Functional prediction of proteins from the human gut archaeome

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1 Abstract

2 The human gastrointestinal tract contains diverse microbial communities, including archaea. Among them, 3 Methanobrevibacter smithii represents a highly active and clinically relevant methanogenic archaeon, being involved in 4 gastrointestinal disorders, such as IBD and obesity. Herein, we present an integrated approach using sequence and 5 structure information to improve the annotation of M. smithii proteins using advanced protein structure prediction and 6 annotation tools, such as AlphaFold2, trRosetta, ProFunc, and DeepFri. Of an initial set of 873 481 archaeal proteins, 7 we found 707 754 proteins exclusively present in the human gut. Having analysed archaeal proteins together with 87 8 282 994 bacterial proteins, we identified unique archaeal proteins and archaeal-bacterial homologs. We then predicted 9 and characterized functional domains and structures of 73 unique and homologous archaeal protein clusters linked the 10 human gut and M. smithii. We refined annotations based on the predicted structures, extending existing sequence 11 similarity-based annotations. We identified gut-specific archaeal proteins that may be involved in defense mechanisms, 12 virulence, adhesion, and the degradation of toxic substances. Interestingly, we identified potential glycosyltransferases 13 that could be associated with N-linked and O-glycosylation. Additionally, we found preliminary evidence for interdomain 14 horizontal gene transfer between Clostridia species and M. smithii, which includes sporulation stage V proteins AE and 15 AD. Our study broadens the understanding of archaeal biology, particularly M. smithii, and highlights the importance of 16 considering both sequence and structure for the prediction of protein function.

17 Introduction

18 In 1977, Woese and Fox, and colleagues discovered the kingdom of Archaebacteria, later renamed Archaea, revealing 19 a new branch in the tree of life [1]-[4]. The discovery of the Asgard superphylum and its close relationship with the 20 eukaryotic branch supports the notion of an archaeal origin for eukaryotes, yet ongoing debates continue regarding 21 whether the archaeal ancestor of eukaryotes belongs within the Asgard superphylum or represents a sister group to all 22 other archaea [5], [6]. Historically, archaea were associated with extreme environments but have since been recognized 23 for their general importance and prevalence [7], [8]. Their ability to thrive in extreme environments and to resist 24 chemicals is attributed, in part, to their unique cell envelope structures. In nature, archaea perform distinctive 25 biogeochemical functions, such as methanogenesis, anaerobic methane oxidation, and ammonia oxidation [9], [10]. By 26 employing diverse ecological strategies for energy production, archaea can inhabit a wide variety of environments [11]. 27 Archaea are also host-associated, such as on plants, in human and animal gastrointestinal tracts [12], [13], on human 28 skin [14], [15], in respiratory airways [16] and in the oral cavity [17]. Based on recent estimates, archaea comprise up 29 to 10% of the human gut microbiota [18].

30 Methanobrevibacter smithii, a ubiquitous and highly active methanogen in the human gut microbiome, has remarkable 31 clinical relevance and is relatively well annotated [19]. It plays an important role in the degradation of complex 32 carbohydrates, leading to the production of methane, which has significant physiological effects on human physiology. 33 Imbalances in the population of M. smithii have been implicated as factors contributing to gastrointestinal disorders 34 such as inflammatory bowel disease (IBD) [20], [21] and obesity [22]-[24]. Given the prevalence of M. smithii in the gut, 35 further research aimed at M. smithii is key to understanding their role in disease. Archaeal proteins, including those of 36 M. smithii, play a crucial role in adapting to diverse environments and showcase their unique biology. The knowledge 37 about diverse archaea, including novel species, in the human gut microbiome has expanded, underscoring their 38 significance [25]. Some host-associated taxa, like Methanomassilicoccales, have potential beneficial effects on human 39 health [26], while others like Methanosphaera stadtmanae have been linked to pro-inflammatory immune processes

[27]. Given the current interest in the role of archaea in human health and disease, understanding the archaeal proteomeis crucial for understanding the functional potential of archaea.

42 Studying archaeal proteins presents challenges both in experimental and computational aspects. Previous research by 43 Lyu et al. has highlighted the potential for biotechnological applications in various archaeal genera [28]. However, 44 genetic toolboxes for targeted genomic modifications are currently limited to mesophilic Methanococcus and 45 Methanosarcina genera [29]. While alternative methods like mass spectrometry-based searches exist, difficulties arise 46 from inaccurate predictions of protein coding sequences due to limited knowledge of ribosomal binding sites and 47 promoter consensus sequences [30]. Another unresolved challenge lies in the isolation and cultivation of archaea under 48 laboratory conditions, although recent progress has been made in this area [31], [32]. To overcome these challenges, 49 metagenomic sequencing has emerged as a promising approach to study archaea and their ecological relationships. 50 Metagenomics has enhanced our understanding of the archaeal branches within the tree of life [31]-[33] whereby 51 assembled sequences allow prediction of protein coding sequences (CDS) and their functional characterization in silico. 52 However, metagenome-assembled genomes (MAGs) face challenges in functional assignment due to incomplete 53 sequences and difficulties in predicting and annotating open-reading frames (ORFs) [34], [35]. Sequence-based protein 54 function annotation, commonly used but limited in cases of distant protein homologies, proves to be not particularly 55 effective [36]. Moreover, the databases containing information about archaeal proteins and functions are not 56 consistently updated, creating a twofold challenge in the sequence-based annotation of archaeal proteins. On one hand, Makarova et al. [37] report that archaeal ribosomal proteins L45 and L47, experimentally identified in 2011 [38] and pre-57 58 rRNA processing and ribosome biogenesis proteins of the NOL1/NOP2/fmu family characterized in 1998 [39] were not 59 added to annotation pipelines by 2019 and were labelled as 'hypothetical'. On the other hand, sequence similarity-60 based approaches fail to capture relationships between highly divergent proteins when aligned with a known database 61 protein [40]-[42]. Archaea, the least characterized domain of life, suffer from spurious and incorrect protein annotations 62 due to insufficient experimental data and outdated databases [43]. Furthermore, the study by Makarova et al. indicates 63 that a substantial proportion of genes within archaeal genomes, estimated to be between 30% and 80%, have not been 64 thoroughly characterized, leading to their classification as archaeal "dark matter" [37]. Poorly annotated proteins limit 65 our study of microbial functionality and their roles in specific processes. However, protein structure prediction represents 66 an alternative strategy to address the gap in sequence-function annotation [44]. It complements sequence-based 67 approaches, particularly when annotations are limited or conflicting across databases, by utilizing the conservation of 68 tertiary structure to infer functional roles [45], [46]. Advanced computational techniques, such as AlphaFold2 [47] and 69 trRosetta [48], offer accurate predictions of three-dimensional structures, thus providing valuable functional insights.

70 In this study, we introduce an integrated in silico approach that aims to refine the functional characterization of proteins, 71 thus enhancing the accuracy of protein annotations in the archaeon M. smithii. Having compared archaeal gut-specific 72 proteins to bacterial gut proteins, we found 73 unique and homologous archaeal protein clusters. Our approach 73 incorporates advanced protein structure prediction and annotation tools, such as AlphaFold2, trRosetta, ProFunc, and 74 DeepFri, into a comprehensive workflow. We predict and characterize the functional domains and structures of 73 gut-75 specific archaeal protein clusters. The predicted functions are linked to the adaptation to changing environments, 76 survival, and nutritional capabilities of M. smithii within the human gut microbiome. We additionally identified sporulation-77 related archaeal proteins, presumably horizontally transferred to archaea from *Clostridium* species.

78 Materials and Methods

79 Selection of gut-specific archaeal proteins

To select specific proteins of gut-associated archaea, we utilized archaeal MAGs obtained from the Genomes from Earth's Microbiomes (GEM) catalog [49] and the Unified Human Gastrointestinal Genome (UHGG) collection [50], along with bacterial MAGs from the UHGG collection (accessed in November 2020). Genomes were extracted based on available metadata and filtered by taxonomy to specifically target archaea.

Gene prediction was performed using Prodigal (V2.6.3: February, 2016) [51] on the archaeal and bacterial MAGs from
 the UHGG collection, while CDSs from the GEM catalog were downloaded from the provided source
 (<u>https://portal.nersc.gov/GEM</u>). Archaeal and bacterial proteins were further separately clustered using MMseqs2
 (MM2) (v12.113e3-2) [52], [53] (Fig. 1) with the following parameters: --cov-mode 0 --min-seq-id 0.9 -c 0.9.

To identify unique functionalities of gut-associated archaea, we selected proteins that are specific to the human gut and encoded by gut-associated archaea. MAGs were selected based on available metadata indicating their sampling location. First, we included protein clusters containing at least one protein from a MAG sampled in the human gut. We then excluded protein clusters that had proteins from MAGs sampled in other environments, as these were outside the scope of the study. The final selection included protein clusters where all proteins were encoded by MAGs sampled exclusively from the human gut.

From the selected gut-specific protein clusters, only those with complete KEGG annotations were included. Fully annotated MM2 clusters were additionally clustered with Sourmash (v4.0.0) into sourmash clusters [54], [55]. Archaeal protein clusters were categorized into two groups: those sharing KEGG Orthology identifiers (KOs) with bacterial proteins (prefix *h*) and those with unique KOs (prefix *u*) (Fig. 1).

98 Protein function annotation

Archaeal and bacterial proteins were annotated with KEGG orthologs (KOs) using Mantis (release 1.5.4) [56] (Fig. 1). 99 100 AlphaFold2 (AF) [47], [57] and TrRosetta (TR) [48] were used as structure-prediction tools. For each tool, the predicted 101 protein structure was then annotated separately. The TR-based model was annotated using templates with the highest 102 identity and coverage features. TR used a template for prediction if it met the criteria of confidence > 0.6, e-value < 103 0.001, and coverage > 0.3. The protein model generated by AF was submitted to the ProFunc (PF) [58] web server for 104 structure-based annotation. Sequence search vs existing PDB entries and 3D functional template searches sections 105 from the ProFunc report were used for structure-based protein annotation. Structure matches were selected according 106 to the reported highest possible likelihood of being correct as follows: certain matches (E-value <10-6), probable 107 matches (10-6 < E-value < 0.01), possible matches (0.01 < E-value < 0.1), and long shots (0.1 < E-value < 10.0). Only 108 certain matches were used for the functional assignment. DeepFri [59] was used as an auxiliary tool, providing broad 109 and general descriptions to verify or refute suggestions from AF and/or TR. DeepFri predictions with a certainty score 110 > 0.7 were considered. Our combined approach integrates multiple methods to enhance the resolution of functional 111 annotation, particularly for challenges faced by traditional sequence-based methods.

112 When TR- and AF-based annotations provided consistent results, the consensus was used as the final annotation of

the protein function. However, when the reports gave different results, we prioritized the result with highest confidence.

For instance, when the confidence of the model predicted by TR was *very high* and template matches were provided, and AF-based ProFunc reported a match with a lower confidence (anything but *certain match*), the template hit by TR

116 was used as the primary source for the annotation. The relationship between PF likelihood and TR Template Modeling

117 scores (TM-scores) generated in our analysis is shown in Table 1. Similarly, any protein with a TR template match was

118 considered as more reliable than an annotation with the *long shot* likelihood. In cases where there were no 3D functional

119 hits, TR annotation was given priority. In cases when PF and TR provided annotations with the same level of

- significance/likelihood, the protein structure with highest coverage and identity was chosen. In this case, we define
- 121 coverage as *coverage* feature in TR and the ratio $\frac{longest fitted segment}{query sequence length}$ as in PF, and for identity we take *identity* as in
- 122 TR and percentage sequence identity as in PF.

123 The appropriateness of an annotation was determined based on the extent to which the assigned function of a protein 124 was found to be directly relevant to archaea and supported by relevant literature. Any other annotations were classified 125 as incorrect. Following this initial step, *sensitivity* was calculated as *sensitivity* = $\frac{N_{str}}{N_{str}+N_{seq}}$, *specificity* as *specificity* = 126 $\frac{N_{seq}}{N_{seq}+N_{str}}$, *positive likelihood ratio (PLR)* as $PLR = \frac{sensitivity}{1-specificity}$, negative likelihood ratio (NLR) as $NLR = \frac{1-sensitivity}{specificity}$, 127 where N_{seq} and N_{str} are the numbers of correct sequence- and structure-based annotations, respectively.

¹²⁸ Protein relative occurrence calculation

129 Relative occurrence or frequency of protein functions in the groups of unique and homologous proteins was calculated. 130 The measure was calculated as the ratio of the number of proteins with a specific KO to the total number of proteins of 131 bacterial or archaeal proteins. For example, the relative occurrence of unique archaeal proteins annotated as *K20411* 132 (sourmash cluster 1) is: $\frac{N_{select}}{N_{total}} * 10^6$, where N_{selet} is the amount of proteins annotated with K20411 and N_{total} is the 133 total number of archaeal proteins. The reason for using a constant factor of 10^6 in the equation is to scale the values 134 and generate numbers better suited for graphical representation.

135 Gene expression analysis

To assess the expression of archaeal proteins in the context of human health and disease, gene expression was verified using an in-house dataset, by mapping metatranscriptomic reads of faecal samples of healthy individuals and patients with type 1 diabetes mellitus (T1DM) [60] to nucleotide sequences of genes of interest using bwa mem [61]. Mapping files were processed with SAMtools (v1.6) [62]. Mosdepth (v0.3.3) [63] was used to calculate mean read coverage per gene of interest.

141 Horizontal gene transfer analysis

142 To assess the stability of gene structures in *M. smithii* genomes, we conducted a horizontal gene transfer (HGT) analysis

using metaCHIP (v1.10.12) [64] on all *M. smithii* MAGs available in the included datasets. One *Methanobrevibacter_A*

144 *oralis* MAG derived from UHGG were also included for the comparison of the number of HGT events.

145 Gene synteny analysis

pyGenomeViz (v0.3.2) [65] was used to build gene synteny for all archaeal genes of interest. Gene coordinates predicted with Prodigal were used as an input. An interval of 10kb up- and downstream of the gene of interest was selected from the protein predictions. KEGG KOs were allocated based on the sequence-based annotations generated using Mantis [56]. The *M. smithii* type strain DSM 861 was used to assess the presence of genes from flanking regions of specific genes in an archaeal culture. In our study, we exclusively focused on *M. smithii*, as our analysis revealed that all the gut-specific proteins encoded by gut-associated archaea were encoded by *M. smithii*.

152 Phylogenetic analysis

In order to build phylogenetic trees for selective sourmash clusters, additional similar sequences were added from Uniprot [66] using BLAST (v2.0.15.153) [67] with default parameters on the consensus sequences representing sourmash clusters of interest, namely *h*9 and *h*20. Furthermore, Uniprot sequences and sourmash cluster sequences were used to build trees. Multiple sequence alignments were built using MAFFT (v7) [68] and trimmed with BMGE (v1.12) [69] using BLOSUM95 similarity matrix and the default cut-off 0.5. Maximum likelihood phylogenetic trees were built with IQ-TREE (v1.6.12) [70] and visualized using the R library *ggtree* (v3.6.2) [71].

159

160 Results & Discussion

161 Our study aimed to analyze the gut-specific proteins encoded by M. smithii in the human gastrointestinal tract. As we 162 focused on identifying archaeal unique proteins and archaea-bacterial homologs, we analysed gut-specific archaeal 163 and gut bacterial proteins together. Having compared the two subsets based on their sequence-based annotation, we 164 categorized archaeal gut-specific proteins into two groups: unique and homologous proteins. To annotate them, we 165 used KEGG KOs due to their consistent functional annotations across organisms and widespread usage. For structurebased functional assignment, we utilized a combination of structure prediction and annotation tools (Fig. 1), leveraging 166 167 the higher prediction accuracy of AlphaFold2 and the rapid and accurate *de novo* predictions obtained via TrRosetta. 168 Utilizing representative sequences of unique and homologous proteins, AlphaFold2 produced protein structures, and 169 subsequent functional annotations were accomplished by integrating ProFunc and DeepFRI. trRosetta was employed 170 to predict structures of unique and homologous proteins showing detectable homologous matches in the Protein Data 171 Bank, which were subsequently used for further structure annotation.

172 Enhancing annotations of proteins encoded by Methanobrevibacter smithii

173 To explore the uncharted functional space of *M. smithii*, we first selected gut-specific proteins of gut-associated archaea. 174 We collected the encoded proteins of a total of 1 190 archaeal and 285 835 bacterial MAGs, resulting in 873 481 175 archaeal proteins and 87 282 994 bacterial proteins (Fig. 1). We focused on proteins associated with archaea of the 176 human gut microbiome, which represented 37% (707 754 proteins) of all predicted archaeal proteins. These proteins 177 were grouped into 61 123 MMsegs2 clusters for archaea (≥2 proteins per cluster) and 1 967 480 MM2 clusters for 178 bacteria (≥10 proteins per cluster). By retaining fully annotated protein clusters, we obtained 55 117 archaeal MM2 clusters and 1 481 580 bacterial MM2 clusters. Using our proposed functional prediction strategy (Fig. 1A), we analyzed 179 180 the gut-associated archaeal proteins alongside bacterial proteins, resulting in 45 homologous sourmash clusters, *i.e.*, 181 shared between archaea and bacteria, and 28 unique sourmash clusters, i.e., composed exclusively of archaeal 182 proteins. The bacterial data served as a reference to distinguish unique proteins encoded and transcribed by archaea, 183 as well as archaeal proteins with homologs to bacterial ORFs. A summary of the annotations is provided in the 184 Supplementary Materials (Supp. Tab. 1-2).

185 All archaeal proteins from the abovementioned sourmash clusters were classified as M. smithii. We thus sought to 186 extend our knowledge of *M. smithii* by exploring functions that could have implications for human health and disease. 187 The investigation of the relative abundance of identified proteins and their associated processes revealed distinct types 188 of functions in unique and homologous protein clusters (Fig. 2). The most frequently identified functions in the unique 189 sourmash clusters were related to adaptation to changing environments and protection mechanisms, e.g., defense 190 against foreign DNA and oxidative stress, while processes such as RNA and DNA regulation, energy metabolism, and 191 cell wall integrity and maintenance were less represented (Supp. Tab. 3). Homologous sourmash clusters showed 192 frequent functions related to adaptation, various protection mechanisms, energy metabolism, and cell structural integrity 193 (Supp. Tab. 4). Analysis of fecal metatranscriptomic data confirmed the transcription of the majority of encoded genes, 194 with some unique and homologous genes exhibiting higher expression levels (Fig. 2). Two unique and 19 homologous 195 sourmash clusters with relatively high expression levels were identified, including genes associated with adaptation to 196 changing environments, defense against foreign DNA and oxidative stress, DNA/RNA regulation, and energy 197 metabolism, while the rest were unannotated (Fig. 2).

Our analysis demonstrated disparity in annotations between sequence- and structure-based approaches. Notably, 46%
 (13 out of 28) and 31% (14 out of 45) of the unique and homologous sourmash clusters, respectively, lacked structure-

200 based annotations, suggesting a reliance on sequence information for their functional annotation thus far. However, 201 literature searches suggest that the KEGG annotations may not provide reasonable or meaningful functional 202 assignments for most of these unannotated proteins. For instance, a protein annotated as mitochondrial import receptor 203 subunit TOM40 by KEGG is predicted to be a putative intimin/invasin-like protein based on its structure, which is more 204 relevant in the context of archaeal biology than being a eukaryotic protein involved in mitochondrial protein import. 205 Similarly, a protein annotated as Endophilin-A, a eukaryotic protein involved in membrane curvature, shows structural 206 similarity to PiIC, a type IVa pilus subunit of a prokaryotic adhesion filament. While the presence of eukaryotic proteins 207 in archaea is not surprising from an evolutionary perspective, the assignment of a protein to its evolutionary homolog 208 from a different kingdom may not provide precise functional assignment of protein function.

209 In general, the agreement between the sequence- and structure-based methods was limited, with 4% (1 out of 28) and 210 25% (11 out of 45) of the unique and homologous proteins showing consistent annotations, respectively (Supp. Tab. 3-211 5). The rest of the proteins exhibited disparity between sequence- and structure-based annotations, which was 212 assessed by comparing their reported functions. For example, unique sourmash cluster u24 yielded different 213 annotations using EGGNOG, KEGG, and Pfam databases which we used to potentially resolve disparities in the 214 annotations (Supp. Tab. 3). However, a consensus structure-based annotation identified it as polypeptide N-215 acetylgalactosaminyltransferase, providing additional annotation beyond sequence analysis. Similarly, the homologous 216 protein clusters h15-h18 had the same functional assignments as novobiocin biosynthesis protein NovC using KEGG. but structure-based annotation revealed further distinctions: h16 and h18 were classified as members of the LytR-217 218 Cps2A-Psr (LCP) protein family, h15 was annotated as 78 kDa glucose-regulated protein, and h17 remained 219 unannotated (Supp. Tab. 4). The incorporation of structural information in protein annotation enables the distinction 220 between closely related sequences, offering additional insights into protein function, which highlights the crucial role of 221 structural data in understanding protein functionality.

222 We further identified glycosyltransferases responsible for N and O-linked glycosylation from clusters h1-h6 as prevalent 223 archaeal gut-specific proteins. These proteins may contribute to the viability and adaptability of archaeal cells in the 224 gut. For instance, the most prevalent unique archaeal glycosyltransferase is 4-amino-4-deoxy-L-arabinose (L-Ara4N) 225 transferase, which is essential for the protection from environmental stress, symbiosis, virulence and resistance against 226 antimicrobial activity [72], [73]. Moreover, one of the six glycosyltransferases is a dolichyl-diphosphooligosaccharide -227 protein glycosyltransferase subunit STT3B (h5) which functions as an accessory protein in N-glycosylation and provides 228 its maximal efficiency [74]. Archaeal N-glycosylation is known to play an important role in the viability and adaptivity of 229 archaeal cells to external conditions such as high salinity [75], elevated temperatures [76] and an acidic environment 230 [77] while also maintaining the structural integrity of cells [78], [79]. Four out of the six identified glycosyltransferases are dolichyl-phosphate-mannose-protein mannosyltransferases 1 (POMT1), which are responsible for O-linked 231 232 glycosylation of proteins in eukaryotes. Another O-glycosylation associated protein, polypeptide N-233 acetylgalactosaminyltransferase, was found in the subset of unique archaeal proteins (u24). M. smithii has been found 234 to decorate its cellular surface with sugar residues mimicking those present in the glycan landscape of the intestinal 235 environment [80]. The presence of human mucus- and epithelial cell surface-associated glycans in M. smithii, along 236 with the coding potential for enzymes involved in O-linked glycosylation in archaeal gut species, suggests that M. smithii 237 cells might have the capability to emulate the surfaces of eukaryotic cells in the intestinal mucus. Beyond their structural 238 role in proteins, O-glycans can also act as regulators of protein interactions, influencing both interprotein and cell-to-239 cell communication processes involved in cell trafficking and environmental recognition [81].

Further findings suggest that 2-aminoethylphosphonate-pyruvate (2-AEP) aminotransferase, transthyretin-like protein and phosphoenolpyruvate-dependent sugar phosphotransferase system (PTS) system encoded by *M. smithii* contribute to energy metabolism. 2-AEP is an enzyme commonly found in bacteria [82] and is known to play a critical role in phosphonate degradation [83], which serves as an important source and production pathway for methane [84].

244 Additionally, cold-shock domains of Unr protein potentially provide M. smithii with adaptation strategies through stress-245 induced control of gene expression [85]. Furthermore, the predicted involvement of proteins such as the Specificity 246 subunit of type I restriction-modification EcoKI enzyme [86] and type II restriction endonuclease BgIII [87] suggests their 247 potential role in host defense strategies employed by M. smithii to protect themselves in the gut environment. 248 Additionally, it is conceivable that archaeal proteins may play a role in protecting against toxicity from other organisms 249 in the gut using propanediol utilization protein pduA [88]-[90], as well as acquiring genes of bacterial origin through 250 horizontal gene transfer. If this is the case, the presence of adhesin-like proteins in archaea could potentially enable 251 them to form symbiotic relationships with bacterial neighbors with diverse metabolic potentials [91]. Figure 5 provides 252 a schematic representation emphasizing specific proteins identified in this study, which could potentially play a 253 significant role in the functional dynamics of archaea within the human intestine. A more detailed description of all 254 identified *M. smithii* proteins is provided in Supplementary Materials.

255 Characterization of select proteins and gene structures in 256 Methanobrevibacter smithii genomes

257 To elucidate the level of conservation among the identified genes recovered in our analyses, we assessed the level of 258 genomic conservation within genomes of 2 strains of M. smithii, 2 strains of Ca. Methanobreviabcter intestini and the 259 related species Methanobrevibacter_A oralis as a reference. Ca. M. intestini has been recently classified as an 260 independent species within the Methanobrevibacter smithii clade [92]. We analysed HGT events and evaluated gene 261 structure stability. Using 1022 available MAGs, we noted an increase in HGT events between 319 genomes of two M. smithii strains: Methanobrevibacter A smithii and GCF 000016525.1 (based on GTDB classification) (Supp. Fig. 1). 262 263 Specifically, 2.6% of the MAGs (n=27) exhibited HGT events involving the transfer of approximately 10±3 genes to 264 other MAGs. Intriguingly, MAGs exhibiting HGT events were sampled in diverse geographical locations such as Austria, 265 France, the UK and the US. Our results suggest that the propensity of these MAGs to exchange genomic segments 266 may be attributed to similarities in their respective local environments [93], including dietary and lifestyle factors of the 267 individuals. Thus, it is plausible that exposure to similar diets or stresses may have influenced the evolution of these 268 MAGs via HGT along comparable trajectories. Conversely, the low occurrence of HGT events among the majority 269 (97.4%) of available M. smithii genomes indicates their overall genomic conservation and stability. This could be 270 explained by the fact that these MAGs were sampled from individuals living under similar dietary and lifestyle conditions. 271 Importantly, our findings support the concept of genomic stability in M. smithii, as we observed a high degree of 272 conservation in the flanking regions of the genes of interest across various M. smithii genomes. Through syntemy analyses, we found compelling evidence of conserved synteny for genes encoded in M. smithii genomes 273 274 (https://doi.org/10.5281/zenodo.8024791).

275 Among the proteins specific for gut-associated archaea, we identified stage V sporulation proteins AE (spoVAE) and 276 AD (spoVAD) (h9 and h20). Using BLAST searches, we extracted 250 bacterial protein sequences for SpoVAE and 277 SpoVAD from Uniprot, including 12 spoVAE and 38 spoVAD proteins from environmental samples and the rest from 278 isolate bacterial genomes belonging to the Firmicutes phylum. Phylogenetic trees demonstrated that proteins from h9 279 and h20 are phylogenetically and compositionally distinct from other sequences and form separate branches (Supp. 280 Fig. 2-3). Gene synteny analyses revealed that sporulation genes are grouped in operons (K06405, K06406 and 281 K06407; Fig. 3). Moreover, the flanking regions around sporulation genes include genes with key archaeal as well as 282 methanogenic functions. In addition, the flanking regions of both spoVAE and spoVAD genes are also encoded in the 283 M. smithii isolate DSM 861 genome (Fig. 4). However, in contrast to our MAGs, the isolate's genome did not encode 284 the spoVAE and spoVAD genes. To assess whether spoVAE and spoVAD genes were acquired by M. smithii via HGT, 285 we performed synteny analysis of bacterial sequences obtained from our human gut dataset that shared similarities 286 with the archaeal sequences in clusters h9 and h20. This analysis revealed that in the bacterial genomes found in the

human intestine, the flanking regions of *spoVAE* and *spoVAD* genes include genes mediating and facilitating HGT, such as a site-specific DNA recombinase (K06400) encoded upstream from *spoVAE* and type IV pilus assembly proteins (K02662, K02664) encoded downstream from *spoVAD* (Supp. Fig. 4-5). Genes originating from clusters *h9* and *h20* are found within bacterial genomes of Firmicutes phylum members, specifically *Clostridium* sp. CAG-302 and CAG-269, which highlights their association with known bacterial taxa in the gut and indicates horizontal gene transfer between these distantly related taxa.

293 While sporulation has been primarily observed in spore-forming bacteria and not in archaea, it is known that non-294 sporulating bacterial species also encode sporulation genes. In these bacterial taxa, the genes likely encode regulatory 295 proteins involved in peptidoglycan (PPG) turnover, thereby playing a role in cell division and/or development [94], [95]. 296 Archaea lack PPG but methanogenic archaea, including Methanobrevibacter species, use pseudopeptidoglycan 297 (pseudo-PPG) instead, which functions similarly to PPG in a bacterial cell and results in Gram positive staining [96]. 298 Certain structural similarities between methanogens and bacteria described above leave open the question of whether 299 sporulation proteins could play a similar role in pseudopeptidoglycan turnover in methanogenic archaea, analogous to 300 their function in non-sporulating bacteria. The identification of these genes holds significant interest, especially in light 301 of the work by Nelson Sathi et al., suggesting that methanogens frequently acquire functionally active genes through 302 horizontal transfer from bacteria. Comprehensive experimental analysis is required to determine their specific functions, 303 but these findings present an exciting opportunity for further exploration [97]. Phylogenetic analysis of spoVAE and 304 spoVAD has demonstrated that sequences from the abovementioned clusters are compositionally homogeneous but 305 phylogenetically distant from other known similar sequences in Uniprot, and therefore might be unique to the human 306 gut environment. Moreover, archaeal and bacterial sequences from sourmash clusters h9 and h20 branch out together, 307 which suggests that sporulation genes encoded in archaea are the result of horizontal gene transfer from bacteria to 308 archaea. This study provides evidence that archaeal genomes exhibit clustered sporulation genes surrounded by genes 309 linked to archaea-specific functions like pyrimidine, thiamine, and methane metabolism. Moreover, genes in flanking 310 regions up- and downstream of spoVAE and spoVAD genes are indeed encoded in the representative M. smithii isolate 311 DSM 861. As bacteria encoding similar spoVAE and spoVAD proteins and bacterial sequences from clusters h9 and 312 h20 belong to various species of the Clostridium genus, HGT probably occurred in the direction from the 313 abovementioned species to M. smithii. Moreover, Ruaud, Esquivel-Elizondo, de la Cuesta-Zuluaga et al. [98] have 314 provided evidence of a syntrophic relationship between Firmicutes bacteria and Methanobrevibacter smithii. The co-315 occurrence of these microorganisms is likely facilitated by physical and metabolic interactions. In addition to this, genes 316 h9 and h20 as well as their surrounding genes are expressed by the archaeal genomes sampled from human faecal 317 samples.

318 Conclusion

319 Our study aimed to uncover the potential functions of archaeal proteins, particularly those encoded by M. smithii, in the 320 human gut. Sequence similarity-based methods, while effective for highly similar proteins (>70-80% identity), may not 321 accurately represent the functions of archaeal proteins due to the lack of experimental validation. More specifically, 322 publicly available databases have limited experimentally validated archaeal sequences compared to bacterial and 323 eukaryotic proteins (~7 000 000 archaeal, ~166 000 000 bacterial and ~70 000 000 eukaryotic proteins, UniProtKB Jun 324 2023) making sequence-based protein annotations applicable to only a subset of archaeal proteins. In contrast, recent 325 deep learning-based methods enable protein structure prediction and annotation without relying on high sequence 326 similarity, allowing for functional similarity beyond close sequence matches. We used structural methods to improve the 327 annotation of archaeal proteins, gaining better insights into their functions compared to traditional sequence-based 328 methods. This approach allowed us to refine some existing annotations and discover new functions for others, giving

329 us valuable insights into the roles of archaeal genes in the human gut. Our findings focus on the characterization of

human-associated and gut-specific proteins identified in *M. smithii*, a metabolically proficient and clinically relevant

331 methanogenic archaeon known to be linked to gastrointestinal disorders, including IBD and obesity. Future work should 332 help in resolving the predicted structures and protein functions using experimental approaches.

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339 Competing Interests

340 The authors declare that they have no competing interests.

³⁴¹ Data Availability Statement

342 Microbial MAGs from UHGG collection are available from the MGnify FTP site at 343 http://ftp.ebi.ac.uk/pub/databases/metagenomics/mgnify genomes/, MAGs from the GEM catalog are accessible at 344 https://portal.nersc.gov/GEM/. Metatranscriptomic sequencing reads are available from NCBI 345 BioProject PRJNA289586 and assembled contigs can be assessed at MG-RAST [100] (submission IDs are indicated 346 in MT assembly RAST ids.xlsx). A description of the analyses including pre-processing steps along with the scripts 347 for the main analysis, archaeal gut-specific unique and homologous sourmash clusters and synteny plots can be found 348 at GitLab: https://gitlab.lcsb.uni.lu/polina.novikova/archaea-in-gut.

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Figure 1. A, *Flowchart demonstrating major steps of the analysis. The Venn diagram demonstrates the number of shared KOs assigned to archaeal and bacterial smash clusters. B*, *Funnels illustrating the protein count at each stage of protein selection. MM2 – Mmseqs2 clusters, SCs – smash clusters.*



Figure 2. Relative occurrence and average metatranscriptomic read coverage of proteins in the **A**, unique and **B**, homologous groups of clusters with archaeal proteins. *MG* – metagenomics, *MT* – metatranscriptomics.



Figure 3. Gene synteny for sporulation stage V genes AE and AD from their respective smash clusters **A**, h9, and **B**, h20. Gene expression of target genes (spoVAE and spoVAD, in red) as well as genes from flanking regions are demonstrated below each sequence and are colored correspondingly. Genes with key archaeal functions: **A**, pyrimidine metabolism (K18678, phytol kinase), methane metabolism (K11781, 5-amino-6-(D-ribitylamino)uracil–L-tyrosine 4-hydroxyphenyl transferase) and thiamine metabolism (K00878, hydroxyethylthiazole kinase; K00788, thiamine-phosphate pyrophosphorylase); **B**, pyrimidine metabolism (K22026, nucleoside kinase; K18678, phytol kinase) and methane metabolism (K11781, 5-amino)uracil–L-tyrosine 4-hydroxyphenyl transferase).



Figure 4. Genomic context of the archaeal flanking regions up- and downstream of the **A**, spoVAE and **B**, spoVAD gene clusters in the M. smithii strain DSM 861.



Figure 5. Schematic proposal highlighting proteins specific to gut-associated archaea with described functions: **u1** - Type II restriction endonuclease BgIII, **u2** - Intimin/invasin-like protein with a Ig-like domain, **u3** - Intimin/invasinlike protein, **u4** - Unr protein, **u22** - Type I restriction-modification EcoKI enzyme, specificity subunit, **u24** -Polypeptide N-acetylgalactosaminyltransferase, **h1** - 4-amino-4-deoxy-L-arabinose transferase or related glycosyltransferases of PMT family, **h2,3,4,6** - Dolichyl-phosphate-mannose – protein mannosyltransferase 1, **h5** - Dolichyl-diphosphooligosaccharide-protein glycosyltransferase subunit STT3B, **h7** - Propanediol utilisation protein pduA, **h11** - Phosphoenolpyruvate-dependent PTS system, IIA component, **h28** - Transthyretin-like protein, **h31** - 2-aminoethylphosphonate-pyruvate aminotransferase.



Figure 1. Heatmaps demonstrating the intensity of HGT events between M. smithii genomes. **A**, HGT between taxonomic groups named as follows: A - Methanobrevibacter_A smithii, B - Methanobrevibacter_A smithii_A (Ca. Methanobreviabcter intestini), C - Methanobrevibacter_A oralis, E - GCF_000016525.1 (M. smithii), F - GCF_002252585.1 (Ca. Methanobreviabcter intestini); **B**, HGT events between individual genomes of same groups. The legend depicts the frequency of HGT events among the genomes of **A**, taxonomic groups and **B**, individual genomes.



Figure 2. Phylogenetic tree of stage V sporulation proteins AE from identified SC h9 and Uniprot. Bacterial and archaeal proteins from cluster h9 are depicted as GUT_bacteria and GUT_archaea in dark blue and pink, respectively.



Figure 3. Phylogenetic tree of stage V sporulation proteins AD from identified SC h20 and Uniprot. Bacterial and archaeal proteins from cluster h9 are depicted as GUT_bacteria and GUT_archaea in dark blue and pink, respectively.

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Figure 4. Gene synteny of homologous bacterial sequences obtained from the human gut dataset that share similarities with the archaeal sequences from cluster h9 encoding stage V sporulation protein AE (spoVAE).



Figure 5. Gene synteny of homologous bacterial sequences obtained from the human gut dataset that share similarities with the archaeal sequences from cluster h20 encoding stage V sporulation protein AD (spoVAD).