- and the co-culture throughout the growth phases, some metabolites showed different patterns.

Tyrosine levels were found to decrease faster in the co-culture, whereas 4-hydroxyphenyl acetic acid, an intermediate of the tyrosine degradation pathway, increased. Hence, tyrosine degradation seems to be accelerated in the co-culture. Both Ca. Micrarchaeum harzensis and Ca. ScheffleriPlasma hospitalis encode enzymes catalysing the conversion of tyrosine to 4-hydroxyphenylpyruvate (Supplementary Data 1) and it seems possible that the presence of the Micrarchaeon leads to faster tyrosine depletion. Of note, the specific genes for the carbonic acid
and dehydrogenase reaction from 4-hydroxyphenylpyruvate to 4-hydroxyphenyl acetic acid are currently unknown. While 2-phenylglycine, a degradation product of phenylalanine that enters the ketoacidipate pathway, decreased, muconic acid, an intermediate of that pathway, increased in the co-culture. Still, we could neither identify genes involved in the degradation of 2-phenylglycine nor for a complete β-ketoadipate pathway in Ca. Micrarchaeum harzensis or Ca. Scheffleriplasma hospitalis (Supplementary Data 1). Moreover, the β-ketoadipate pathway operates under oxic conditions and the organisms were cultivated in the absence of oxygen. Hence, so far, we cannot explain the consumption and production of 2-phenylglycine and muconic acid, respectively. The analyses also revealed increased levels of gluconic acid in the co-culture, which may be a product of sugar-degradation, for instance from the biofilm EPS matrix. Genomic information indicates that Ca. Scheffleriplasma hospitalis is able to degrade glucose into glucono-1,5-lactone (KO: K18124), which can spontaneously be converted into gluconic acid (Supplementary Data 1). Both organisms possess the enzymes to convert gluconic acid to glycerate, as discussed above. Overall, our results reveal that the pattern of metabolites do not seem to deviate between isolated Ca. Scheffleriplasma hospitalis and co-culture and that the kinetics of consumption show minor differences towards faster consumption of some compounds in the co-culture. Hence, either both organisms employ similar pathways and compounds or, perhaps more likely, Ca. Micrarchaeum harzensis predominantly uses metabolites provided by Ca. Scheffleriplasma hospitalis. The latter assumption is in agreement with the sparsity of transporters encoded in the Ca. Micrarchaeum harzensis genome, which may indicate that it relies on direct cell-cell interaction for nutrient and metabolite exchange.

Evidence for direct cell-cell interactions between Ca. Micrarchaeum harzensis and Ca. Scheffleriplasma hospitalis. Due to the pleomorphic morphology and great variability in cell size of members of the Thermoplasmatales including Ca. Scheffleriplasma hospitalis, it was previously challenging to clearly distinguish symbiont and host cells on electron micrographs. Recently, Gfrerer et al. revealed that Ca. Micrarchaeum harzensis cells are characterised by the presence of an S-layer that can be observed on electron micrographs of freeze-etched, Platinum-Carbon shadowed samples. Here, using electron microscopy, we could show the attachment of several Ca. Micrarchaeum harzensis to Ca. Scheffleriplasma hospitalis cells, suggesting direct cell-cell interactions between these organisms (Fig. 7a), as was previously shown for N. equitans and I. hospitalis and in environmental studies containing Micrarchaeota. However, we also observed a large number of Ca. Micrarchaeum harzensis cells that were not in contact with their potential host organism (Fig. 7b), which is in agreement with observations from microscopic images of CARD-FISH-stained cultures. While we cannot exclude that this is (to some degree) a result of sample preparation, it is possible that growth in the biofilm enables a more dynamic interaction between Ca. Micrarchaeum harzensis and Ca. Scheffleriplasma hospitalis than observed for N. equitans and I. hospitalis, as the risk of detaching from the host is mitigated by growth within the biofilm matrix. Moreover, Ca. Micrarchaeum harzensis has a larger genome and in turn greater metabolic flexibility than N. equitans and may in turn be less dependent on permanent attachment to host cells. Finally, we detected several unattached Ca. Micrarchaeum harzensis cells in the process of cell division (Fig. 7c). This could either be an artefact of sample preparation or suggest that Ca. Micrarchaeum harzensis can store a sufficient amount of building blocks to divide without being in direct cell-cell contact with Ca. Scheffleriplasma hospitalis.

Cryo-electron tomography (cryo-ET) shows attachment sites of Ca. Micrarchaeum harzensis and Ca. Scheffleriplasma hospitalis. To further investigate the physical interaction and characteristics of the two organisms, we used cryo-ET investigations of specimens preserved in vitreous ice in a near-native environment. Cryo-ET images of Ca. Micrarchaeum harzensis showed a cell membrane surrounded by a prototypical archaeal S-layer with clustered ribosomes abundant in the cytoplasm. We interpret the ribosome-free space as ‘nucleoid’, most likely containing the DNA. Cryo-ET further showed that Ca. Scheffleriplasma hospitalis cells are surrounded by a single cell membrane with clustered ribosomes abundant in the cytoplasm. Its cytoplasm is not homogeneous. At the sites of interactions of the two species, we detected filamentous structures between host and Micrarchaeon cells, which appear to contact the S-layer of Ca. Micrarchaeum harzensis cells but do not penetrate it. We also detected surface remoulding of

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Table 2 Differentially expressed genes of Ca. Scheffleriplasma hospitalis in pure culture in comparison to Ca. Micrarchaeum harzensis. The table indicates the gene ID, the description of the expressed protein, the log2-fold change and the p-value (<0.01; two-tailed Wald test, Bonferroni corrected).

| ID          | Description                                         | log2-foldchange | p-value |
|-------------|-----------------------------------------------------|-----------------|---------|
| Thermo_01860 | Hypothetical protein                                | −5.41           | 0       |
| Thermo_01798 | Hypothetical protein                                | −5.23           | 0       |
| Thermo_00025 | Hypothetical protein                                | −4.87           | 0       |
| Thermo_00444 | 41kDa archaellin                                    | −4.44           | 0       |
| Thermo_01859 | 41kDa archaellin                                    | −4.37           | 0       |
| Thermo_01877 | Hypothetical protein                                | −4.95           | 0       |
| Thermo_00445 | Putative archaellin-related protein C               | −3.98           | 0       |
| Thermo_00680 | Putative aminopeptidase 1                          | −3.96           | 0       |
| Thermo_00502 | Hexuronic acid methyltransferase AgIP               | −3.83           | 0       |
| Thermo_00446 | Putative archaellin-related protein D/E             | −3.58           | 0       |
| Thermo_01799 | Hypothetical protein                                | −3.23           | 0       |
| Thermo_01744 | Pink FeS protein                                    | −3.03           | 0       |
| Thermo_00581 | Trehalose/maltose import ATP-binding protein MalK    | −2.22           | 0       |
| Thermo_01800 | Hypothetical protein                                | −2.22           | 4,1592E−08 |
| Thermo_01811 | Trehalose/maltose import ATP-binding protein MalK    | −1.17           | 0       |
| Thermo_00582 | ABC-type transport system involved in multi-copper enzyme maturation, permease component | −2.12 | 0 |
| Thermo_01732 | Putative membrane protein                           | 2.12            | 0       |
Micrarchaeota and Ca. Schefleriplasma hospitalis cells, which were in direct contact with each other, specifically at the attachment site (Fig. 8d). This suggests that interaction of both organisms might be established by small filamentous structures leading to surface remodelling of the interacting cells. Direct contacts between the cytoplast of both interaction partners as described for N. equitans and I. hospitalis24,25 were not observed in the investigated samples so far. We addressed the question whether Ca. Micrarchaeum harzensis is able to divide independently and performed quantitative analysis by ultrastructural characterisation of associated and un-associated dividing Micrarchaeota cells. An analysis of a high-throughput cryo-EM dataset (2,000 images) revealed that 77.8% of the dividing cells are associated with a host cell (Fig. 8e) supporting the hypothesis that Ca. Micrarchaeum harzensis predominantly relies on host interactions for cell division.

Integration of results reveals a specific regulatory response towards co-culturing providing future research question. The detailed characterisation of our co-culture in comparison with pure host cultures indicate a specific regulatory response of Ca. Schefleriplasma hospitalis as a consequence of growth with Ca. Micrarchaeum harzensis. The combination of genomic analyses with comparative metabolomics, lipidomics and determination of EPS composition revealed that the growth of Ca. Micrarchaeum harzensis is dependent on interaction with its host Ca. Schefleriplasma hospitalis, whose growth seemed little impaired by the symbiont. Cryo-ET micrographs and comparative transcriptomics indicate that Ca. Schefleriplasma hospitalis might initiate the interaction by biofilm formation, change of EPS composition and development of a filamentous structure, which was observed in contact with Micrarchaeota cells. A physical interaction involving surface remodelling is established, which is likely crucial for the uptake of various metabolites and building blocks for, among others, membrane formation. Due to the limited metabolic capabilities of Ca. Micrarchaeum harzensis and its dependency on Ca. Schefleriplasma hospitalis for growth, it is very likely that the DPANN member initiates the interaction. The close cell-cell interactions between acidophilic Micrarchaeota and Thermoplasmales may also provide a route for horizontal gene transfer among these DPANN symbionts and their hosts.

It will be interesting to compare cell-cell interactions underlying this system to those characterising the various other members of the extremely diverse DPANN archaea and establish unique and common characteristics12-14,16,17,41. To this end, similar observations were published while this manuscript was under consideration41. A thermoacidophilic co-culture between a Micrarchaeon and a new isolate belonging to the genus Metallosphaera also revealed direct interactions between the Micrarchaeon and its host. Nevertheless, the results also reveal the physiological variability of Micrarchaeota members, as the co-culture enriched by Sakai et al. was not derived from a biofilm but acidic spring water and grew aerobically. Further studies will reveal whether these differences can be traced back to genomic or transcriptomic variations. On a broader scale, insights into symbiont-host interactions involving DPANN representatives will be important to improve our understanding of their role in ecological networks, considering their prevalence in most environments on Earth2,8,10,42.

**Methods**

**Culturing conditions.** The pure culture of Ca. Schefleriplasma hospitalis (JCM 39074) and the co-culture of Ca. Schefleriplasma hospitalis and Ca. Micrarchaeum harzensis were cultivated under anoxic conditions in a modified Picrophilus medium at a pH of 2.0 or 2.5 and at 22 °C. The pH of Picrophilus salt solution was adjusted with H2SO4 and oxygen was eliminated from the solution via boiling and subsequent exchanging of the headspace with N2. After autoclaving the medium was complemented with anoxic, sterile filtered solutions of yeast extract (final concentration 0.1%), casein hydrolysate (final concentration 0.1%) and ferric sulphate (final concentration 20 mM). Cultivation took place in anoxic flasks with a headspace containing 5% H2 and 95% N2. Transfers of the cultures were conducted with 20% of the pre-culture. The growth phase was assessed by following the

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**Fig. 6 Heatmap showing significantly different metabolite levels (p < 0.05 false discovery rate, FDR corrected) between corresponding growth phases of pure culture (Ca. Schefleriplasma hospitalis) and co-culture (Ca. Micrarchaeum harzensis-Ca. Schefleriplasma hospitalis).** Normalisation was done using a z-score, and significance was calculated by a two-tailed t-test. Source data are provided as a Source Data file.
reduction of ferric iron via the ferrozine assay. Furthermore, the cultures were monitored regarding their activity via CARD-FISH (see below) and regarding their composition via PCR with organism-specific primers (Supplementary Data 9). The cultures typically reached the exponential growth phase after incubation for 2–8 weeks.

CARD-FISH and lectin staining. Samples were fixed for 1 h in 4% formaldehyde, washed twice in phosphate-buffered saline (PBS), and stored at −20°C in 50:50 PBS/ethanol for CARD-FISH analysis or at 4°C in 100% PBS for lectin staining. The fixed cells were immobilised on a PTFE-coated slide (Thermo Fisher Scientific Inc., Schwerte, Germany). Inherent peroxidases were inactivated via a 10 min incubation in 0.1 M HCl, followed by a quick wash step with ddH₂O. Hybridisation of specific probes was carried out via incubating the dried cells with 0.3 ng/µL HRP-labelled probe in appropriate hybridisation buffer for 1.5 h at 46°C followed by three wash steps (quick dip in appropriate washing buffer, 15 min at 48°C in washing buffer, 15 min in saline sodium citrate (SSC) buffer). Amplification of signals involved a 15 min incubation of cells at 37°C with 0.5 µL fluorese-coupled tyramide (Thermo Fisher Scientific Inc., Schwerte, Germany) in 500 mL amplification buffer (4 mL 10 × PBS, 16 mL 5% NaCl, 4 g dextran sulphate, 0.4 mL 10% blocking reagent (Sigma-Aldrich, Steinheim, Germany), 19.6 mL ddH₂O) mixed with 5 µL H₂O₂. For subsequent wash steps incubation of the slide for 15 min in SSC was repeated once followed by a quick dip in ddH₂O. For double hybridisation the previous steps were repeated starting with the inactivation of peroxidases. Thereafter, the dried slides were treated with DAPI solution (1 mg/mL) for 5 min and subsequently incubated at ddH₂O for 2×1 min. At last, cells were mounted in embedding buffer (10 mL Citifluor (Citifluor Limited, London, UK), 2 mL Vectashield (Vector Laboratories, Burlingame, CA, USA), 1 mL phosphate-buffered saline (PBS)) and stored at −20°C until further analysis. Labelling was conducted using HRP-labelled 16S rDNA probes ARCH915 (Archaea domain, GTG CTC CCC CGC CAA TTC CT, 20% formamide, purchased at biomers.net GmbH, Ulm, Germany;25, TH1187 (Thermoplasmatales, GTA CTC ACC TGC GGT CAG A, 20% formamide, purchased at biomers.net GmbH, Ulm, Germany;26) and ARM089 (ARMAN, GCC GTC GCT TCT GGT AAT, 30% formamide, purchased at biomers.net GmbH, Ulm, Germany;27). For standard CARD-FISH staining, Alexa546 and Alexa647 fluorophores were used, and counterstaining was conducted with DAPI. CARD-FISH staining for lectin analysis was conducted using Alexa546 and Alexa647 fluorophores; see below for more details. Slides were visualised on a Leica DM 5500B microscope (objective lens 100×: HCX PL FLUOTAR, 1.4, oil immersion and objective lens 64×: HCY PL APO; eyepiece 10×: HC PLAN s (25) M), and images were taken with a Leica DFC 360 FX CCD camera and the corresponding Leica LAS AF 6000 software. Lectin staining was conducted according to a protocol of ref. 49. The positive lectin results were compared with CARD-FISH staining of the same slides, using a protocol of ref. 49. CARD-FISH staining was performed as described above with the following modifications: 1) all ethanol washing steps were omitted, as these would negatively affect the lectin staining; 2) cells were not counterstained with DAPI. After fixation and staining via CARD-FISH, the samples were dried at 37°C, and 100 µL lectin solution (0.1 mg/mL) was added and incubated for 30 min at room temperature in the dark. After washing with PBS solution, the slides were dried at 37°C and mounted in the same embedding buffer as used for CARD-FISH. Please refer to Supplementary Data 2 for an overview of lectins used. Imaging was conducted using a Zeiss Axiosvert 200 M fluorescence microscope equipped with the software Axios Vision 4.7.

DNA/RNA isolation and quantitative PCR analysis. Isolation of DNA for quantitative (qPCR) analysis was conducted with the Invisorb Spin Forensic Kit following the manufacturer’s instructions (Invitek, Berlin, Germany). DNA for Illumina and PacBio sequencing was isolated as described by ref. 50. In total 50 mL of three co-cultures were spun down at 15,000 × g for 10 min. 3 mL of Tris-EDTA (50 mM Tris-HCl pH 7.5, 50 mM EDTA pH 8.0) was used for resuspending of the cells. After addition of 0.6 mL STEP buffer (0.5% SDS, 50 mM Tris-HCl pH 7.5, 400 mM EDTA, 1 mg/mL protease K, 0.5% Sarkosy) the lysate was incubated at 50°C overnight. Subsequent phenol-chloroform extraction involved the addition of 3.6 mL phenol-chloroform isooamyl and hand-shaking for 5 min followed by a centrifugation step at 6000 × g for 10 min. The water fraction was transferred to another tube and the extraction procedure was repeated for a total of three times. Precipitation of DNA was conducted by adding an equal volume of isopropanol and 0.1× the volume of sodium acetate (3 M) to the solution and incubation at −20°C overnight. Precipitated DNA was centrifugated at 15,000 × g for 30 min, washed once with 30% ethanol (30 min 15,000 × g) and dried at room temperature.
before subsequent resuspension in TE buffer at 4 °C overnight. RNA isolation and library preparation was conducted by IMGM laboratories GmbH using the RNeasy Micro Kit (Qiagen, Hilden, Germany) and TruSeq Stranded total RNA LT kit according to the manufacturer’s instructions. Illumina, Berlin, Germany). All sequencing analyses were conducted with samples of exponentially growing pure host cultures as well as symbiont-host co-cultures. The cell number of the host cultures as well as symbiont-host co-cultures. The cell number of the

**Fig. 8 Cryo-ET study of a Ca. Micrarchaeum harzensis-Ca. Scheffleriplasma hospitalis co-culture.** Shown are (a) a tomographic slice through a Ca. Micrarchaeum harzensis cell, (b) a tomographic slice through a Ca. Scheffleriplasma hospitalis cell, (c) attachment sites and (d) cell surface remodelling of both organisms, as well as (e) results of a quantification of associated and un-associated dividing Ca. Micrarchaeum harzensis cells. Arrows highlight a filamentous structure (panel c) and attachment sites undergoing surface remodelling (panel d). Scale bars equal 100 nm. Additional Tomograms displaying Ca. Micrarchaeum harzensis and Ca. Scheffleriplasma hospitalis cells, as well as the corresponding movie to Fig. 8c are part of the supplements (Supplementary Movies 1–3). The experiment was performed two times with equivalent results.

**Metagenome analysis.** A SMRTbell™ template library was prepared according to the instructions from PacificBiosciences, Menlo Park, CA, USA, following the Procedure & Checklist – Greater Than 10 kb Template Preparation. Briefly, for preparation of 15 kb libraries, DNA was end-repaired and ligation overnight to hairpin adaptors applying components from the DNA/Polymerase Binding Kit P6 from Pacific Biosciences, Menlo Park, CA, USA. Reactions were carried out according to the manufacturer’s instructions. BluePippin™ Size-Selection to greater than 4 kb was performed according to the manufacturer’s instructions (Sage Science, Beverly, MA, USA). Conditions for annealing of sequencing primers and binding of polymerase to purified SMRTbell™ template were assessed with the Calculator in RS Remote, PacificBiosciences, Menlo Park, CA, USA. One SMRT cell was sequenced on the PacBio RSII (PacificBiosciences, Menlo Park, CA, USA), taking one 240-min movie. Libraries for sequencing on the Illumina platform were prepared to apply the Nextera XT DNA Library Preparation Kit (Illumina, San Diego, USA) with modifications according to ref. 51 and sequenced on Illumina NextSeq® 500. Genome assembly was performed by applying the RS_HGAP_Assembly.3 protocol included in the SMRT Portal version 2.3.0. The assembly revealed two major contigs. Potentially misassembled artificial contigs with low coverage and included in other replicons were removed from the assembly. Redundancies at the ends of the two major contigs allowed them to be circularised. Replicons were adjusted to smc (chromosome partition protein Smc) as the first gene. Error-correction was performed by mapping of the Illumina short reads onto finished genomes using the Burrows-Wheeler Aligner bwa 0.6.2 in paired-end mode using default settings52 with subsequent variant and consensus calling using VarScan 2.3.6 (Parameters: mpileup2cns--min-coverage 10 --min-reads2 6 --min-avg-qual 20 --min-var-freq 0.8 --min-freq-for-hom 0.75 --p-value 0.01 --strand-filter 1 --variants 1 --output-vcf 1)53. Automated genome annotation was carried out using Prokka 1.84.

**Genome annotations.** For further functional annotation, the protein files from the two complete genomes as well as 22 Micrarchaea and 11 Thermoplasmatales reference genomes (Supplementary Data 8) were compared against several databases, including the Archaeal Clusters of Orthologous Genes (arCOGs55; version from 2018), the KO profiles from the KEGG Automatic Annotation Server.
Phylogenetic analyses. We performed phylogenetic analyses of membrane lipid biosynthetic proteins, whenever homologues of relevant aCOGs were present in both Ca. Micrarchaeum harzensis and Ca. Schefleriplasmas hospitalis to assess the extent of horizontal gene transfer (HGT) affecting these proteins (Supplementary Data 7). In particular, we extracted homologues of corresponding arCOGs for all of these proteins from Ca. Micrarchaeum harzensis, Ca. Schefleriplasmas hospitalis, a reference set of 566 archaeal genomes (archaeal-only analysis) as well as from an additional set of 1029 eukaryotic and 100 eukaryotic genomes (universal analysis) (Supplementary Data 8). The reference genomes were annotated as described above. For the archaeal-only analysis, the individual homologues for each protein family were aligned using MAFFT v7-LINS-i and v7.407 (settings: –reorder2, trimmed with BMGE v1.12 (settings: –t 1 -a -m BLOSUM62 -h 0.535)). Phylogenetic trees were inferred using IQ-TREE (v1.6.10, settings: –m LG+C+F-R+F+RA –gtree reflap –topology-plot –drawtree –c 0.95 –bootstrap 1000 –cbs 0). For the universal analysis, MAFFT v7-LINS-i and v7.407 and MAFFT v7.407 were used to align protein families with less/equal (≤) or more (≥) than 1000 homologues, respectively. BMGE v1.12 was used for trimming all alignments (settings: –t AA -m BLOSUM30 -h 0.535) and phylogenetic trees were inferred using IQ-TREE (v1.6.10, settings: –m LG+C+F-R+F+RA –gtree reflap –topology-plot –drawtree –c 0.95 –bootstrap 1000 –cbs 0). Due to the large number of sequences affiliating with arCOG00570 when including bacterial and eukaryotic homologues (i.e., 13381), sequences for this column of the same material (Waters Chromatography Europe BV, Etten-Leur, Netherlands) and maintained at 30 °C. Archaeal CLs were eluted isocratically for 10 min with 10% B, followed by a linear gradient to 18% B in 20 min, then a linear gradient to 100% B in 20 min, where A is hexane and B is hexane/isopropanol (9:1, v/v ratio). The m/z values of the protonated molecules of archaeol and GDGTs were monitored. GDGTs were quantified by adding a C48 GDGT internal standard. A response factor derived from an archaeol:GDGT-0 standard (1:1) was used to correct for the difference in ionisation between archaeol and GDGTs.

The Bligh and Dyer extract (non-hydrolysed) and the acid-hydrolysed Bligh and Dyer extract were also analysed using ultra-high-performance liquid chromatography coupled to positive ion atmospheric pressure chemical ionisation/Time-of-Flight mass spectrometry (UHPLC-APCI/ToFMS) on an Agilent 1290 Infinity II UHPLC, equipped with an automatic injector, coupled to a 6230 Agilent TOF MS and Mass Hunter software. This additional analysis was performed to detect other archaeal lipids that were not included in the SIM method on the 6310 quadrupole MSD mentioned above. Separation of the archaeal lipids was achieved according to ref. 72 with some modifications using two silica BEH HILIC columns in series (2.1 × 150 mm, 1.7 µm; Waters Chromatography Europe BV, Etten-Leur, Netherlands) at a temperature of 25 °C. The injection volume was 10 µL. Compounds were isocratically eluted with 90% A and 10% B for the first 10 min, eluted with a linear gradient to 18% B in 20 min, followed by a linear gradient to 100% B in 30 min. A = hexane and B = hexane/isopropanol (9:1, v/v) and the flow rate was 0.2 mL/min. The typical injection volume was 10 µL of a 1 mg/mL solution (weighed dried Bligh and Dyer extract dissolved in hexane/isopropanol (99:1, v/v ratio)). The m/z values of the protonated molecules of archaeol and GDGTs were monitored. GDGTs were quantified by adding a C48 GDGT internal standard. A response factor derived from an archaeol:GDGT-0 standard (1:1) was used to correct for the difference in ionisation between archaeol and GDGTs.

Metabolome analysis. A pure culture of Ca. Schefleriplasmas hospitalis and a culture of Ca. Micrarchaeum harzensis and Ca. Schefleriplasmas hospitalis were inoculated as described above for 42 (pure culture) or 49 days (co-culture) until all available ferric iron was reduced and stationary phase was reached. Experiments were performed in triplicates. For metabolomic examination, 1 mL culture was sampled and stored at −80 °C until further analyses. Samples were taken every 7 days, along with samples for ferrous iron quantification to estimate growth, and samples for DNA extraction and CARD-FISH for further detection of Ca. Micrarchaeum harzensis and Ca. Schefleriplasmas hospitalis. Due to the low pH of the culture medium, 500 µL samples were amended by inducing sulphur precipitation through the addition of a spatula tip of CaCO3, to be completed at 5 min for 2000 rpm at room temperature. This treatment also led to cell lysis so that the analysis included also intracellular metabolites. After centrifugation for 5 min at 17,000 × g at room temperature, 50 µL of the supernatant was transferred to glass vials and dried under vacuum at 4 °C. Dried samples were stored at −80 °C until further analysis.

Online metabolite derivatization was performed using a Gerstel MPS2 autosampler (Muelheim, Germany). Dried metabolites were dissolved in 15 µL of 2% methoxyamine hydrochloride in pyridine at 40 °C under shaking. After 90 min, an equal volume of N-Methyl-N-(trimethylsilyl)trifluoroacetamide (MSTFA) was added and held for 30 min at 40 °C. One µL of the sample was injected into an SSL injector at 270 °C in splitless mode. Gas chromatography/mass spectrometry (GC/MS) analysis was performed using an Agilent 7890 A GC equipped with a 30-m DB-35MS 5% 2-Methylbutylphenyl/5% diphenyl column (Agilent Technologies, USA) in series, fitted with an FID detector and a MS detector (Waters Chromatography Europe BV, Etten-Leur, Netherlands) in series, fitted with an FID detector and a JEOL JMS-T100 spectrometer (Jeol, Cressington, Watford, UK; p < 10−5 mbar). At T = 176 °C, samples were evaporated using a cold knife (T = 90 K); after sublimation of about 400 nm of surface water, the samples were shadowed with Platinum-Carbon at an angle of 45 degrees (1.5 mm), and an additional layer of pure Carbon (about 15 nm; both by electron- beam deposition). Replics were deposited for 15 h, followed by three times on nitrogen. The samples were treated on 700 mesh (hex) grids and air-dried.

Structural analysis by electron microscopy. For freeze-initching, both a pure culture of Ca. Schefleriplasmas hospitalis and a co-culture of Ca. Micrarchaeum harzensis and Ca. Schefleriplasmas hospitalis were concentrated by centrifugation (3000 × g). The concentrated cell pellet (1.5 µL) was applied onto a gold carrier, coated with carbon followed by a layer of carbon dioxide, and the mounted grid was transferred to the column of the same material (Waters Chromatography Europe BV, Etten-Leur, Netherlands) and maintained at 30 °C. Archaeal CLs were eluted isocratically for 10 min with 10% B, followed by a linear gradient to 18% B in 20 min, then a linear gradient to 100% B in 20 min, where A is hexane and B is hexane/isopropanol (9:1, v/v). The flow rate was 0.2 mL/min. After 24 h, the cell grid was washed using 80% methanol/hexane. Source settings were identical to ref. 73. The typical injection volume was 10 µL of a 1 mg/mL solution (weighed dried Bligh and Dyer extract dissolved in hexane/isopropanol (99:1, v/v ratio)). The m/z values of the protonated molecules of archaeol and GDGTs were monitored. GDGTs were quantified by adding a C48 GDGT internal standard. A response factor derived from an archaeol:GDGT-0 standard (1:1) was used to correct for the difference in ionisation between archaeol and GDGTs. 

For electron microscopy analysis at 200 kV, a transmission electron microscope JEM-
2100F (JEOL GmbH, Freising, Germany), equipped with a F416 CMOS camera (TVIPS, Gauting, Germany) under control of SerialEM v. 3.8.6 was used.

**Cryo-EM sample preparation.** For cryo-EM grid preparation, 100 µl of anaerobically grown culture was removed aseptically from the growth flask. Then, 2.5 µl of the culture was immediately applied to a freshly glow-discharged Quantifoil R2/2 Cu/Rh 200 mesh or R3/5.1 Au 200 mesh grid, adsorbed for 10 s, blotted for 4–5 s and plunge-frozen into liquid ethane using a Vitrobot Mark IV (Thermo Fisher Scientific Inc., Waltham, Massachusetts, USA), while the blotting chamber was maintained at 100% humidity at 10 °C. For half of the prepared grids 10 nm protein-A gold (CMC Utrecht, Netherlands) was pelleted by centrifugation (100,000 g x 1 h, 4 °C) resuspended in 1:10 dilution of the growth medium and additionally added to the samples immediately prior to grid preparation.

**Cryo-ET data collection and analysis.** For tomographic data collection, the SerialEM software66 was used on a Titan Krios G3 microscope using the Quantum energy filter (slit width 20 eV) and the K3 direct electron detector running in counting mode. Tilt series (46 in total) were collected in two sessions with a defocus range of –5 to –12 µm, collected between 260° in a grouped dose symmetric scheme77 with a 2° tilt increment. A total dose of 100 or 177 e−/Å2 was applied over the entire series, and image data were sampled at a pixel size of 3.468 Å with 4 fractionated frames per tilt image. Unaligned tilt-movie frames were motion-corrected in IMOD and tilt series alignment using gold fiducials was performed in IMOD78. Tilt-series alignment without gold fiducials was performed using 250 ×250 pixel² patches with a fractional overlap of 33% within IMOD. Contrast transfer functions (CTFs) of the motion-corrected tilt series were estimated using CTFFind479 and tomographic reconstruction of CTF-corrected aligned tilt-series was carried out using the SIRT algorithm implemented within Tomo3D200. Figure panels containing cryo-ET images were prepared using Tomo3D200.

**Cryo-ET data analysis.** Cryo-ET images were aligned and motion-corrected in IMOD and tilt series alignment using gold fiducials was performed in IMOD78. Tilt-series alignment without gold fiducials was performed using 250 ×250 pixel² patches with a fractional overlap of 33% within IMOD. Contrast transfer functions (CTFs) of the motion-corrected tilt series were estimated using CTFFind479 and tomographic reconstruction of CTF-corrected aligned tilt-series was carried out using the SIRT algorithm implemented within Tomo3D200. Figure panels containing cryo-ET images were prepared using Tomo3D200.

**Data availability.** The genome sequences generated in this study including annotations have been deposited at NCBI Genbank under Accession Numbers CP060530 and CP060531. Raw reads of transcriptomic data generated in this study are available as SRA deposited at NCBI Genbank under control of SerialEM v. 3.8.6 was used. The genome sequences generated in this study including annotations have been deposited at NCBI Genbank under Accession Numbers CP060530 and CP060531. Raw reads of transcriptomic data generated in this study are available as SRA deposited at NCBI Genbank under control of SerialEM v. 3.8.6 was used.

**Code availability.** All custom scripts used are available at Zenodo [https://zenodo.org/record/3839790].

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Author contributions S.K. and S.G. isolated the organisms, performed growth experiments, interpreted metagenomic and metatranscriptomic data and conducted together with T.R.N. and U.K. the lectin staining experiments. S.K., S.G. and I.G. prepared the figures and wrote the manuscript. I.G. designed the scientific study. T.R.N. and U.K. contributed to the writing of the manuscript. B.B. and C.P. performed PacBio and Illumina sequencing and were involved in bioinformatic analysis of the data. A.V.K. and T.A.M.B. performed cryo-ET analysis, interpretation of the respective data and contributed to the writing of the manuscript. R.R. conducted together with S.G. the electron microscopic analysis, interpreted the data and was involved in writing the manuscript. T.A.M.B. performed the analysis of the Toll-like receptor expression and contributed to the writing of the manuscript. K.H. and K.H. contributed to metabolome analysis and writing of the manuscript. A.S. and N.D. analysed genome data, constructed phylogenetic trees and contributed to the writing of the manuscript. L.V. performed and interpreted the lipid analysis and contributed to the writing of the manuscript.

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