

Open issues for protein function assignment in *Haloferax volcanii* and other halophilic archaea.

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1 Abstract

2

3 **Background:** Annotation ambiguities and annotation errors are a general challenge in
4 genomics. While a reliable protein function assignment can be obtained by
5 experimental characterization, this is expensive and time-consuming, and the number
6 of such Gold Standard Proteins (GSP) with experimental support remains very low
7 compared to proteins annotated by sequence homology, usually through automated
8 pipelines. Even a GSP may give a misleading assignment when used as a reference:
9 the homolog may be close enough to support isofunctionality, but the substrate of the
10 GSP is absent from the species being annotated. In such cases the enzymes cannot be
11 isofunctional. Here, we examine a variety of such issues in halophilic archaea (class
12 Halobacteria), with a strong focus on the model haloarchaeon *Haloferax volcanii*.

13 **Results:** Annotated proteins of *Hfx. volcanii* were identified for which public
14 databases tend to assign a function that is probably incorrect. In some cases, an
15 alternative, probably correct, function can be predicted or inferred from the available
16 evidence but this has not been adopted by public databases because experimental
17 validation is lacking. In other cases, a probably invalid specific function is predicted
18 by homology, and while there is evidence that this assigned function is unlikely, the
19 true function remains elusive. We list 50 of those cases, each with detailed
20 background information so that a conclusion about the most likely biological function
21 can be drawn. For reasons of brevity and comprehension, only key aspects are listed
22 in the main text, with detailed information being provided in a corresponding section
23 of the Supplementary Material.

24 **Conclusions:** Compiling, describing and summarizing these open annotation issues
25 and functional predictions will benefit the scientific community in the general effort
26 to improve the evaluation of protein function assignments and more thoroughly detail
27 them. By highlighting the gaps and likely annotation errors currently in the databases,
28 we hope this study will provide a framework for experimentalists to systematically
29 confirm (or disprove) our function predictions or to uncover yet unexpected functions.

30

31 **Keywords:** haloarchaea, genome annotation, Gold Standard Protein, *Haloferax*
32 *volcanii*, annotation error

33

1 INTRODUCTION

Haloflex volcanii is a model organism for halophilic archaea (Hartman et al., 2010, Schulze et al., 2020, Leigh et al., 2011, Perez-Arnaiz et al., 2020, Soppa, 2011, Haque et al., 2020), for which an elaborate set of genetic tools has been developed (Allers et al., 2010, Allers and Mevarech, 2005, Kiljunen et al., 2014). Its genome has been sequenced and carefully annotated (Hartman et al., 2010, Pfeiffer et al., 2008a, Pfeiffer and Oesterhelt, 2015). A plethora of biological aspects have been successfully tackled in this species, with examples including DNA replication (Perez-Arnaiz et al., 2020); cell division and cell shape (Turkowsky et al., 2020, Walsh et al., 2019, de Silva et al., 2021, Duggin et al., 2015, Liao et al., 2021); metabolism (Brasen and Schonheit, 2001, Johnsen et al., 2009, Pickl et al., 2012, Sutter et al., 2016, Reinhardt et al., 2019, Kuprat et al., 2021, Kuprat et al., 2020, Sutter et al., 2020, Tästensen et al., 2020); protein secretion (Abdul-Halim et al., 2020, Abdul Halim et al., 2018, Abdul Halim et al., 2013, Storf et al., 2010); motility and biofilms (Schiller et al., 2020, Pohlschroder and Esquivel, 2015, Li et al., 2020, Collins et al., 2020, Quax et al., 2018, Nussbaum et al., 2020); mating (Shalev et al., 2017); signalling (Braun et al., 2019); virus defence (Maier et al., 2019); proteolysis (Reuter and Maupin-Furlow, 2004, Reuter et al., 2010, Prunetti et al., 2014, Cerletti et al., 2018, Cerletti et al., 2014, Costa et al., 2018); posttranslational modification (N-glycosylation; SAMPylation)(Cao et al., 2015, Kaminski and Eichler, 2014, Tripepi et al., 2012, Schulze et al., 2021, Shalev et al., 2018, Kandiba et al., 2016); gene regulation (Qi et al., 2016, Rawls et al., 2010, Hattori et al., 2016, Hwang et al., 2017, Johnsen et al., 2015, Tästensen et al., 2020, Reinhardt et al., 2019); microproteins (Zahn et al., 2021, Nagel et al., 2019, Kubatova et al., 2020) and small noncoding RNAs (sRNAs) (Straub et al., 2009, Heyer et al., 2012, Babski et al., 2014, Wyss et al., 2018).

Genome annotations are frequently compromised by annotation errors (Schnoes et al., 2009, Pfeiffer and Oesterhelt, 2015, Promponas et al., 2015, Danchin et al., 2018). Many of these errors are caused by invalid annotation transfer between presumed homologs, which, once introduced, are further spread by annotation robots. This problem can be partially overcome by using a Gold Standard Protein (GSP) based annotation strategy (Pfeiffer and Oesterhelt, 2015). Because the GSP has itself been subjected to experimental analysis, its annotation cannot be caused by an invalid

annotation transfer process. The GSP strategy had already been applied to a detailed analysis of the metabolism of halophilic archaea (Falb et al., 2008). However, with decreasing level of sequence identity, the assumption of isofunctionality becomes increasingly insecure. Although this may be counterbalanced by additional evidence, e.g. gene clustering, experimental confirmation would be the best option for validation of the annotation.

There are additional and much more subtle genome annotation problems. In some cases, GSPs are true homologs and the annotated function in the database is correct. Nevertheless, the biological context in the query organism makes it unlikely that the homologs are isofunctional, e.g. when the substrate of the GSP is lacking in the query organism. Also, paralogs may have distinct but related functions, which cannot be assigned by sequence analysis but may be assigned based on phylogenetic considerations. Here, again, experimental confirmation is the preferred option for validation of the annotation. Lack of experimental confirmation may keep high-level databases like KEGG or the SwissProt section of UniProt from adopting assignments based on well-supported bioinformatic analyses, so that the database entries continue to provide information that is probably incorrect. We refer to annotation problems in these databases solely to underscore that the biological issues raised by us are far from trivial. There is no intention to question the exceedingly high quality of the SwissProt and KEGG databases (UniProt, 2021, Kanehisa et al., 2019) and their tremendous value for the scientific community. We have actively supported them by providing feedback and encourage others to do the same, e.g. with the recently implemented “Add a publication” functionality in UniProt entries, that allows users to connect a protein to a publication that describes its experimental characterization (<https://community.uniprot.org/bbsub/bbsubinfo.html>).

In this study, we describe a number of annotation issues for haloarchaea with a strong emphasis on *Hfx. volcanii*. We denote such cases as ‘open annotation issues’ with the hope of attracting members of the *Haloferax* community and other groups working with halophilic archaea to apply experimental analyses to elucidate the true function(s) of these proteins. This will increase the number of Gold Standard Proteins

which originate from *Hfx. volcanii* or other haloarchaea, reduce genome annotation ambiguities, and perhaps uncover novel metabolic processes.

2 MATERIALS AND METHODS

2.1 Curation of genome annotation and Gold Standard Protein identification

The Gold Standard Protein based curation of haloarchaeal genomes has been described (Pfeiffer and Oesterhelt, 2015). Since then, a systematic comparison to KEGG data was performed for a subset of the curated genomes (Pfeiffer et al., 2020). The *Hfx. volcanii* genome annotation is continuously scrutinized, especially when a closely related genome is annotated (Tittes et al., 2021). The 16 haloarchaeal genomes that are currently under survey are listed in Suppl. Table S11.

2.2 Additional bioinformatics tools

Key databases were UniProtKB/Swiss-Prot (UniProt, 2021), InterPro (Hunter et al., 2009), KEGG (Kanehisa et al., 2019), and OrthoDB (Kriventseva et al., 2019). The SyntTax server was used for inspecting conservation of gene neighbourhood analysis (Oberto, 2013). As general tools the BLAST suite of programs (Johnson et al., 2008, Altschul et al., 1997) was used for genome comparisons.

3 RESULTS

Open issues are organised below under the subsections (3.1), the respiratory chain and oxidative decarboxylation; (3.2), amino acid metabolism; (3.3), heme and cobalamin biosynthesis; (3.4), coenzyme F420; (3.5), tetrahydrofolate as opposed to methanopterin; (3.6), NAD and riboflavin; (3.7), lipid metabolism; (3.8), genetic information processing, and (3.9), stand-alone (miscellaneous) cases.

3.1 The respiratory chain and oxidative decarboxylation

In the respiratory chain, coenzymes that have been reduced during catabolism (e.g. glycolysis) are reoxidized, with the energy being saved as an ion gradient. The textbook example of a respiratory chain are the five mitochondrial complexes (Rich and Marechal, 2010, Guo et al., 2018): complex I (NADH dehydrogenase), complex II (succinate dehydrogenase), complex III (cytochrome bc_1 complex), complex IV (cytochrome-c oxidase as prototype for a terminal oxidase) and complex V (F-type ATP synthase). In mitochondria, a significant part of the NADH which feeds into the respiratory chain originates from oxidative decarboxylation: conversion of pyruvate to acetyl-CoA by the pyruvate dehydrogenase complex and conversion of alpha-ketoglutarate to succinyl-CoA by the homologous 2-oxoglutarate dehydrogenase complex. While complex I and II transfer reducing elements to a lipid-embedded two-electron carrier (ubiquinone), the bc_1 complex transfers the electrons to the one-electron carrier cytochrome-c, a heme (and thus iron) protein, which then transfers electrons to the terminal oxidase.

Bacteria like *Escherichia coli* and *Paracoccus* have related complexes and enzymes: NADH dehydrogenase (encoded by the *nuo* operon), succinate dehydrogenase (encoded by *sdhABCD*) and the related fumarate reductase (encoded by *frdABCD*) (Crofts et al., 2013), several terminal oxidases (e.g. products of *cyoABCDE*, *cydABC*), and an F-type ATP synthase (encoded by *atp* genes). *E. coli* lacks a bc_1 complex, which, however, occurs in *Paracoccus denitrificans* (Kaila and Wikstrom, 2021). *E. coli* contains the canonical complexes of oxidative decarboxylation (pyruvate dehydrogenase complex, encoded by *aceEF+lpdA*, and 2-oxoglutarate dehydrogenase complex, encoded by *sucAB+lpdA*).

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150 The respiratory chain of *Hfx. volcanii* and other haloarchaea deviates considerably
 151 from those of mitochondria and bacteria such as *Paracoccus* and *E. coli* (reviewed by
 152 (Schafer et al., 1999)), and a number of questions remain unresolved. We focus on the
 153 equivalents of complex I, III and IV, because these have unresolved issues. We also
 154 cover some aspects relevant for NADH levels (oxidative decarboxylation enzymes
 155 and type II NADH dehydrogenase). We do not cover complexes that have already
 156 been studied in haloarchaea: complex II (succinate dehydrogenase) (Scharf et al.,
 157 1997, Sreeramulu et al., 1998, Gradin et al., 1985) and complex V (ATP synthase)
 158 (Steinert et al., 1997, Nanba and Mukohata, 1987).

159

160 (a) In haloarchaea, oxidative decarboxylation is not linked to reduction of NAD to
 161 NADH but to reduction of a ferredoxin (encoded by *fdx*, e.g. OE_4217R, HVO_2995)
 162 which has a redox potential similar to that of the NAD/NADH pair (Kerscher and
 163 Oesterhelt, 1977). The enzymes for oxidative decarboxylation are pyruvate--
 164 ferredoxin oxidoreductase (*porAB*, e.g. OE_2623R/2622R, HVO_1305/1304) and 2-
 165 oxoglutarate--ferredoxin oxidoreductase (*korAB*, e.g. OE_1711R/1710R,
 166 HVO_0888/0887), and these have been characterized from *Halobacterium salinarum*
 167 (Plaga et al., 1992, Kerscher and Oesterhelt, 1981b, Kerscher and Oesterhelt, 1981a).

168

169 (b) In *Hfx. volcanii*, ferredoxin Fdx (HVO_2995) plays an essential role in nitrate
 170 assimilation (Zafrilla et al., 2011). It may well be involved in additional metabolic
 171 processes and it is yet unresolved how ferredoxin Fdx is reoxidised, but this might be
 172 achieved by the Nuo complex.

173

174 (c) The *nuo* cluster of haloarchaea resembles that of *E. coli*, with genes and gene
 175 order highly conserved, and just a few domain fissions and fusions. However,
 176 haloarchaea lack NuoEFG (Falb et al., 2005), which is a subcomplex mediating
 177 interaction with NADH (Leif et al., 1995, Braun et al., 1998). Thus, the haloarchaeal
 178 *nuo* complex is unlikely to function as NADH dehydrogenase, despite its annotation
 179 as such in KEGG (as of April 2021).

180

181 (d) Other catabolic enzymes generate NADH, which also must be reoxidized. Based
182 on inhibitor studies in *Hbt. salinarum*, NADH is not reoxidized by a type I but rather
183 by a type II NADH dehydrogenase (Sreeramulu et al., 1998). A gene has been
184 assigned for *Natronomonas pharaonis* (Falb et al., 2008). However, for reasons
185 detailed in Suppl.Text.S1, this assignment is highly questionable, so that this issue
186 calls for experimental analysis.

187

188 (e) About one-third of the haloarchaea, especially the *Natrialbales*, do not code for a
189 complex III equivalent (cytochrome bc₁ complex encoded by *petABC*) according to
190 OrthoDB analysis. The bc₁ complex is required to transfer electrons from the lipid-
191 embedded two-electron carrier (menaquinone in haloarchaea) to the one-electron
192 carrier associated with terminal oxidases (probably halocyanin). How electrons flow
193 in the absence of a complex III equivalent is currently unresolved.

194

195 The haloarchaeal *petABC* genes resemble those of the chloroplast b6-f complex rather
196 than those of the mitochondrial bc₁ complex (see Suppl.Text S1 Section 1 for more
197 details).

198

199 (f) A bc cytochrome has been purified from *Nmn. pharaonis*, but with an atypical 1:1
200 ratio between the b-type and c-type heme (Scharf et al., 1997). The complex is
201 heterodimeric, with subunits of 18 kDa and 14 kDa. The 18 kDa subunit carries the
202 covalently attached heme group (Scharf et al., 1997). An attempt was made to identify
203 the genes coding for these subunits (Mattar, 1996) (for details see Suppl.Text S1
204 Section 1). Two approaches were used to obtain protein sequence data, one being N-
205 terminal protein sequencing of the two subunits extracted from a SDS-polyacrylamide
206 gel. In the other attempt, peptides from the purified complex were separated by
207 HPLC, and a peptide absorbing at 280 nm (protein) as well as 400 nm (heme) was
208 isolated. Absorption at 400 nm clearly indicates covalent attachment of the heme
209 group to the peptide. The sequences from the two approaches overlapped and resulted
210 in a contiguous sequence of 41 aa, with only the penultimate position remaining
211 undefined (Mattar, 1996). Based on this information, a PCR probe was generated

(designated “cyt-C Sonde”) that allowed the gene to be identified and sequenced, including its genomic neighbourhood. It turned out that the genes coding for the four subunits of succinate dehydrogenase (*sdhCDBA*) had been isolated. The obtained protein sequence corresponds to the N-terminal region of *sdhD* (with the initiator methionine cleaved off) and only 2 sequence discrepancies in addition to the unresolved penultimate residue.

In the PhD thesis (Mattar, 1996), this unambiguous result was rated to be a failure (and the data were never formally published). The reason is that SdhD is free of cysteine residues, while textbook knowledge states that a pair of cysteines is required for covalent heme attachment (Kletzin et al., 2015). The lack of the required cysteine pair was taken to indicate that the results were incorrect and that the identified genes did not encode the cytochrome bc that the study had been seeking (Mattar, 1996). In contrast, we speculate that the results were completely correct, despite being in conflict with the cysteine pair paradigm. In our view, a paradigm shift is required. The obtained results call for a yet unanticipated novel mode of covalent heme attachment, exemplified by the 18 kDa subunit of *Natronomonas* succinate dehydrogenase subunit SdhD. It should be noted that the 41 aa protein sequence, which had been obtained, turned out to contain three histidine residues upon translation of the gene, but none of these had been detected upon Edman degradation.

In *Halobacterium*, a small c-type cytochrome was purified (cytochrome _{C552}, 14.1 kDa) (Sreeramulu, 2003). Heme staining after SDS-PAGE indicated a covalent heme attachment, but no sequence or composition data were reported, so that it is not possible to identify the protein based on the available information. We speculate that the *Halobacterium* cytochrome _{C552} also represents SdhD (as detailed in Suppl. Text S1 Section 1). In that case, the proposed novel type of covalent heme attachment would not be restricted to *Nmn. pharaonis* but might be a general property of haloarchaea. This would also solve the “*Halobacterium* paradox” (Kletzin et al., 2015).

(g) The haloarchaeal one-electron carrier is the copper protein halocyanin rather than the iron-containing heme protein cytochrome-c. A halocyanin from *Nmn. pharaonis* (NP_3954A) has been characterized, including its redox potential (Mattar et al., 1994, Scharf and Engelhard, 1993, Hildebrandt et al., 1994). A gene fusion supports the close connection of a halocyanin with a subunit of a terminal oxidase. For further details see Suppl.Text S1 Section 1.

(h) Terminal oxidases are highly diverse in haloarchaea and we restrict our analysis to three species (*Nmn. pharaonis*, *Hfx. volcanii*, and *Hbt. salinarum*) because in each of these at least one terminal oxidase has been experimentally studied (Table 1). Details are described in Suppl. Text S1 Section 1.

(i) NAD-dependent oxidative decarboxylation is a canonical reaction to convert pyruvate into acetyl-CoA, and alpha-ketoglutarate into succinyl-CoA. In haloarchaea, the conversion of pyruvate to acetyl-CoA and alpha-ketoglutarate to succinyl-CoA is dependent on ferredoxin, not on NAD (see above). Nevertheless, most haloarchaeal genomes also code for homologs of enzymes catalyzing NAD-dependent oxidative decarboxylation, such as the *E. coli* pyruvate dehydrogenase complex. In most cases, the substrates could not be identified, an exception being a paralog involved in isoleucine catabolism (Sisignano et al., 2010). In several cases the enzymes were found not to show catalytic activity with pyruvate or alpha-ketoglutarate (see Suppl.Text S1 Section 1 for details). Also, a conditional lethal *porAB* mutant was unable to grow on glucose or pyruvate, thus excluding that alternative enzymes for conversion of pyruvate to acetyl-CoA exist in *Hfx. volcanii* (Kuprat et al., 2021). Nonetheless, despite experimental results to the contrary, pyruvate is assigned as substrate for some of the homologs of the pyruvate dehydrogenase complex in KEGG (as of April 2021).

Section	Code	Gene	Gold Standard Protein				Reference	PMID	Comment
			isofunc	%seq_id	Locus tag	UniProt			
1a	HVO_1305 HVO_1304	<i>porAB</i>	yes	67% 80%	OE2623R OE2622R	B0R4X6 B0R4X5	(Plaga et al., 1992) (Kerscher and Oesterhelt, 1981b) (Kerscher and Oesterhelt, 1981a)	1555599 6266826 6266827	
1a	HVO_0888 HVO_0887	<i>korAB</i>	yes	77% 77%	OE1711R OE1710R	B0R3G0 B0R3F9	(Kerscher and Oesterhelt, 1981b) (Kerscher and Oesterhelt, 1981a)	6266826 6266827	
1a/1b	HVO_2995	<i>fdx</i>	yes	88%	OE4217R	B0R7I9	(Kerscher and Oesterhelt, 1976) (Kerscher et al., 1976) (Kerscher and Oesterhelt, 1977)	964365 188650 201489	role in oxidative decarboxylation
1a/1b	HVO_2995 (cont.)				self	D4GY89	(Zafrilla et al., 2011)	22103537	role in nitrate assimilation
1c	HVO_0979 (complex)	<i>nuoB</i>	possibly	50%	tlr0705	Q8DKZ4	(Zhang et al., 2005) (Schuller et al., 2019) (Pan et al., 2020)	15910282 30573545 32001694	reoxidizes ferredoxin
1c	HVO_0979 (cont.)		no	48%	b2287	P0AFC7	(Leif et al., 1995) (Braun et al., 1998)	7607227 9485311	reoxidizes NADH in <i>E.coli</i>
1d	NP_3508A	<i>ndh1</i>	special	26% (N-term 140 aa)	-	Q7ZAG8			function of Q7ZAG8 was reassigned (from <i>ndh1</i> to <i>sqr</i>) after annotation transfer
1d	NP_3508A (cont.)		possibly	30%	BpOF4_04810	A7LKG4	(Liu et al., 2008)	18359284	type II NADH dehydrogenase
1e	HVO_2620 HVO_0842 HVO_0841	<i>petABD</i>	yes	39%	SYNPCC7002_A0842	P28056	(Lee et al., 2001)	11245788	HVO_0842 (<i>petB</i>) related to cytochrome b6
1f	HVO_2810	<i>sdhD</i>	yes	66%	NP_4268A	Q3INS7	(Scharf et al., 1997) (Mattar, 1996)	9109654 PhD_Mattar	
1g	HVO_0943	<i>cbaD</i>	yes	57%	NP_2966A	A0A1U7EWW4	(Mattar and Engelhard, 1997)	9428682	
	HVO_0943 (cont.)		-	63%	OE_4073R (C-term)	B0R7A9		-	halocyanin/cbaD fusion protein, uncharacterized

1g	HVO_2150	<i>hcpG</i>	-	44%	OE_4073R (N-term)	B0R7A9		-	halocyanin/cbaD fusion protein, uncharacterized
1h	HVO_0945 (complex)	<i>cbaA</i>	yes	64%	NP_2966A	A0A1U7EWW4	(Mattar and Engelhard, 1997)	9428682	
1h	HVO_0907 (complex)	<i>coxA1</i>			self		(Tanaka et al., 2002)	11790755	
1h	HVO_0907 (cont.)		yes	70%	VNG_0657G (OE_1979R)	P33588	(Fujiwara et al., 1989) (Denda et al., 1991)	2542239 1659810	
1h	HVO_1645 (complex)	<i>coxAC2</i>	yes	43%	APE_0793.1	Q9YdX6	(Ishikawa et al., 2002)	12471503	
1h	HVO_0462 HVO_0461	<i>cydAB</i>	yes	32% 24%	- -	Q09049 Q05780	(Moshiri et al., 1991)	1655703	
1h	HVO_0462 HVO_0461 (cont.)		yes	30% 27%	b0733 b0734	P0ABJ9 P0ABK2	(Miller and Gennis, 1983)	6307994	
1h	NP_4296A NP_4294A	<i>coxA3</i> <i>coxB3</i>	yes	28% 33%	TTHA1135 TTHA1134	Q5SJ79 Q5SJ80	(Zimmermann et al., 1988) (Keightley et al., 1995)	2842747 7657607	
1i	HVO_2958 HVO_2959	<i>oadhAB1</i>			self	D4GY15 D4GY17	(Sisignano et al., 2010)	19910413	Ile indirectly assigned as substrate
1i	HVO_2958 HVO_2959 (cont.)				self		(Jolley et al., 2000) (Al-Mailem et al., 2008) (van Ooyen and Soppa, 2007)	10832633 17571210 17906130	no substrate was identified;pyruvate and alphaKG excluded
1i	HVO_2595 HVO_2596	<i>oadhAB2</i>			self		(Wanner and Soppa, 2002) (van Ooyen and Soppa, 2007) (Sisignano et al., 2010)	12003954 17906130 19910413	no substrate was identified;pyruvate and alphaKG excluded
1i	HVO_0669 HVO_0668	<i>oadhAB3</i>			self		(van Ooyen and Soppa, 2007) (Sisignano et al., 2010)	17906130 19910413	no substrate was identified;pyruvate and alphaKG excluded
1i	HVO_2209	<i>oadhA4</i>			self				not yet analyzed experimentally
1i	HVO_2958 HVO_2959 (cont.)		yes/no	38% 52%	TA1438 TA1437	Q9HIA3 Q9HIA4	(Heath et al., 2007)	17894823	substrates are Ile, Leu, Val

1i	HVO_2595 HVO_2596 (cont.)		no	41% 41%	- -	Q57102 Q57041	(Oppermann et al., 1991)	1898934	substrate is acetoin
1i	HVO_2595 HVO_2596 (cont.)		unknown	40% 43%	BSU08060 BSU08070	O31404 O34591	(Huang et al., 1999)	10368162	substrate is acetoin
1i	HVO_0669 HVO_0668 (cont.)		unknown	54% 47%	BSU08060 BSU08070	O31404 O34591	(Huang et al., 1999)	10368162	substrate is acetoin
1i	HVO_0669 HVO_0668 (cont.)		unknown	49% 43%	- -	Q57102 Q57041	(Oppermann et al., 1991)	1898934	substrate is acetoin
1i	HVO_2209 (cont.)		unknown	38%	TA1438	Q9HIA3	(Heath et al., 2007)	17894823	substrates are Ile, Leu, Val

Table 1: **Proteins with open annotation issues and their Gold Standard Protein homologs (Section 1).** The column Section refers to the Table listing the protein and to the section in the Results and in Suppl. Text S1. As an example, 2c covers topic (c) from the decimal-numbered Results subsection 3.2 Amino Acid Biosynthesis. In Suppl. Text S1, this is covered under Section S2 subsection S2.c. The corresponding proteins are listed in Table 2. For a few proteins, two sections are indicated (e.g. 1a/1b). The column Code refers to a haloarchaeal protein by its locus tag, which is mainly from *Haloferax volcanii* (HVO), but also from *Halobacterium salinarum* (OE), *Natronomonas pharaonis* (NP) and *Halohasta litchfieldiae* (halTADL). When the reconstruction of a complete pathway is presented, the unassigned genes are indicated as a “pathway gap”. In one case we indicate the absence of a haloarchaeal ortholog by a dash. In the case of a complex, we either list more than one code, or we list only one subunit together with the term (complex). All subunits of these complexes are listed groupwise in Table S10. A protein may be shown in more than one row. From the 2nd row onwards, this is indicated by the term (cont.). The column Gene lists the assigned gene or a dash if no gene has been assigned. The assigned gene is only indicated in the first row of a protein. A set of four columns is used to relate a query protein to an experimentally characterized homolog, a GSP (Gold Standard Protein) (isofunc, %seq_id, Locus tag, UniProt). The column

283 isofunc indicates if the query protein and its Gold Standard Protein homolog are isofunctional. The meaning of the terms used in this column in
284 Tables 1-9 (yes, no, yes/no, probably, possibly, unclear, unknown, prediction, special, “-“) is described at the end of this legend. The column
285 %seq_id indicates the protein sequence identity between the query protein and the homologous GSP. The column Locus tag contains the locus
286 tag, if assigned. The column UniProt contains the UniProt accession of the GSP. GSPs have been experimentally characterized as described in a
287 publication. The column Reference links to the reference list of the manuscript. The column PMID lists the PubMed ID of the publication, if
288 available. Otherwise, this is indicated as “not in PubMed”. Also, one PhD thesis is indicated (PhD_Matter). The column Comment provides
289 various types of additional information. The terms used in the column isofunc in Tables 1-9 have the following meaning: The term “yes”
290 indicates that we consider the two proteins as isofunctional and annotate the query protein accordingly. The term “no” is used when we conclude
291 that the proteins differ in function. Additional terms are used for more difficult cases. The term “yes/no” is used for GSPs which are
292 multifunctional, and we assign only one a subset of these functions to the query protein. The term “probably” is used when we consider
293 isofunctionality likely and annotated the query protein accordingly (with the term probable added to the protein name). The term “possibly” is
294 used when we see a good chance that the proteins are isofunctional, but consider it too speculative to annotate the protein accordingly. The term
295 “unclear” is used when we consider it likely that the same overall reaction is catalyzed, but when reaction details, e.g. the energy-providing
296 compound, is unresolved. The term “unknown” is used when it is not possible to predict the substrate of the query protein. The term “prediction”
297 is used if a function assignment is based on bioinformatic analyses but not yet on an experimentally characterized homologous protein. The term
298 “special” is used when multiple arguments have to be considered with full details provided in the corresponding section of Suppl. Text S1.
299 Finally, a dash (“-“) is used when isofunctionality does not apply, e.g. when a homologous Gold Standard Protein could not be identified.

3.2 Amino acid metabolism

While most amino acid biosynthesis and degradation pathways can be reliably reconstructed, a few open issues remain, which are discussed below.

(a) The first and last steps of arginine biosynthesis deal with blocking and unblocking of the alpha-amino group of the substrate (glutamate) and a product intermediate (ornithine). As detailed in Suppl. Text S1 Section 2, it is highly likely that glutamate is attached to the gamma-carboxyl group of a carrier protein, and ornithine is released from that carrier protein. This is based on characterized proteins from *Thermus thermophilus* (Horie et al., 2009), *Thermococcus kodakarensis* (Yoshida et al., 2016) and *Sulfolobus acidocaldarius* (Ouchi et al., 2013). The assignment is strongly supported by clustering of the arginine biosynthesis genes. Some of the homologs are bifunctional, being involved in arginine biosynthesis but also in lysine biosynthesis via the prokaryotic variant of the alpha-aminoadipate pathway. This ambiguity is not assumed to occur in haloarchaea, which use the diaminopimelate pathway for Lys biosynthesis (Hochuli et al., 1999) (see Suppl. Text S1 Section 2 for further discussion of this issue).

Expanding the above, we provide full details underlying our reconstruction of arginine and lysine biosynthesis in *Hfx. volcanii* in Table 2.

Section	Code	Gene	Gold Standard Protein				Reference	PMID	Comment
			isofunc	%seq_id	Locus tag	UniProt			
2a	HVO_0047	<i>argW</i>	no	54%	TT_C1544	Q72HE5	(Yoshida et al., 2015)	25392000	for Arg, not for Lys biosynthesis
2a	HVO_0047 (cont.)		yes/no	39%	Saci_0753	Q4J AQ0			only for Arg, not for Lys biosynthesis
2a	HVO_0047 (cont.)		yes/no	61%	TK0279	Q5JFV9	(Yoshida et al., 2016)	27566549	only for Arg, not for Lys biosynthesis
2a	HVO_0046	<i>argX</i>	no	44%	TT_C1543	Q72HE6	(Horie et al., 2009)	19620981	for Arg, not for Lys biosynthesis
2a	HVO_0046 (cont.)		yes	30%	Saci_1621	Q4J8E7			only for Arg, not for Lys biosynthesis
2a	HVO_0046 (cont.)		yes/no	37%	TK0278	Q5JFW0	(Yoshida et al., 2016)	27566549	only for Arg, not for Lys biosynthesis
2a	HVO_0044	<i>argB</i>	no	41%	TT_C1541	O50147	(Horie et al., 2009) (Yoshida et al., 2015)	19620981 25392000	for Arg, not for Lys biosynthesis
2a	HVO_0044 (cont.)		yes/no	33%	Saci_0751	Q4J AQ2	(Ouchi et al., 2013)	23434852	only for Arg, not for Lys biosynthesis
2a	HVO_0044 (cont.)		yes/no	32%	TK0276	Q5JFW2	(Yoshida et al., 2016)	27566549	only for Arg, not for Lys biosynthesis
2a	HVO_0045	<i>argC</i>	no	48%	TT_C1542	O50146	(Horie et al., 2009) (Shimizu et al., 2016)	19620981 26966182	for Arg, not for Lys biosynthesis
2a	HVO_0045 (cont.)		yes/no	42%	Saci_0750	Q4J AQ3	(Ouchi et al., 2013)	23434852	only for Arg, not for Lys biosynthesis
2a	HVO_0045 (cont.)		yes/no	46%	TK0277	Q5JFW1	(Yoshida et al., 2016)	27566549	only for Arg, not for Lys biosynthesis
2a	HVO_0043	<i>argD</i>	no	45%	TT_C1393	Q93R93	(Miyazaki et al., 2001)	11489859	for Arg, not for Lys biosynthesis
2a	HVO_0043 (cont.)		yes/no	40%	Saci_0755	Q4J AP8	(Ouchi et al., 2013)	23434852	only for Arg, not for Lys biosynthesis
2a	HVO_0043 (cont.)		yes/no	42%	TK0275	Q5JFW3	(Yoshida et al., 2016)	27566549	only for Arg, not for Lys biosynthesis

2a	HVO_0042	<i>argE</i>	no	36%	TT_C1396	Q8VUS5	(Horie et al., 2009) (Fujita et al., 2017)	19620981 28720495	for Arg, not for Lys biosynthesis
2a	HVO_0042 (cont.)		yes/no	29%	Saci_0756	Q4JAP7	(Ouchi et al., 2013)	23434852	only for Arg, not for Lys biosynthesis
2a	HVO_0042 (cont.)		yes/no	37%	TK0274	Q5JFW4	(Yoshida et al., 2016)	27566549	only for Arg, not for Lys biosynthesis
2a	HVO_0041	<i>argF</i>	yes	50%	P18186	BSU11250	(Issaly and Issaly, 1974)	4216455	
2a	HVO_0041 (cont.)		yes	47%	OE_5205R	B0R9X3	(Ruepp et al., 1995)	7868583	
2a	HVO_0049	<i>argG</i>	yes	35%	-	P00966	(Shaheen et al., 1994)	8792870	human
2a	HVO_0049 (cont.)		yes	23%	b3172	P0A6E4	(Lemke et al., 1999)	10666579	<i>E. coli</i>
2a	HVO_0048	<i>argH</i>	yes	38%	MMP0013	O74026	(Cohen-Kupiec et al., 1999)	10220900	
2a	HVO_0008	<i>lysC</i>	yes	32%	BSU28470	P08495	(Kato et al., 2004)	15033471	
2a	HVO_2487	<i>asd</i>	yes	51%	MJ0205	Q57658	(Faehnle et al., 2005)	16225889	
2a/9e	HVO_1101	<i>dapA</i>	yes	45%	PA1010	Q9I4W3	(Kaur et al., 2011)	21396954	
2a	HVO_1100	<i>dapB</i>	yes	33%	b0031	P04036	(Reddy et al., 1995)	7893644	
2a	HVO_1099	<i>dapD</i>	yes	32%	b0166	P0A9D8	(Simms et al., 1984)	6365916	
2a	HVO_1096	<i>dapE</i>	yes	29%	b2472	P0AED7	(Lin et al., 1988)	3276674	function supported by gene clustering
2a	HVO_1097	<i>dapF</i>	yes	35%	b3809	P0A6K1	(Wiseman and Nichols, 1984)	6378903	
2a	HVO_1098	<i>lysA</i>	yes	38%	b2838	P00861	(White and Kelly, 1965)	14343156	
2a	HVO_A0634	-	unknown	25%	b2472	P0AED7	(Lin et al., 1988)	3276674	function assigned to HVO_1096 in <i>dap</i> cluster
2b	HVO_0790	<i>fba2</i>	special	67%	OE_1472F	B0R334	(Gulko et al., 2014)	25216252	EC 2.2.1.10 activity of OE_1472F not yet confirmed in vitro
2b	HVO_0790 (cont.)		special	45%	MJ0400	Q57843	(White, 2004)	15182204	substrate uncertain
2b	HVO_0792	<i>aroB</i>	yes	69%	OE_1475F	B0R336	(Gulko et al., 2014)	25216252	OE_1475F only partially characterized
2b	HVO_0792 (cont.)		yes	44%	MJ1249	Q58646	(White, 2004)	15182204	
2b	HVO_0602	<i>aroD1</i>	yes	44%	OE_1477R	B0R338	(Gulko et al., 2014)	25216252	

2b	HVO_0602 (cont.)		yes	31%	MMP1394	Q6LXF7	(Porat et al., 2004)	15262931	
2c	HVO_0009	<i>tnaA</i>	yes	41%	b3708	P0A853	(Phillips and Gollnick, 1989) (Newton et al., 1965)	2659590 14284727	
2d	HVO_A0559	<i>hutH</i>	yes	42%	BSU39350	P10944	(Oda et al., 1988) (Hartwell and Magasanik, 1963)	2454913 14066617	
2d	HVO_A0562	<i>hutU</i>	yes	62%	BSU39360	P25503	(Kaminskas et al., 1970)	4990470	
2d	HVO_A0560	<i>hutI</i>	yes	42%	BSU39370	P42084	(Yu et al., 2006)	16990261	
2d	HVO_A0561	<i>hutG</i>	yes	33%	BSU39380	P42068	(Kaminskas et al., 1970)	4990470	
2e	HVO_0431	-	-						no GSP available
2e	HVO_0644	<i>leuA1</i>	yes/no	47%	MJ1392	Q58787	(Howell et al., 1999)	9864346	HVO_0644 monofunc (CimA) or bifunc (CimA+LeuA); MJ1392 CimA
2e	HVO_0644 (cont.)		unclear	44%	MJ1195	Q58595	(Howell et al., 1998)	9665716	HVO_0644 monofunc (CimA) or bifunc (CimA+LeuA); MJ1195 LeuA
2e/2f	HVO_1510	<i>leuA2</i>	yes	47%	MJ1195	Q58595	(Howell et al., 1998)	9665716	HVO_1510 LeuA; MJ1195 LeuA
2e/2f	HVO_1510 (cont.)		no	41%	MJ1392	Q58787	(Howell et al., 1999)	9864346	HVO_1510 LeuA MJ1392 CimA
2e	HVO_A0489	-	no	31%	MJ1392	Q58787	(Howell et al., 1999)	9864346	HVO_A0489 general function only; MJ1392 CimA
2e	HVO_A0489 (cont.)		no	30%	MJ1195	Q58595	(Howell et al., 1998)	9665716	HVO_A0489 general function only; MJ1195 LeuA
2e	HVO_1153	-	-						function unassigned; no GSP

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322 Table 2: **Proteins with open annotation issues and their Gold Standard Protein homologs (Section 2).** For a description of this table see the
323 legend to Table 1.

(b) Archaea use a different precursor for aromatic amino acid biosynthesis than the classical pathway. This has been resolved for *Methanocaldococcus jannaschii* and for *Methanococcus maripaludis* (White, 2004, Porat et al., 2006). However, the initial steps may differ from those reported for *Methanocaldococcus* in that fructose-1,6-disphosphate rather than 6-deoxy-5-ketofructose might be a substrate (Gulko et al., 2014). Up to now, a clean deletion of the corresponding enzymes and confirmation with in vitro assays has not yet been achieved (for details see Suppl. Text S1 Section 2).

(c) The gene for tryptophanase (*tpa*) is stringently regulated in *Haloferax*, which is the basis for using its promoter in the toolbox for regulated gene expression (Large et al., 2007). The shutdown of this gene avoids tryptophan degradation when supplies are scarce. Tryptophanase cleaves tryptophan into indole, pyruvate and ammonia. The fate of indole is, however, yet unresolved.

(d) A probable histidine utilization cluster exists, based on characterized homologs from *Bacillus subtilis*, but has not yet been experimentally verified.

(e) Among 16 auxotrophic mutants observed in a *Hfx. volcanii* transposon insertion library (Kiljunen et al., 2014), some could grow only in the presence of one (or several) supplied amino acids. In many cases, the affected genes were known to be involved in the corresponding pathway, but the others may lead to novel function assignments. One affected gene resulted in histidine auxotrophy and the product of this gene (HVO_0431) is an interesting candidate. The InterPro domain assignment (HAD family hydrolase) fits to the only remaining pathway gap in histidine biosynthesis (histidinol-phosphatase). In this context it should be noted that the enzyme which catalyzes the preceding reaction (encoded by *hisC*) is part of a highly conserved three-gene operon involved in polar lipid biosynthesis (see below). For details see Suppl. Text S1 Section 2. One affected gene resulted in isoleucine auxotrophy. The product of this gene (HVO_0644) is currently annotated to catalyze two reactions, one being an early step in isoleucine biosynthesis (EC 2.3.1.182), the

other being the first step after leucine biosynthesis branches off from valine biosynthesis (EC 2.3.3.13) (see below, f) (for details see Suppl. Text S1 Section 2).

(f) *Hfx. volcanii* codes for two paralogs with an attributed function as 2-isopropylmalate synthase (EC 2.3.3.13). This is the first reaction specific to leucine biosynthesis, when the pathway branches off valine biosynthesis. One paralog, HVO_0644, is annotated as bifunctional, also catalyzing a chemically similar reaction which is an early step in isoleucine biosynthesis (EC 2.3.1.182). When the gene encoding HVO_0644 is disrupted by transposon integration, cells cannot grow in the absence of isoleucine. It is unclear if the protein is really bifunctional and is really involved in leucine biosynthesis, catalyzing the reaction of EC 2.3.3.13. The other paralog, HVO_1510, belongs to an ortholog set with major problems concerning start codon assignment. The ortholog set from the 16 genomes listed in Suppl. Table S11 were analyzed. When only canonical start codons are considered (ATG, GTG, TTG), then the orthologs from *Haloferax mediterranei*, *Nmn. pharaonis*, *Natronomonas moolapensis* and *Halohasta litchfieldiae* either lack a long highly conserved N-terminal region, or they are disrupted (pseudogenes), being devoid of a potential start codon. The gene from *Hfx. volcanii* has a start codon (GTG) which is consistent to that of *Haloferax gibbonsii* strain LR2-5 (but a GTA in *Hfx. gibbonsii* strain ARA6). In this region, the gene from *Hfx. mediterranei* is closely related but has in-frame stop codons. HVO_1510 is considerably longer than the orthologs from *Haloquadratum walsbyi*, *Haloarcula hispanica*, and *Natrialba magadii*. The first alternative start codon for HVO_1510 codes for Met-93. This protein was proteomically identified in three ArcPP datasets (Schulze et al., 2020), and peptides upstream of Met-93 were identified. This gene might be translated from an atypical start codon, either an in-frame CTG, or an out-of-frame ATG, which would require ribosomal slippage (for details see Suppl. Text S1 Section 2). It is tempting to speculate that translation occurs only when leucine is not available.

3.3 Coenzymes I: cobalamin and heme

The classical heme biosynthesis pathway branches off cobalamin biosynthesis at the level of uroporphyrinogen III. The alternative heme biosynthesis pathway (Bali et al., 2011), which is used by haloarchaea, has an additional common step, the conversion

of uroporphyrinogen III to precorrin-2. For heme biosynthesis, precorrin-2 is converted to siroheme. This pathway has been reconstructed (Siddaramappa et al., 2012), except for the iron insertion step. For de novo cobalamin biosynthesis, haloarchaea use the cobalt-early pathway with a cobalt-dependent key reaction being catalyzed by CbiG (Moore et al., 2013). Several aspects of heme and cobalamin biosynthesis in haloarchaea are yet unresolved. This is illustrated in Figure 1.

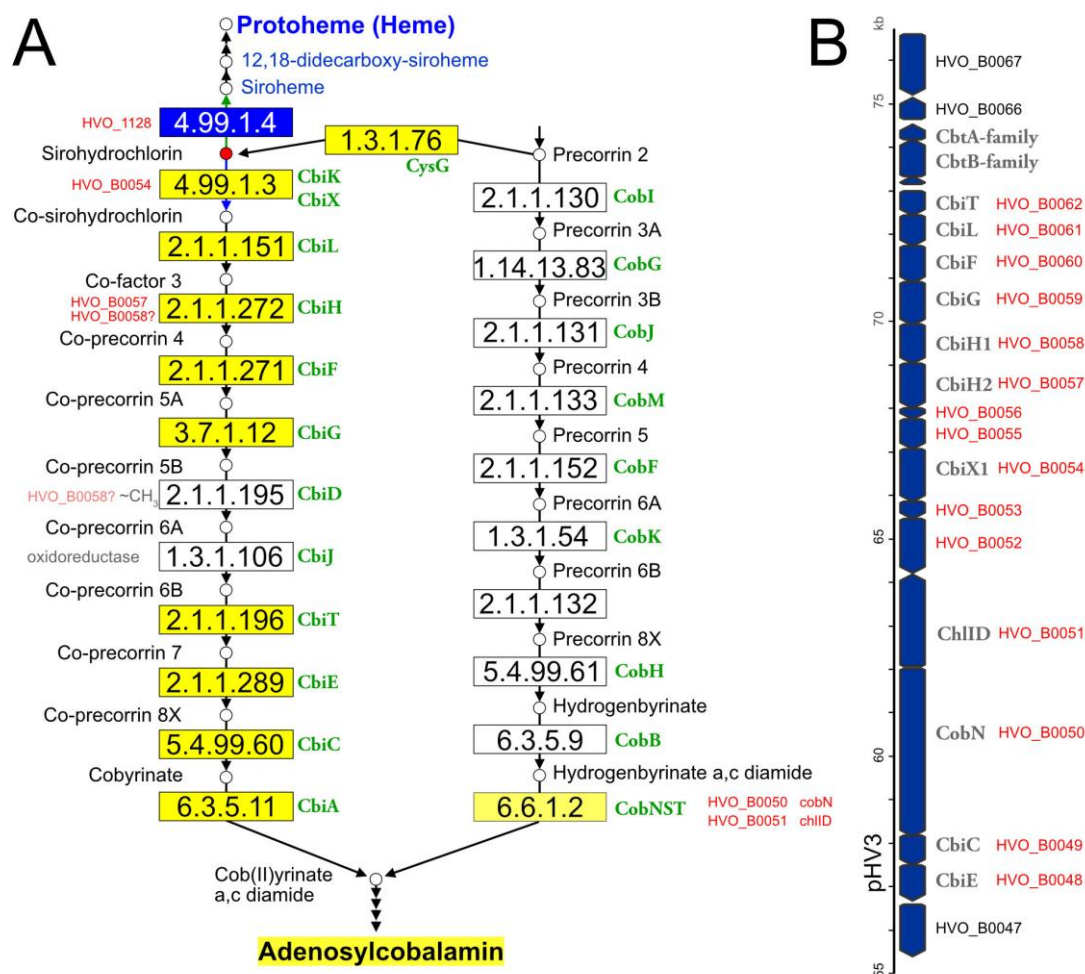


Figure 1. Illustration of the haloarchaeal cobalamin and heme biosynthesis pathways and of the major cobalamin biosynthesis gene cluster. (A) Biosynthesis pathways. This illustration is based on the corresponding KEGG map 00860. Small circles represent pathway intermediates and have their names assigned. Pathway intermediates upstream of Precorrin 2 are not displayed. The circle for sirohydrochlorin is highlighted in red as this is the branchpoint for heme and cobalamin biosynthesis in haloarchaea. Enzymatic reactions are shown by arrows, EC numbers being provided in rectangular boxes. Rectangles are colored when the enzyme has been reconstructed for haloarchaea (blue: heme biosynthesis; dark

yellow: de novo cobalamin biosynthesis; light yellow: late cobaltochelatase which may be a salvage reaction). Gene names in green are adopted from KEGG and represent those from bacterial model pathways. Consecutive arrowheads indicate reaction series which are not shown in detail for space reasons. For enzymatic reactions which are considered to be open issues, the *Hfx. volcanii* locus tags are provided. For two pathway gaps (white boxes in the cobalt-early pathway), the type of reaction is indicated (oxidoreductase and ~CH₃, indicating a methylation reaction). The question mark after HVO_B0058 indicates that this protein, currently co-attributed to EC 2.1.1.272, is a candidate for the yet unassigned EC 2.1.1.195 reaction. We note that haloarchaea might use a deviating biosynthesis pathway, e.g. by swapping the methylation and oxidoreductase reactions (not illustrated). (B) The major cobalamin cluster, encoded on megaplasmid pHV3. Arrows are used to indicate the coding strand and are roughly drawn to scale. If assigned, the gene name is provided in addition to the *Hfx. volcanii* locus tag. Locus tags in red indicate genes that are part of the cobalamin cluster.

(a) *Hfx. volcanii* contains two annotated *cblX* genes. For reasons detailed in Suppl.Text S1 Section 3, we predict that one is a cobaltochelatase, involved in cobalamin biosynthesis, while the other is a ferrochelatase, responsible for conversion of precorrin-2 to siroheme in the alternative heme biosynthesis pathway.

(b) De novo cobalamin biosynthesis has been extensively reconstructed upon curation of the genome annotation (Pfeiffer and Oesterhelt, 2015). All enzymes of the pathway and their associated GSPs are listed in Table 3. Only two pathway gaps remained, and because these are consecutive, it may be possible that the haloarchaeal pathway is non-canonical and proceeds via a novel biosynthetic intermediate. There are only four genes with yet unassigned function in the *Hfx. volcanii* cobalamin gene cluster, and their synteny is well conserved in the majority of haloarchaeal genomes. Thus, these genes are obvious candidates for filling the pathway gaps (for details see Suppl.Text S1 Section 3).

(c) The cobalamin biosynthesis and salvage reactions (those beyond ligand cobyrrinate a,c diamide) involve “adenosylation of the corrin ring, attachment of the aminopropanol arm, and assembly of the nucleotide loop that bridges the lower ligand

dimethylbenzimidazole and the corrin ring” (Rodionov et al., 2003). The enzymes of these branches of cobalamin biosynthesis and their associated GSPs are listed in Table 3. Only two pathway gaps remain open. For one of these, a candidate was proposed upon detailed bioinformatic analysis (Rodionov et al., 2003) (for further details see Suppl.Text S1 Section 3).

(d) Haloarchaea may code for a late cobaltochelatase of the heterotrimeric type. Distantly related GSPs are either cobalt or magnesium chelataes. A late cobaltochelatase is not required for de novo cobalamin biosynthesis via the cobalt-early pathway. We speculate that it may be involved in cobalamin salvage. The chelatase has a mosaic subunit structure as also reported previously (Rodionov et al., 2003) (see Suppl.Text S1 Section 3 for details).

(e) In the alternative heme biosynthesis pathway, siroheme is decarboxylated to 12,18-didecarboxysiroheme, which is attributed to the proteins encoded by *ahbA* and *ahbB*. These are homologous to each other and are organized as two two-domain proteins. It is unclear if AhbA and AhbB function independently or if they form a complex.

Section	Code	Gene	Gold Standard Protein				Reference	PMID	Comment
			isofunc	%seq_id	Locus tag	UniProt			
3a	HVO_B0054	<i>cbiX1</i>	yes	30%	-	O87690	(Raux et al., 2003)	12408752	cobaltochelatase
3a	HVO_B0054 (cont.)		yes	27%	MTH_1397	O27448	(Brindley et al., 2003)	12686546	cobaltochelatase
3a	HVO_1128	<i>cbiX2</i>	no	29%	AF0721	O29537	(Yin et al., 2006)	16835730	cobaltochelatase
3a	HVO_1128 (cont.)		no	28%	MTH_1397	O27448	(Brindley et al., 2003)	12686546	cobaltochelatase
3a	HVO_1128 (cont.)		no	29%	AF0721	O29537	(Yin et al., 2006)	16835730	cobaltochelatase
3a	NP_0734A	<i>cbiX3</i>	-						function unassigned; no GSP; distantly related to paralogs
3a	HVO_2312	<i>sirC</i>	yes/no	31%	Mbar_A1461	Q46CH4	(Storbeck et al., 2010)	21197080	precorrin-2 DH; no analysis for Fe-chelatase
3a	HVO_2312 (cont.)		yes/no	29%	STM3477	P25924	(Stroupe et al., 2003) (Pennington et al., 2020)	14595395 32054833	matches to the N-term domain which is bifunctional as precorrin-2 DH and Fe-chelatase
3a	HVO_2312 (cont.)		yes/no	29%	-	P61818	(Raux et al., 2003) (Schubert et al., 2008)	12408752 18588505	precorrin-2 DH; devoid of Fe-chelatase activity
3b	HVO_B0061	<i>cbiL</i>	no	32%	STM2024	Q05593	(Roessner et al., 1992)	1451790	equivalent reaction on cobalt-free substrate
3b	HVO_B0057	<i>cbiH2</i>	yes	45%	-	O87689	(Moore et al., 2013)	23922391	corresponds to N-term of O87689 which has a C-term extension
3b	HVO_B0057 (cont.)		no	40%	STM2027	Q05590	(Santander et al., 1997) (Santander et al., 2006)	9331403 16198574	equivalent reaction on cobalt-free substrate
3b	HVO_B0058	<i>cbiH1</i>	special	32%	-	O87689	(Moore et al., 2013)	23922391	corresponds to N-term of O87689 which has a C-term extension;

									more distant to O87689 than CbiH2
3b	HVO_B0058 (cont.)		no	30%	STM2027	Q05590	(Santander et al., 1997) (Santander et al., 2006)	9331403 16198574	equivalent reaction on cobalt-free substrate
3b	HVO_B0060	<i>cbiF</i>	no	40%	STM2029	P0A2G9	(Roessner et al., 1992) (Kajiwara et al., 2006)	1451790 16866557	equivalent reaction on cobalt-free substrate
3b	HVO_B0060 (cont.)		yes	38%	-	O87686	(Moore et al., 2013)	23922391	
3b	HVO_B0059	<i>cbiG</i>	yes	24%	-	O87687	(Moore et al., 2013)	23922391	
3b	pathway gap								EC 2.1.1.195
3b	pathway gap								EC 1.3.1.106
3b	HVO_B0062	<i>cbiT</i>	yes	36%	-	O87694	(Moore et al., 2013)	23922391	corresponds to the C-term of bifunctional O87694
3b	HVO_B0048	<i>cbiE</i>	yes	28%	-	O87694	(Moore et al., 2013)	23922391	corresponds to the N-term of bifunctional O87694
3b	HVO_B0049	<i>cbiC</i>	yes	33%	-	O87692	(Moore et al., 2013)	23922391	
3b	HVO_A0487	<i>cbiA</i>	no	37%	STM2035	P29946	(Fresquet et al., 2004)	15311923	equivalent reaction on cobalt-free substrate
3b	HVO_B0052	-	-						function unassigned; no GSP
3b	HVO_B0053	-	-						function unassigned; no GSP
3b	HVO_B0055	-	-						function unassigned; no GSP
3b	HVO_B0056	-	-						function unassigned; no GSP
3c	HVO_A0488	<i>cobA</i>	yes	31%	MM_3138	Q8PSE1	(Buan et al., 2006)	16672609	
3c	HVO_A0488 (cont.)		yes	30%	STM1718	P31570	(Fonseca et al., 2002)	12080060	
3c	HVO_2395	<i>pduO</i>	yes	37%	-	Q9XDN2	(Johnson et al., 2001)	11160088	PduO and CobA are isofunctional; In Q9XDN2, the PduO domain (N-term) is

									fused to a DUF336 domain
3c	HVO_A0553	<i>cbiP</i>	yes	63%	VNG_1576G OE_3246F	Q9HPL5 B0R5X2	(Woodson et al., 2003b)	14645280	
3c	HVO_0587	<i>cbiB</i>	yes	58%	VNG_1578H OE_3253F	Q9HPL3 B0R5X4	(Woodson et al., 2003b)	14645280	
3c	HVO_0592	<i>cbiZ</i>	yes	57%	VNG_1583C OE_3261F	Q9HPL3 B0R5X8	(Woodson and Escalante-Semerena, 2004)	14990804	
3c	HVO_0589	<i>cobY</i>	yes	47%	VNG_1581C OE_3257F	Q9HPL1 B0R5X6	(Woodson et al., 2003a)	12486068	
3c	HVO_0588	<i>cobS</i>	yes	30%	STM2017	Q05602	(Zayas and Escalante-Semerena, 2007)	17209023	
3c	-				STM0643	P39701	(O'Toole et al., 1994)	7929373	EC 3.1.3.73; CobC; no homolog in haloarchaea
3c	HVO_0586	-	prediction	-	-	-	(Rodionov et al., 2003)	12869542	EC 3.1.3.73; prediction for HSL01294 (VNG_1577C)
3c	pathway gap								EC 2.7.1.177
3c	HVO_0591	<i>cobD1</i>	yes	31%	STM0644	P97084	(Brushaber et al., 1998)	9446573	
3c	HVO_0593	<i>cobD2</i>	yes						no GSP; 51% seq_id to HVO_0591 (<i>cobD1</i>)
3c	HVO_0590	<i>cobT</i>	prediction				(Rodionov et al., 2003)	12869542	prediction for VNG_1572C
3c	halTADL_3045	<i>cobT</i>	yes	39%	STM0644	Q05603	(Trzebiatowski et al., 1994)	8206834	
3d	HVO_B0051	<i>cobN</i>	yes	34%	-	P29929	(Debussche et al., 1992)	1429466	
3d	HVO_B0051 (cont.)		no	29%	-	Q55284	(Jensen et al., 1996) (Jensen et al., 1998)	8663186 9716491	Mg chelatase
3d	HVO_B0050	<i>chlID</i>	no	46%	slr1030	P51634	(Jensen et al., 1996) (Jensen et al., 1998)	8663186 9716491	match to N-term; Mg chelatase
3d	HVO_B0050 (cont.)		no	33%	slr1777	P52772	(Jensen et al., 1996) (Jensen et al., 1998)	8663186 9716491	match to complete sequence, incl distant match to N-term; Mg chelatase
3e	HVO_1121	<i>ahbC</i>	yes	47%	Mbar_A1793	Q46BK8	(Bali et al., 2011)	21969545	

							(Kuhner et al., 2014)	24669201	
3e	HVO_2144	<i>ahbD</i>	yes	42%	Mbar_A1458	Q46CH7	(Kuhner et al., 2014)	24669201	
3e	HVO_2227	<i>ahbA</i>	yes	35%	-	I6UH61	(Bali et al., 2011)	21969545	
3e	HVO_2313	<i>ahbB</i>	yes	32%	-	I6UH61	(Bali et al., 2011)	21969545	

456

457 Table 3: **Proteins with open annotation issues and their Gold Standard Protein homologs (Section 3)**. For a description of this table see the
458 legend to Table 1.

3.4 Coenzymes II: coenzyme F420

Even though coenzyme F420 is predominantly associated with methanogenic archaea (Eirich et al., 1979, Jaenchen et al., 1984), it occurs also in bacteria and a small amount of this coenzyme has been detected in non-methanogenic archaea, including halophiles (Lin and White, 1986). The genes required for the biosynthesis of this coenzyme are encoded in haloarchaeal genomes, but the origin and attachment of the phospholactate moiety are not completely resolved (see below). To the best of our knowledge, only a single coenzyme F420 dependent enzymatic reaction has yet been reported for halophilic archaea (de Wit and Eker, 1987). Thus, the importance of this coenzyme in haloarchaeal biology is currently enigmatic and awaits experimental analysis.

(a) The pathway that creates the carbon backbone of this coenzyme has been reconstructed. We list the enzymes with their associated GSPs in Table 4. Coenzyme F420 contains a phospholactate moiety, which was reported to originate from 2-phospho-lactate (Grochowski et al., 2008), but this compound is not well connected to the remainder of metabolism. As summarized in Suppl.Text S1 Section 4, there are various new insights regarding this pathway from recent studies in other prokaryotes (Bashiri et al., 2019, Braga et al., 2019). To the best of our knowledge, the haloarchaeal coenzyme F420 biosynthesis pathway has never been experimentally analyzed.

(b) The prediction of coenzyme F420-specific oxidoreductases in *Mycobacterium* and actinobacteria has been reported (Selengut and Haft, 2010), leading to patterns and domains that are also found in haloarchaea. Several such enzymes are described in Suppl.Text S1 Section 4.

(c) HVO_1937 might be a coenzyme F420-dependent 5,10-methylenetetrahydrofolate reductase (see also below, C1 metabolism, and Suppl.Text S1 Section 4).

489 (d) The precursor for coenzyme F420 may be used by a photo-lyase involved in DNA
490 repair.

Section	Code	Gene	Gold Standard Protein				Reference	PMID	Comment
			isofunc	%seq_id	Locus tag	UniProt			
4a	HVO_2198	<i>cofH</i>	yes	35%	MJ1431	Q58826	(Graham et al., 2003) (Philmus et al., 2015)	14593448 25781338	
4a	HVO_2201	<i>cofG</i>	yes	43%	MJ0446	Q57888	(Graham et al., 2003) (Decamps et al., 2012) (Philmus et al., 2015)	14593448 23072415 25781338	
4a	HVO_2202	<i>cofC</i>	yes	25%	MJ0887	Q58297	(Grochowski et al., 2008) (Bashiri et al., 2019) (Braga et al., 2019)	18260642 30952857 31469543	
4a	HVO_2479	<i>cofD</i>	yes	39%	MM_1874	Q8PVT6	(Forouhar et al., 2008) (Braga et al., 2019)	18252724 31469543	
4a	HVO_2479 (cont.)		yes	32%	MJ1256	Q58653	(Graupner et al., 2002a)	11888293	
4a	HVO_1936	<i>cofE</i>	yes	47%	AF_2256	O28028	(Nocek et al., 2007)	17669425	
4a	HVO_1936 (cont.)		yes	38%	MJ0768	Q58178	(Li et al., 2003)	12911320	
4b	HVO_0433	<i>npdG</i>	yes	38%	AF_0892	O29370	(Kunow et al., 1993)	not in PubMed	
4b	HVO_B0113	-	no	27%	Rv0132c	P96809	(Purwantini and Mukhopadhyay, 2013)	24349169	too distant to assume isofunctionality
4b	HVO_B0342	-	unknown	29%	-	O93734	(Klein et al., 1996) (Aufhammer et al., 2004)	8706724 15016352	too distant to assume isofunctionality
4b	NP_1902A	-	no	28%	-	Q9UXP0	(Haase et al., 1992) (Westenberg et al., 1999)	1735436 9933933	too distant to assume isofunctionality
4b	NP_4006A	-	no	27%	MJ0870	Q58280	(Johnson and Mukhopadhyay, 2005)	16048999	too distant to assume isofunctionality
4c/5c	HVO_1937	<i>mer</i>	no	38%	MTH_1752	O27784	(te Brömmelstroet et al., 1990) (Vaupel and Thauer, 1995) (Shima et al., 2000)	2298726 7649177 10891279	
4d	HVO_2911	<i>phr2</i>	yes	62%	VNG_1335G OE_2907R	Q9HQ46 B0R5D6	(Takao et al., 1989) (McCready and Marcello, 2003)	2681164 12773185	

4d	HVO_2843	<i>phr1</i>	no	45%	sl1629	P77967	(Brudler et al., 2003)	12535521	sl1629 implicated in transcription regulation
4d	HVO_2843 (cont.)		possibly	45%	At5g24850	Q84KJ5	(Kleine et al., 2003) (Selby and Sancar, 2006)	12834405 17062752	mediates photorepair of ssDNA
4d	HVO_1234	<i>phr3</i>	possibly	40%	Atu4765	A9CH39	(Zhang et al., 2013)	23589886	

492

493 Table 4: **Proteins with open annotation issues and their Gold Standard Protein homologs (Section 4)**. For a description of this table see the
494 legend to Table 1.

3.5 Coenzymes III: coenzymes of C1 metabolism: tetrahydrofolate in haloarchaea, methanopterin in methanogens

Halophilic and methanogenic archaea use distinct coenzymes as one-carbon carrier (C1 metabolism): tetrahydrofolate in haloarchaea and methanopterin in methanogens (White, 1988, Maden, 2000). Several characterized methanogenic proteins that act on or with methanopterin have comparably close homologs in haloarchaea (Table 5), which results in misannotation of haloarchaeal proteins (e.g. in SwissProt) as being involved in methanopterin biology. We assume that the haloarchaeal proteins function with the haloarchaeal one-carbon carrier tetrahydrofolate and that this shift in coenzyme specificity is possible due to the structural similarity between methanopterin and tetrahydrofolate (a near-identical core structure consists of a pterin heterocyclic ring linked via a methylene bridge to a phenyl ring; both also have a polyglutamate tail). A detailed review on the many variants of the tetrahydrofolate biosynthetic pathway is available (de Crecy-Lagard, 2014).

(a) Folate biosynthesis requires aminobenzoate. We had proposed candidates for a pathway from chorismate to para-aminobenzoate (Falb et al., 2008, Pfeiffer et al., 2008b) (for details see Suppl.Text S1 Section 5). However, these predictions have not been adopted by KEGG (accessed April 2021) and without experimental confirmation this is unlikely to ever happen.

(b) GTP cyclohydrolase MptA (HVO_2348) catalyzes a reaction in the common part of tetrahydrofolate and methanopterin biosynthesis. The enzymes specific for methanopterin biosynthesis are absent from haloarchaea and thus the assignment of HVO_2348 to the methanopterin biosynthesis pathway in UniProt is invalid (accessed March 2021).

The next common pathway step (EC 3.1.4.56) has been resolved in *M. jannaschii* (MJ0837) but still is a pathway gap in halophilic archaea. MJ0837 is very distantly related to HVO_A0533, which thus is a promising candidate for experimental analysis.

526

527 HVO_2628 shows 30% protein sequence identity to the enzyme catalyzing the first
528 committed step to methanopterin biosynthesis. As detailed in Suppl.Text S1 Section
529 5, we consider it likely that it does not catalyze that reaction.

530

531 (c) Two enzymes that alter the oxidation level of the coenzyme-attached one-carbon
532 compound probably function with tetrahydrofolate, even though their methanogenic
533 homologs function with methanopterin. In contrast to their assignments in KEGG and
534 UniProt (as of March 2021), their probable functions are thus
535 methenyltetrahydrofolate cyclohydrolase (HVO_2573) and 5,10-
536 methylenetetrahydrofolate reductase (HVO_1937) (see Suppl.Text S1 Section 5)

537

Section	Code	Gene	Gold Standard Protein				Reference	PMID	Comment
			isofunc	%seq_id	Locus tag	UniProt			
5a	HVO_0709	<i>pabA</i>	no	47%	TTHA1843	P05379	(Sato et al., 1988)	2844259	Trp biosynthesis
5a	HVO_0709 (cont.)		yes/no	39%	BSU00750	P28819	(Slock et al., 1990)	2123867	TrpG works with TrpE and with PabB
5a	HVO_0710	<i>pabB</i>	no	46%	TTHA1844	P05378	(Sato et al., 1988)	2844259	Trp biosynthesis
5a	HVO_0710 (cont.)		yes	44%	BSU00740	P28820	(Schadt et al., 2009)	19275258	PabB; para-aminobenzoate biosynthesis
5a	HVO_0708	<i>pabC</i>	no	36%	AF_0933	O29329	(Isupov et al., 2019)	30733943	branched-chain amino acids
5b	HVO_2348	<i>mptA</i>			self		(El Yacoubi et al., 2009)	19478918	gene deletion phenotypes
5b	HVO_2348 (cont.)		yes	41%	MJ0775	Q58185	(Grochowski et al., 2007)	17497938	common part of methanopterin and tetrahydrofolate biosynthesis
5b	HVO_A0533	-	unknown	27%	MJ0837	Q58247	(Mashhadi et al., 2009)	19746965	if isofunctional would resolve a pathway gap
5b	HVO_2628	-	no	31%	AF_2089	O28190	(Scott and Rasche, 2002)	12142414	first committed step to methanopterin biosynthesis
5b	HVO_2628 (cont.)		no	26%	MJ1427	Q58822	(Dumitru and Ragsdale, 2004)	15262968	first committed step to methanopterin biosynthesis
5c	HVO_2573	<i>mch</i>	no	45%	MK0625	P94954	(Vaupel et al., 1998)	9676239	acts on a one-carbon attached to methanopterin
4c/5c	HVO_1937	<i>mer</i>	no	38%	MTH_1752	O27784	(te Brömmelstroet et al., 1990) (Vaupel and Thauer, 1995) (Shima et al., 2000)	2298726 7649177 10891279	acts on a one-carbon compound attached to methanopterin

538

539 Table 5: **Proteins with open annotation issues and their Gold Standard Protein homologs (Section 5)**. For a description of this table see the
540 legend to Table 1.

3.6 Coenzymes IV: NAD and FAD (riboflavin)

(a) The energy source for NAD kinase may be ATP or polyphosphate. This is unresolved for the two paralogs of probable NAD kinase (HVO_2363, *nadK1*, HVO_0837, *nadK2*). These show only 25% protein sequence identity to each other (see Suppl. Text S1 Section 6). Polyphosphate was not found in exponentially growing *Hfx. volcanii* cells (Zerulla et al., 2014), so that ATP is the more likely energy source.

(b) HVO_0782 is an enzyme involved in NAD biosynthesis, which is encoded in most haloarchaeal and archaeal genomes. The adjacent gene, HVO_0781, is encoded in nearly all haloarchaeal genomes according to OrthoDB, and with very strong syntenic coupling revealed by SyntTax analysis. Thus, HVO_0781 is a candidate to also be involved in NAD biosynthesis. Characterized homologs to HVO_0781 decompose S-adenosyl-methionine into methionine and adenosine, a reaction that seems wasteful and might not be expected to be highly conserved with respect to existence and gene clustering (see Suppl. Text S1 Section 6).

(c) We describe the reconstruction of riboflavin biosynthesis based on a detailed bioinformatic reconstruction (Rodionova et al., 2017). The enzymes and their associated GSPs are listed in Table 6. Three pathway gaps remain, with candidate genes predicted for two of these (Rodionova et al., 2017) (for details see Suppl. Text S1 Section 6).

563

Section	Code	Gene	Gold Standard Protein				Reference	PMID	Comment
			isofunc	%seq_id	Locus tag	UniProt			
6a	HVO_2363	<i>nadK1</i>	unclear	37%	Rv1695	P9WHV7	(Kawai et al., 2000)	11006082	can use ATP and PP
6a	HVO_2363 (cont.)		unclear	31%	AF_2373	O30297			ATP or PP usage unresolved
6a	HVO_0837	<i>nadK2</i>	unclear	28%	Rv1695	P9WHV7			can use ATP and PP
6a	HVO_0837 (cont.)		unclear	partial	AF_2373	O30297			ATP or PP usage unresolved
6b	HVO_0782	<i>nadM</i>	yes	53%	MJ0541	Q57961	(Raffaelli et al., 1997) (Raffaelli et al., 1999)	9401030 10331644	
6b	HVO_0781	-	unknown	42%	Sare_1364	A8M783	(Eustaquio et al., 2008)	18720493	
6b	HVO_0781 (cont.)		unknown	35%	PH0463	O58212	(Deng et al., 2008)	18551689	
6b	HVO_0327	<i>ribB</i>	yes	43%	MJ0055	Q60364	(Fischer et al., 2002)	12200440	
6b	HVO_0974	<i>ribH</i>	yes	45%	MJ0303	Q57751	(Haase et al., 2003)	12603336	
6b	HVO_1284	<i>arfA</i>		self			(Phillips et al., 2012)	21999246	gene deletion leads to riboflavin auxotrophy
6b	HVO_1284 (cont.)		yes	44%	MJ0145	Q57609	(Graham et al., 2002)	12475257	
6b	HVO_1235	-	prediction				(Rodionova et al., 2017)	28073944	<i>arfB</i> candidate
6b	HVO_1341	<i>arfC</i>	yes	36%	MJ0671	Q58085	(Graupner et al., 2002b) (Romisch-Margl et al., 2008)	11889103 18671734	
6b	HVO_2483	-	prediction	34%	MJ0699	Q58110	(Rodionova et al., 2017)	28073944	predicted also for MJ0699
6b	pathway gap								EC 3.1.3.104
6b	HVO_0326	<i>rbkR</i>	yes	37%	TA1064	Q9HJA6	(Rodionova et al., 2017)	28073944	bifunctional as gene regulator and enzyme
6b	HVO_0326 (cont.)		yes/no	32%	MJ0056	Q60365	(Ammelburg et al., 2007)	18073108	enzyme only; lacks an N-terminal HTH domain
6b	HVO_1015	<i>ribL</i>	yes	50%	MJ1179	Q58579	(Mashhadi et al., 2010)	20822113	

564

565 Table 6: **Proteins with open annotation issues and their Gold Standard Protein homologs (Section 6)**. For a description of this table see the
566 legend to Table 1.

3.7 Biosynthesis of membrane lipids, bacterioruberin and menaquinone

Archaeal membrane lipids contain ether-linked isoprenoid side chains (see (Caforio and Driessen, 2017) and references cited therein). The isoprenoid precursor isopentenyl diphosphate is synthesized in haloarchaea by a modified version of the mevalonate pathway (Vannice et al., 2014). Isoprenoid units are then linearly condensed to the C20 compound geranylgeranyl diphosphate. The haloarchaeal core lipid, archaeol, consists of 2,3-sn-glycerol with two C20 isoprenoid side chains attached by ether linkages. In some archaea, especially alkaliphiles, C25 isoprenoids are also found (see e.g. (De and Gambacorta, 1988, Dawson et al., 2012)). Also, a number of distinct headgroups are found in polar lipids (phospholipids) (reviewed in (Caforio and Driessen, 2017)). Even though polar lipids are used as important taxonomic markers (Oren et al., 1997) their biosynthetic pathways are not completely resolved.

Haloarchaea typically have a red color, which is due to carotenoids, mainly the C50 carotenoid bacterioruberin (Oren, 2002, Kushwaha et al., 1975, Yang et al., 2015). For carotenoid biosynthesis, two molecules of geranylgeranyl diphosphate, a C20 compound, are linked head to head to generate phytoene, which is desaturated to lycopene (Falb et al., 2008, Giani et al., 2020). The pathway from lycopene to the C50 compound bacterioruberin has been experimentally characterized (Dummer et al., 2011, Yang et al., 2015).

(a) We assigned HVO_2725 (*idsA1*, paralog of NP_3696A) and HVO_0303 (*idsA2*, paralog of NP_0604A) for the linear isoprenoid condensation reactions resulting in a C20 isoprenoid (EC 2.5.1.10, 2.5.29, short chain isoprenyl diphosphate synthase) (see also Suppl.Text S1 Section 7). Some archaea, mainly haloalkaliphiles, also contain C25 isoprenoid side chains. Geranylgeranyl diphosphate synthase, the enzyme which generates the C25 isoprenoids, has been purified and enzymatically characterized from *Nmn. pharaonis* (Tachibana, 1994), but data that allow the assignment to a specific gene have not been collected. Three paralogous genes from *Nmn. pharaonis* are candidates for this function (NP_0604A, NP_3696A, and NP_4556A). Because NP_0604A and NP_3696A have orthologs in *Hfx. volcanii*, a species devoid of C25

lipids, we assign the synthesis of C25 isoprenoids (geranylgeranyl diphosphate synthase activity) to the third paralog, NP_4556A. UniProt assigns C25 biosynthesis activity to NP_3696A for undescribed reasons (as of April 2021) and KEGG does not make this assignment for any of the three paralogs (as of April 2021). Our assignments are supported by analysis of key residues which determine the length of the isoprenoid chain (Bale et al., 2019). These authors label the cluster containing NP_3696A (WP011323557.1) as “C15/C20” and the cluster containing NP_4556A (WP011323984.1) as “C20->C25->C30?”.

(b) Typical polar lipids in haloarchaea are phosphatidylglycerophosphate methyl ester (PGP-Me) and phosphatidylglycerol (PG), but also phosphatidylglycerosulfate (PGS) (Kates, 1993, Kates et al., 1993, Bale et al., 2019). Other polar lipids are archaetidylserine and its decarboxylation product archaetidylethanolamine, both of which are found in rather low quantities in *Haloferax* (Kellermann et al., 2016). A third group of polar lipids has a headgroup derived from myo-inositol. The biosynthetic pathway of the head groups is only partially resolved. One CDP-archaeol 1-archaetidyltransferase that belongs to a highly conserved three-gene operon may attach either glycerol phosphate or myo-inositol phosphate. In Suppl. Text S1 Section 7 we summarize arguments in favor of each of these candidates, but the true function can only be decided by experimental analysis.

(c) Carotenoid biosynthesis involves the head-to-head condensation of the C20 isoprenoid geranylgeranyl diphosphate to phytoene, which is desaturated to lycopene (Falb et al., 2008, Giani et al., 2020). The *crtB* gene product (e.g. HVO_2524) catalyzes the head-to-head condensation. It is yet uncertain which gene product is responsible for the desaturation of phytoene to lycopene. The further pathway from lycopene to bacterioruberin has been experimentally characterized in *Haloarcula japonica* (Yang et al., 2015). A three gene cluster (*crtD-lyeJ-cruF*) codes for the three enzymes of this pathway. The synteny of this three gene cluster is strongly conserved according to SyntTax analysis. Several genes which are certainly or possibly involved in carotenoid biosynthesis are encoded in the vicinity of this cluster (for details see Suppl. Text S1 Section 7).

632 (d) Halophilic archaea contain menaquinone as a lipid based two electron carrier of
 633 the respiratory chain (Elling et al., 2016, Kellermann et al., 2016). We describe the
 634 reconstruction of the menaquinone biosynthesis pathway (Table 7), with two pathway
 635 gaps remaining open (see Suppl.Text S1 Section 7 for details).

Section	Code	Gene	Gold Standard Protein				Reference	PMID	Comment
			isofunc	%seq_id	Locus tag	UniProt			
7a	NP_0604A	<i>idsA2</i>	yes	32%	GACE_1337	A0A0A7GEY4	(Petrova et al., 2018)	30062607	ortholog of HVO_0303 (66%); produces a C20 isoprenoid (same assignment for NP_0604A)
7a	NP_0604A (cont.)	<i>idsA2</i>	no	30%	APE_1764	Q9YB31 Q9UWR6	(Tachibana et al., 2000)	10632701	produces a C25 isoprenoid (C20 assigned to NP_0604A)
7a	NP_3996A	<i>idsA3</i>	yes	44%	GACE_1337	A0A0A7GEY4	(Petrova et al., 2018)	30062607	ortholog of HVO_2725 (67%); produces a C20 isoprenoid (same assignment for NP_3996A)
7a	NP_3996A (cont.)	<i>idsA2</i>	no	36%	APE_1764	Q9YB31 Q9UWR6	(Tachibana et al., 2000)	10632701	produces a C25 isoprenoid (C20 assigned to NP_3996A)
7a	NP_4556A	<i>idsA1</i>	no	34%	GACE_1337	A0A0A7GEY4	(Petrova et al., 2018)	30062607	no ortholog in <i>Hfx. volcanii</i> ; produces a C20 isoprenoid (C25 assigned to NP_4556A)
7a	NP_4556A (cont.)	<i>idsA1</i>	yes	29%	APE_1764	Q9YB31 Q9UWR6	(Tachibana et al., 2000)	10632701	produces a C25 isoprenoid (same assignment for NP_4556A)
7b	HVO_0332	<i>carS</i>	yes	45%	AF_1740	O28537	(Jain et al., 2014)	25219966	
7b	HVO_1143	<i>assA</i>	yes	32%	MTH_1027	O27106	(Morii and Koga, 2003)	12562787	gene synonym: <i>pgsA3</i>
7b	HVO_1297	<i>aisA</i>	yes	25%	MTH_1691	O27726	(Morii et al., 2009)	19740749	gene synonym: <i>pgsA2</i>

7b	HVO_1136	<i>pgsA1</i>	-						only distant partial matches to GSPs
7b	HVO_1971	<i>pgsA4</i>	unclear	26%	MTH_1027	O27106	(Morii and Koga, 2003)	12562787	MTH_1027 is less distant to HVO_1143
7b	HVO_0146	<i>asd</i>	no	39%	SMc00551	Q9FDI9	(Vences-Guzman et al., 2008)	18708506	equivalent function for the bacterial lipid
7b	HVO_1295	<i>hisC</i>		self			(Conover and Doolittle, 1990)	2345144	complements a His auxotrophy mutant
7b	HVO_1295 (cont.)		yes	31%	b2021	P06986	(Grisolia et al., 1985)	2999081	weak support, see text
7b	HVO_1296	<i>adk2</i>	unclear	34%	PAB0757	Q9UZX4	(Loc'h et al., 2014)	24823650	<i>Pyrococcus</i> : involved in ribosome biogenesis
7b	HVO_1296 (cont.)		unclear	32%	-	Q9Y3D8	(Ren et al., 2005)	15630091	human: adenylate kinase; HVO_1296 may be inositol kinase
7b	HVO_2496	<i>adk1</i>	yes	45%	BSU01370	P16304	(Moon et al., 2019)	31111079	<i>Bacillus</i> : adenylate kinase
7b	HVO_B0213	-	yes	43%	AF_1794	O28480	(Chen et al., 2000) (Neelon et al., 2011)	11015222 22261071	<i>Archaeoglobus</i> : adenylate kinase
7b	HVO_1135	-	-						a SAM-dependent methyltransferase
7c	HVO_2524	<i>crtB</i>		self			(Kiljunen et al., 2014) (Maurer et al., 2018)	25488358 29038254	<i>crtB</i> mutants are colorless
7c	HVO_2524 (cont.)		yes	32%	Synpcc7942_1984	P37269	(Chamovitz et al., 1992)	1537409	
7c	HVO_2527	<i>lyeJ</i>		self			(Dummer et al., 2011)	21840984	
7c	HVO_2527 (cont.)		yes	65%	VNG_1682C OE_3380R	Q9HPD9 B0R651	(Dummer et al., 2011)	21840984	
7c	HVO_2527 (cont.)		yes	61%	C444_12922	M0L7V9	(Yang et al., 2015)	25712483	
7c	HVO_2528	<i>crtD</i>		self			(Maurer et al., 2018)	29038254	a HVO_2528 mutant was white
7c	HVO_2528 (cont.)		yes	71%	C444_12917	A0A0A1GKA2	(Yang et al., 2015)	25712483	
7c	HVO_2526	<i>cruF</i>	yes	59%	C444_12927	A0A0A1GNF2	(Yang et al., 2015)	25712483	

7d	HVO_1470	<i>menF</i>	yes	38%	PA4231	Q51508	(Serino et al., 1995)	7500944	
7d	HVO_1469	<i>menD</i>	yes	37%	BSU30820	P23970	(Dawson et al., 2010)	20600129	
7d	pathway gap								EC 4.2.99.20
7d	HVO_1461	<i>menC</i>	no	29%	BSU12980	O34508	(Schmidt et al., 2001)	11747447	Ala/Glu epimerase
7d	HVO_1461 (cont.)		yes	24%	BSU30780	O34514	(Palmer et al., 1999)	10194342	o-succinylbenzoate synthase
7d	HVO_1375	<i>menE</i>	yes	36%	BSU30790	P23971	(Chen et al., 2016)	27933791	
7d	HVO_1465	<i>menB</i>	yes	66%	Rv0548c	P9WNP5	(Jiang et al., 2010)	20643650	
7d	pathway gap								EC 3.1.2.28
7d	HVO_1462	<i>menA</i>	yes	37%	b3930	P32166	(Suvarna et al., 1998)	9573170	
7d	HVO_0309	<i>menG</i>	yes/no	44%	At3g63410	Q9LY74	(Cheng et al., 2003)	14508009	<i>A.thaliana</i> enzyme also involved in tocopherol biosynthesis
7d	HVO_0309 (cont.)		yes	27%	-	O86169	(Koike-Takeshita et al., 1997)	9139683	

637

638 Table 7: **Proteins with open annotation issues and their Gold Standard Protein homologs (Section 7)**. For a description of this table see the
639 legend to Table 1.

3.8 Issues concerning RNA polymerase, protein translation components and signal peptide degradation

(a) Haloarchaeal RNA polymerase consists of a set of canonical subunits (encoded by *rpoA1A2B1B2DEFHKLNP*). *Hbt. salinarum* and a subset of other haloarchaea contain an additional subunit called epsilon (Leffers et al., 1989, Madon and Zillig, 1983). Purified RNA polymerase containing the epsilon subunit transcribes native templates efficiently, in contrast to RNA polymerase devoid of this subunit (Madon and Zillig, 1983). The biological relevance of this subunit is enigmatic (see Suppl.Text S1 Section 8).

(b) Two distant paralogs are found for haloarchaeal ribosomal protein S10 (uS10) in nearly all haloarchaeal genomes. It is uncertain if both occur in the ribosome, whether they occur together or are mutually exclusive. The latter distribution would result in heterogeneity of the ribosomes. Alternatively, one of the paralogs may exclusively have a non-ribosomal function.

In a subset of archaea, two distant paralogs are found for haloarchaeal ribosomal protein S14 (uS14) (ca 20% of the genomes, e.g. in *Nmn. pharaonis*). For more details see Suppl.Text S1 Section 8.

(c) The ribosomal protein L43e (eL43) shows heterogeneity with respect to the present of C2-C2 type zinc finger motif. This zinc finger is found in L43e from all *Halobacteriales* and all euryarchaeal proteins outside the order *Halobacteria*, but is not found in *Haloferacales* and very rare in *Natrialbales*. Eukaryotic orthologs (e.g. from rat and yeast) contain this zinc finger and its biological importance has been experimentally shown for the yeast protein (Rivlin et al., 1999) (for details see Suppl.Text S1 Section 8).

(d) Diphthamide is a complex covalent modification of a histidine residue of translation elongation factor α -EF2. The pathway has been reconstructed (Table 8), based on distant homologs (enzymes encoded by *dph2*, *dph5*) and by detailed

671 bioinformatic analysis (enzyme encoded by *dph6*) (de Crecy-Lagard et al., 2012) (for
672 details see Suppl.Text S1 Section 8). These uncertain function assignments await
673 experimental confirmation.

674

675 (e) N-terminal signal sequences target proteins to the secretion machinery.
676 Subsequent to membrane insertion or transmembrane transfer, the signal sequence is
677 cleaved off by signal peptidase. After cleavage, the signal peptide must be degraded
678 to avoid clogging of the membrane. Degradation is catalyzed by signal peptide
679 peptidase. Candidates for this activity have been predicted from two protein families
680 (Ng et al., 2007, Raut et al., 2021) (for details see Suppl.Text S1 Section 8).

681

Section	Code	Gene	Gold Standard Protein				Reference	PMID	Comment
			isofunc	%seq_id	Locus tag	UniProt			
8a	OE_1279R	<i>rpoeps</i>		self			(Leffers et al., 1989) (Madon and Zillig, 1983)	2495365 6852054	
8b	HVO_0360	<i>rps10a</i>	yes	94%	rrnAC2405	P23357	(Arndt et al., 1991)	1764513	
8b	HVO_1392	<i>rps10b</i>	-						no GSP; 24% seq_id to HVO_0360 (<i>rps10a</i>)
8b	NP_4882A	<i>rps14a</i>	yes	72%	rrnAC1597.1	P26816	(Scholzen and Arndt, 1991)	1832208	full-length similarity; <i>Haloarcula</i> protein was not isolated or characterized
8b	NP_4882A (cont.)		yes	57%	YDL061C	P41058	(Otaka et al., 1984)	18782943	yeast YS29B; N-term 20 aa divergent
8b	NP_1768A	<i>rps14b</i>	unclear	80%	rrnAC1597.1	P26816	(Scholzen and Arndt, 1991)	1832208	N-term 20 aa divergent
8c	OE_1373R	<i>rpl43e</i>	yes	69%	rrnAC1669	P60619	(Ban et al., 2000)	10937989	
8c	OE_1373R (cont.)		yes	39%	YPR043W	P0CX25	(Rivlin et al., 1999) (Dresios et al., 2002)	10588896 11866512	
8c	HVO_0654	<i>rpl43e</i>	yes	54%	rrnAC1669	P60619	(Ban et al., 2000)	10937989	<i>Haloarcula</i> : has zinc finger; <i>Haloferax</i> : lacks zinc finger
8d	HVO_1631	<i>dph2</i>	yes	35%	PH1105	O58832	(Zhu et al., 2011)	20931132	
8d	HVO_0916	<i>dph5</i>	yes	39%	PH0725	O58456	(Zhu et al., 2010)	20873788	
8d	HVO_1077	<i>dph6</i>	yes	31%	YLR143W	Q12429	(Su et al., 2012) (Uthman et al., 2013)	23169644 23468660	
8e	HVO_0881	<i>sppA1</i>	yes	33%	BSU19530	O34525	(Bolhuis et al., 1999) (Nam et al., 2012)	10455123 22472423	
8e	HVO_1987	<i>sppA2</i>	probably	23%	BSU19530	O34525	(Bolhuis et al., 1999) (Nam et al., 2012)	10455123 22472423	
8e	HVO_1107	-	prediction						no GSP

682

683 Table 8: **Proteins with open annotation issues and their Gold Standard Protein homologs (Section 8)**. For a description of this table see the
684 legend to Table 1.

3.9 Miscellaneous metabolic enzymes and proteins with other functions

Here, we list a few other enzymatic or non-enzymatic functions for which candidate genes have been assigned, but without experimental validation.

(a) Ketohexokinase from *Haloarcula vallismortis* has been experimentally characterized (Rangaswamy and Altekari, 1994). However, the activity was not assigned to a gene. Detailed bioinformatic analyses have been made (Anderson et al., 2011, Williams et al., 2019) and point to a small set of orthologs represented by Hmuk_2662, the ortholog of HVO_1812 (for further details see Suppl. Text S1 Section 9).

(b) The assignment of fructokinase activity to the *Hht. litchfieldiae* candidate gene halTADL_1913 (UniProt:A0A1H6QYL4) is based on differential proteomic analysis (Williams et al., 2019) (see Suppl. Text S1 Section 9 for details). Very close homologs are rare in haloarchaea. For this protein family (carbohydrate kinase) it is unclear if more distant homologs (with about 50% protein sequence identity) are isofunctional.

(c) A candidate gene for glucoamylase is HVO_1711 for reasons described in Suppl. Text S1 Section 9. The enzyme from *Halorubrum sodomense* has been characterized (Chaga et al., 1993) but the activity was not yet assigned to a gene.

(d) A strong candidate for having glucose-6-phosphate isomerase activity is *Hfx. volcanii* (HVO_1967, *pgi*), based on 36% protein sequence identity to the characterized enzyme from *M. jannaschii* (MJ1605) (Rudolph et al., 2004) (Table 9).

Section	Code	Gene	Gold Standard Protein				Reference	PMID	Comment
			isofunc	%seq_id	Locus tag	UniProt			
9a	HVO_1812	-	prediction						no GSP
9b	halTADL_1913	-	yes	37%	-	P26984	(Aulkemeyer et al., 1991)	1809835	
9b	halTADL_1913 (cont.)	-	yes	31%	OCC_03567	Q7LYW8 H3ZP68	(Qu et al., 2004)	15138858	
9c	HVO_1711	-	probably	33%	-	P29761	(Ohnishi et al., 1992)	1633799	P29761 matches to C-term half of HVO_1711
9c	HVO_1711 (cont.)	-	probably	51%	SAMN04487937_2677	A0A1I6HD35	(Chaga et al., 1993)	8305855	correlation between PMID:8305855 and A0A1I6HD35 likely (see text)
9d	HVO_1967	<i>pgi</i>	yes	36%	MJ1605	Q59000	(Rudolph et al., 2004)	14655001	
9e	OE_1665R	<i>kdgA</i>	no	31%	PA1010	Q9I4W3	(Kaur et al., 2011)	21396954	GSP for <i>dapA</i> (see under 2a)
9e	OE_1665R (cont.)		probably	30%	TTX_1156.1 TTX_1156a	G4RJQ2	(Ahmed et al., 2005)	15869466	
9e	OE_1665R (cont.)		probably	25%	SSO3197	Q97U28	(Ahmed et al., 2005)	15869466	
9f	HVO_1692	<i>ludB</i>		self			(Reinhardt et al., 2019)	30707467	
9f	HVO_1692 (cont.)		probably	35%	BSU34040	O07021	(Chai et al., 2009)	19201793	matches up to HVO_1692 pos 490 of 733
9f	HVO_1692 (cont.)		probably	35%	PST_3338	O4VPR6	(Gao et al., 2015)	25917905	matches up to HVO_1692 pos 400 of 733
9f	HVO_1693	<i>ludC</i>		self			(Reinhardt et al., 2019)	30707467	
9f	HVO_1693 (cont.)		probably	30%	BSU34030	O32259	(Chai et al., 2009)	19201793	
9f	HVO_1693 (cont.)		probably	33%	PST_3339	O4VPR7	(Gao et al., 2015)	25917905	partial match
9f	HVO_1697	-	unclear	24%	PST_3340	O4VPR8	(Gao et al., 2015)	25917905	

9f	HVO_1696	<i>lctP</i>	probably	44%	PST_3336	O4VPR4	(Gao et al., 2015)	25917905	
9g	HVO_B0300	<i>pucL1</i>	yes	49%	BSU32450	O32141	(Pfrimer et al., 2010)	20168977	<i>Bacillus</i> : bifunctional, matches to C-term
9g	HVO_B0299	<i>pucM</i>	yes	43%	BSU32460	O32142	(Lee et al., 2005)	16098976	
9g	HVO_B0301	<i>pucL2</i>	yes	43%	BSU32450	O32141	(Kim et al., 2007)	17567580	<i>Bacillus</i> : bifunctional, matches to N-term
9g	HVO_B0302	<i>pucH1</i>	no	33%	-	Q8VTT5	(Xu et al., 2002)	12148274	paper in Chinese, abstract in English; pyrimidine degradation
9g	HVO_B0302 (cont.)		yes	30%	STM0523	Q7CR08	(Ho et al., 2013)	23287969	purine degradation
9g	HVO_B0302 (cont.)		yes	29%	BSU32410	O32137	(Schultz et al., 2001)	11344136	purine degradation
9g	HVO_B0306	<i>amaB4</i>	no	39%	-	Q53389	(Martinez-Rodriguez et al., 2012)	22904279	carbamoyl-AA hydrolysis
9g	HVO_B0306 (cont.)		yes	34%	At5g43600	Q8VXY9	(Werner et al., 2010) (Werner et al., 2013)	19935661 23940254	purine degradation
9g	HVO_B0308	<i>coxS</i>	no	46%	Saci_2270	Q4J6M5	(Kardinahl et al., 1999)	10095793	GAPDH
9g	HVO_B0308 (cont.)		no	41%	-	P19915	(Kang and Kim, 1999)	10482497	CO-DH
9g	HVO_B0308 (cont.)		yes	39%	b2868	Q46801	(Xi et al., 2000)	10986234	xanthine DH
9g	HVO_B0309	<i>coxL</i>	yes	33%	b2866	Q46799	(Xi et al., 2000)	10986234	xanthine DH
9g	HVO_B0309 (cont.)		no	28%	-	P19913	(Kang and Kim, 1999)	10482497	CO-DH
9g	HVO_B0309 (cont.)		no	26%	Saci_2271	Q4J6M3	(Kardinahl et al., 1999)	10095793	GAPDH
9g	HVO_B0310	<i>coxM</i>	no	31%	Saci_2269	Q4J6M6	(Kardinahl et al., 1999)	10095793	GAPDH
9g	HVO_B0310 (cont.)		no	31%	-	P19914	(Kang and Kim, 1999)	10482497	CO-DH
9g	HVO_B0310 (cont.)		yes	25%	b2867	Q46800	(Xi et al., 2000)	10986234	xanthine DH
9g	HVO_B0303	<i>uraA4</i>	yes	38%	b3654	P0AGM9	(Karatza and Frillingos, 2005)	16096267	
9h	HVO_0197	-	possibly	39%	lp_0105	F9UST0	(Desguin et al., 2016)	27114550	LarB family protein
9h	HVO_2381	-	possibly	31%	lp_0106 /	F9UST1	(Desguin et al., 2016)	27114550	LarC family protein

					lp_0107				
9h	HVO_0190	-	possibly	34%	lp_0109	F9UST4	(Desguin et al., 2016)	27114550	LarE family protein
9i	HVO_1660	<i>dacZ</i>		self			(Braun et al., 2019)	30884174	
9i	HVO_0756	-	prediction				(He et al., 2020)	32095817	
9i	HVO_0990	-	prediction				(He et al., 2020)	32095817	
9i	HVO_1690	-	prediction				(He et al., 2020)	32095817	
9j	HVO_2763	-		self				22350204	no function could be assigned
9j	HVO_2763 (cont.)		no	27%	HVO_0144	D4GZ88	(Spath et al., 2008)	18437358	Rnase Z
9k	HVO_2410	<i>dabA</i>	yes	33%	Hneap_0211	D0KWS7	(Desmarais et al., 2019)	31406332	
9k	HVO_2411	<i>dabB</i>	yes	31%	Hneap_0212	D0KWS8	(Desmarais et al., 2019)	31406332	

711

712 Table 9: **Proteins with open annotation issues and their Gold Standard Protein homologs (Section 9)**. For a description of this table see the

713 legend to Table 1.

(e) A candidate gene for specifying an enzyme with 2-dehydro-3-deoxy-(phospho)gluconate aldolase activity is *Hbt. salinarum kdgA* (OE_1665R). It is rather closely related (36% protein sequence) to *Hfx. volcanii* HVO_1101 (encoded by *dapA*), which is involved in lysine biosynthesis, a biosynthetic pathway that is absent from *Hbt. salinarum*. The function assignment is based on distant homologs from *Saccharolobus (Sulfolobus) solfataricus* and *Thermoproteus tenax* which have been characterized (Ahmed et al., 2005) (for details see Suppl. Text S1 Section 9).

(f) Haloarchaea may contain an NAD-independent L-lactate dehydrogenase, LudBC (HVO_1692 and HVO_1693). Deletion of this gene pair impairs growth on rhamnose, which is catabolized to pyruvate and lactate (Reinhardt et al., 2019). There is a very distant relationship (for details see Suppl. Text S1 Section 9) to the LldABC subunits of the characterized L-lactate dehydrogenase from *Pseudomonas stutzeri* A1501 (Gao et al., 2015) and to the LutABC proteins from *B. subtilis*, which have been shown to be involved in lactose utilization (Chai et al., 2009).

(g) *Hfx. volcanii* may be able to convert urate to allantoin, using the gene cluster HVO_B0299-HVO_B0302. This could be part of a complete degradation pathway for purines, which, however, has to be considered highly speculative (see Suppl. Text S1 Section 9).

(h) *Hfx. volcanii* may contain an enzyme having a "nickel-pincer cofactor". The biogenesis of this cofactor may be catalyzed by *larBCE* (as detailed in Suppl. Text S1 Section 9).

(i) cyclic diguanylate hydrolysis

Cyclic di-AMP (c-di-AMP) is an important nucleotide signalling molecule in bacteria and archaea. It is generated from two molecules of ATP by diadenylate cyclase (encoded by *dacZ*) and is degraded to pApA by phosphodiesterases (Corrigan and Grundling, 2013). The level of this signalling molecule is strictly controlled (Gundlach et al., 2015, Commichau et al., 2019), thus requiring a sophisticated interplay of cyclase and phosphodiesterase. *DacZ* from *Hfx. volcanii* has been

characterized and it was shown that c-di-AMP levels must be tightly regulated (Braun et al., 2019). The degrading enzyme, however, has not yet been identified in *Haloferax* but candidates have been proposed (Corrigan and Grundling, 2013, He et al., 2020, Yin et al., 2020) (see Suppl. Text S1 Section 9).

(j) homolog to RNaseZ

HVO_2763 is distantly related to RNase Z (HVO_0144, *rnz*). The experimental characterization of HVO_2763 (Fischer et al., 2012) excluded activity as exonuclease but did not reveal its physiological function. Upon transcriptome analysis, the downregulation of several genes was detected. Several of these were uncharacterized at the time of the experiment but have since been shown to be involved in the minor N-glycosylation pathway that was initially detected under low salt conditions (see Suppl. Text S1 Section 9 for further details) .

(k) A pair of genes (*dabAB*) is predicted to function as CO₂ transporter.

HVO_2410 and HVO_2411 probably function as carbon dioxide transporter, based on the identification of such transporters in *Halothiobacillus neapolitanus* (Desmarais et al., 2019). Being a member of the proton-conducting membrane transporter family, this protein may be misannotated as a subunit of the *nuo* or *mrp* complex (see Suppl. Text S1 Section 9 for further details).

4. Conclusion

We describe a large number of cases where protein function cannot be correctly predicted when restricting considerations to computational analyses without taking the biological context into account. An example is the switch from methanopterin to tetrahydrofolate as C1 carrier in haloarchaea. Homologous enzymes, inherited from the common ancestor, have adapted to the new C1 carrier, rather being replaced by non-homologous proteins. Function prediction tools may misannotate haloarchaeal proteins to work with methanopterin. Another example is the *nuo* complex and its misannotation as a type I NADH dehydrogenase. In other cases, even distant sequence similarity may allow a valid function prediction if additional evidence (e.g. from gene

777 neighbourhood analysis or from detailed evaluation of metabolic pathway gaps) is
 778 taken into account. Examples include cobalamin cluster proteins, which probably
 779 close the two residual pathway gaps, and the predicted degradation pathway for
 780 purines. In all these cases, we have presented reasonable hypotheses based on current
 781 knowledge, and in many cases these are so well supported as to be compelling, but to
 782 be certain, experimental data are required. With this overview, we attempt to arouse
 783 the curiosity of colleagues, hoping that they will confirm or disprove our speculations
 784 and thus advance the knowledge about haloarchaeal biology. *Hfx. volcanii* is a model
 785 species for halophilic archaea, and the more complete and correctly its genome is
 786 annotated, the higher will be its value for systems biology analyses (modelling) and
 787 for synthetic biology (metabolic engineering) and biotechnology.

Acknowledgments

We thank all members of the *Haloferax* community for more than a decade of fruitful cooperation and for sharing their deep knowledge about the biology of this model species. We thank our colleagues for kindly reading and providing thoughtful comments on the manuscript before submission: Sonja-Verena Albers, Thorsten Allers, Maria Jose Bonete, Rosana de Castro, Sebastien Ferreira-Cerca, Anita Marchfelder, Mechthild Pohlschroder, Joerg Soppa. We thank Birgit Scharf for insightful discussions on the *Natronomonas* respiratory chain.

Data Availability Statement

None.

Funding

This research received no specific grant from any funding agency in the public, commercial, or non-for-profit sectors.

Authors contribution

Conceptualization, F.P.

Data curation, F.P.

Project administration, F.P.

Formal analysis, F.P., M.D-S.

Investigation F.P., M.D-S.

Validation, F.P., M.D-S.

Visualization, F.P., M.D-S.

Writing – original draft, F.P., M.D-S.;

Writing – review & editing, F.P., M.D-S.

Conflict of interest

None declared.

817

818 **Ethics statement**

819 Not applicable

REFERENCES

- ABDUL HALIM, M. F., PFEIFFER, F., ZOU, J., FRISCH, A., HAFT, D., WU, S.,
TOLIC, N., BREWER, H., PAYNE, S. H., PASA-TOLIC, L. &
POHLSCHRODER, M. 2013. *Haloferax volcanii* archaeosortase is required
for motility, mating, and C-terminal processing of the S-layer glycoprotein. *Mol
Microbiol*, 88, 1164-75.
- ABDUL HALIM, M. F., RODRIGUEZ, R., STOLTZFUS, J. D., DUGGIN, I. G. &
POHLSCHRODER, M. 2018. Conserved residues are critical for *Haloferax
volcanii* archaeosortase catalytic activity: Implications for convergent
evolution of the catalytic mechanisms of non-homologous sortases from
archaea and bacteria. *Mol Microbiol*, 108, 276-87.
- ABDUL-HALIM, M. F., SCHULZE, S., DILUCIDO, A., PFEIFFER, F., BISSON
FILHO, A. W. & POHLSCHRODER, M. 2020. Lipid anchoring of
archaeosortase substrates and midcell growth in haloarchaea. *mBio*, 11,
e00349-20.
- AHMED, H., ETTEMA, T. J., TJADEN, B., GEERLING, A. C., VAN DER OOST, J. &
SIEBERS, B. 2005. The semi-phosphorylative Entner-Doudoroff pathway in
hyperthermophilic archaea: a re-evaluation. *Biochem J*, 390, 529-40.
- AL-MAILEM, D. M., HOUGH, D. W. & DANSON, M. J. 2008. The 2-oxoacid
dehydrogenase multienzyme complex of *Haloferax volcanii*. *Extremophiles*,
12, 89-96.
- ALLERS, T., BARAK, S., LIDDELL, S., WARDELL, K. & MEVARECH, M. 2010.
Improved strains and plasmid vectors for conditional overexpression of His-
tagged proteins in *Haloferax volcanii*. *Appl Environ Microbiol*, 76, 1759-69.
- ALLERS, T. & MEVARECH, M. 2005. Archaeal genetics - the third way. *Nat. Rev.
Genet.*, 6, 58-73.
- ALTSCHUL, S. F., MADDEN, T. L., SCHAFFER, A. A., ZHANG, J., ZHANG, Z.,
MILLER, W. & LIPMAN, D. J. 1997. Gapped BLAST and PSI-BLAST: a new
generation of protein database search programs. *Nucleic Acids Res*, 25,
3389-402.
- AMMELBURG, M., HARTMANN, M. D., DJURANOVIC, S., ALVA, V., KORETKE, K.
K., MARTIN, J., SAUER, G., TRUFFAULT, V., ZETH, K., LUPAS, A. N. &
COLES, M. 2007. A CTP-dependent archaeal riboflavin kinase forms a bridge
in the evolution of cradle-loop barrels. *Structure*, 15, 1577-90.
- ANDERSON, I., SCHEUNER, C., GOKER, M., MAVROMATIS, K., HOOPER, S. D.,
PORAT, I., KLENK, H. P., IVANOVA, N. & KYRPIDES, N. 2011. Novel

857 insights into the diversity of catabolic metabolism from ten haloarchaeal
858 genomes. *PLoS One*, 6, e20237.

859 ARNDT, E., SCHOLZEN, T., KROMER, W., HATAKEYAMA, T. & KIMURA, M. 1991.
860 Primary structures of ribosomal proteins from the archaebacterium
861 *Halobacterium marismortui* and the eubacterium *Bacillus stearothermophilus*.
862 *Biochimie*, 73, 657-68.

863 AUFHAMMER, S. W., WARKENTIN, E., BERK, H., SHIMA, S., THAUER, R. K. &
864 ERMLER, U. 2004. Coenzyme binding in F420-dependent secondary alcohol
865 dehydrogenase, a member of the bacterial luciferase family. *Structure*, 12,
866 361-70.

867 AULKEMEYER, P., EBNER, R., HEILENMANN, G., JAHREIS, K., SCHMID, K.,
868 WRIEDEN, S. & LENGELER, J. W. 1991. Molecular analysis of two
869 fructokinases involved in sucrose metabolism of enteric bacteria. *Mol*
870 *Microbiol*, 5, 2913-22.

871 BABSKI, J., MAIER, L. K., HEYER, R., JASCHINSKI, K., PRASSE, D., JAGER, D.,
872 RANDAU, L., SCHMITZ, R. A., MARCHFELDER, A. & SOPPA, J. 2014.
873 Small regulatory RNAs in Archaea. *RNA Biol*, 11, 484-93.

874 BALE, N. J., SOROKIN, D. Y., HOPMANS, E. C., KOENEN, M., RIJPSTRA, W. I. C.,
875 VILLANUEVA, L., WIENK, H. & SINNINGHE DAMSTE, J. S. 2019. New
876 insights into the polar lipid composition of extremely halo(alkali)philic
877 euryarchaea from hypersaline lakes. *Front Microbiol*, 10, 377.

878 BALI, S., LAWRENCE, A. D., LOBO, S. A., SARAIVA, L. M., GOLDING, B. T.,
879 PALMER, D. J., HOWARD, M. J., FERGUSON, S. J. & WARREN, M. J.
880 2011. Molecular hijacking of siroheme for the synthesis of heme and d1
881 heme. *Proc Natl Acad Sci U S A*, 108, 18260-5.

882 BAN, N., NISSEN, P., HANSEN, J., MOORE, P. B. & STEITZ, T. A. 2000. The
883 complete atomic structure of the large ribosomal subunit at 2.4 Å resolution.
884 *Science*, 289, 905-20.

885 BASHIRI, G., ANTONEY, J., JIRGIS, E. N. M., SHAH, M. V., NEY, B., COPP, J.,
886 STUTELEY, S. M., SREEBHAVAN, S., PALMER, B., MIDDLEDITCH, M.,
887 TOKURIKI, N., GREENING, C., SCOTT, C., BAKER, E. N. & JACKSON, C.
888 J. 2019. A revised biosynthetic pathway for the cofactor F420 in prokaryotes.
889 *Nat Commun*, 10, 1558.

890 BOLHUIS, A., MATZEN, A., HYYRYLAINEN, H. L., KONTINEN, V. P., MEIMA, R.,
891 CHAPUIS, J., VENEMA, G., BRON, S., FREUDL, R. & VAN DIJL, J. M. 1999.
892 Signal peptide peptidase- and ClpP-like proteins of *Bacillus subtilis* required

893 for efficient translocation and processing of secretory proteins. *J Biol Chem*,
894 274, 24585-92.

895 BRAGA, D., LAST, D., HASAN, M., GUO, H., LEICHNITZ, D., UZUM, Z., RICHTER,
896 I., SCHALK, F., BEEMELMANN, C., HERTWECK, C. & LACKNER, G.
897 2019. Metabolic pathway rerouting in *Paraburkholderia rhizoxinica* evolved
898 long-overlooked derivatives of coenzyme F420. *ACS Chem Biol*, 14, 2088-94.

899 BRASEN, C. & SCHONHEIT, P. 2001. Mechanisms of acetate formation and acetate
900 activation in halophilic archaea. *Arch Microbiol*, 175, 360-8.

901 BRAUN, F., THOMALLA, L., VAN DER DOES, C., QUAX, T. E. F., ALLERS, T.,
902 KAEVER, V. & ALBERS, S. V. 2019. Cyclic nucleotides in archaea: Cyclic di-
903 AMP in the archaeon *Haloferax volcanii* and its putative role.
904 *Microbiologyopen*, 8, e00829.

905 BRAUN, M., BUNGERT, S. & FRIEDRICH, T. 1998. Characterization of the
906 overproduced NADH dehydrogenase fragment of the NADH:ubiquinone
907 oxidoreductase (complex I) from *Escherichia coli*. *Biochemistry*, 37, 1861-7.

908 BRINDLEY, A. A., RAUX, E., LEECH, H. K., SCHUBERT, H. L. & WARREN, M. J.
909 2003. A story of chelatase evolution: identification and characterization of a
910 small 13-15-kDa "ancestral" cobaltochelatase (CbiXS) in the archaea. *J Biol*
911 *Chem*, 278, 22388-95.

912 BRUDLER, R., HITOMI, K., DAIYASU, H., TOH, H., KUCHO, K.-I., ISHIURA, M.,
913 KANEHISA, M., ROBERTS, V. A., TODO, T., TAINER, J. A. & GETZOFF, E.
914 D. 2003. Identification of a new cryptochrome class. *Molecular Cell*, 11, 59-
915 67.

916 BRUSHABER, K. R., O'TOOLE, G. A. & ESCALANTE-SEMERENA, J. C. 1998.
917 CobD, a novel enzyme with L-threonine-O-3-phosphate decarboxylase
918 activity, is responsible for the synthesis of (R)-1-amino-2-propanol O-2-
919 phosphate, a proposed new intermediate in cobalamin biosynthesis in
920 *Salmonella typhimurium* LT2. *J Biol Chem*, 273, 2684-91.

921 BUAN, N. R., REHFELD, K. & ESCALANTE-SEMERENA, J. C. 2006. Studies of the
922 CobA-type ATP:Co(I)rrinoid adenosyltransferase enzyme of *Methanosarcina*
923 *mazei* strain Go1. *J Bacteriol*, 188, 3543-50.

924 CAFORIO, A. & DRIESSEN, A. J. M. 2017. Archaeal phospholipids: Structural
925 properties and biosynthesis. *Biochim Biophys Acta Mol Cell Biol Lipids*, 1862,
926 1325-39.

927 CAO, S., HEPOWIT, N. & MAUPIN-FURLOW, J. A. 2015. Ubiquitin-like protein
928 SAMP1 and JAMM/MPN+ metalloprotease HvJAMM1 constitute a system for

929 reversible regulation of metabolic enzyme activity in Archaea. *PLoS One*, 10,
930 e0128399.

931 CERLETTI, M., MARTINEZ, M. J., GIMENEZ, M. I., SASTRE, D. E., PAGGI, R. A. &
932 DE CASTRO, R. E. 2014. The LonB protease controls membrane lipids
933 composition and is essential for viability in the extremophilic haloarchaeon
934 *Haloferax volcanii*. *Environ Microbiol*, 16, 1779-92.

935 CERLETTI, M., PAGGI, R., TROETSCH, C., FERRARI, M. C., GUEVARA, C. R.,
936 ALBAUM, S., POETSCH, A. & DE CASTRO, R. 2018. LonB protease is a
937 novel regulator of carotenogenesis controlling degradation of phytoene
938 synthase in *Haloferax volcanii*. *J Proteome Res*, 17, 1158-71.

939 CHAGA, G., PORATH, J. & ILLENI, T. 1993. Isolation and purification of
940 amyloglucosidase from *Halobacterium sodomense*. *Biomed Chromatogr*, 7,
941 256-61.

942 CHAI, Y., KOLTER, R. & LOSICK, R. 2009. A widely conserved gene cluster
943 required for lactate utilization in *Bacillus subtilis* and its involvement in biofilm
944 formation. *J Bacteriol*, 191, 2423-30.

945 CHAMOVITZ, D., MISAWA, N., SANDMANN, G. & HIRSCHBERG, J. 1992.
946 Molecular cloning and expression in *Escherichia coli* of a cyanobacterial gene
947 coding for phytoene synthase, a carotenoid biosynthesis enzyme. *FEBS*
948 *Letters*, 296, 305-10.

949 CHEN, L., ZHOU, C., YANG, H. & ROBERTS, M. F. 2000. Inositol-1-phosphate
950 synthase from *Archaeoglobus fulgidus* is a class II aldolase. *Biochemistry*, 39,
951 12415-23.

952 CHEN, Y., JIANG, Y. & GUO, Z. 2016. Mechanistic insights from the crystal structure
953 of *Bacillus subtilis* o-succinylbenzoyl-CoA synthetase complexed with the
954 adenylate intermediate. *Biochemistry*, 55, 6685-95.

955 CHENG, Z., SATTler, S., MAEDA, H., SAKURAGI, Y., BRYANT, D. A. &
956 DELLAPENNA, D. 2003. Highly divergent methyltransferases catalyze a
957 conserved reaction in tocopherol and plastoquinone synthesis in
958 cyanobacteria and photosynthetic eukaryotes. *Plant Cell*, 15, 2343-56.

959 COHEN-KUPIEC, R., KUPIEC, M., SANDBECK, K. & LEIGH, J. A. 1999. Functional
960 conservation between the argininosuccinate lyase of the archaeon
961 *Methanococcus maripaludis* and the corresponding bacterial and eukaryal
962 genes. *FEMS Microbiol Lett*, 173, 231-8.

963 COLLINS, M., AFOLAYAN, S., IGIRANEZA, A. B., SCHILLER, H., KRESPAN, E.,
964 BEITING, D. P., DYALL-SMITH, M., PFEIFFER, F. & POHLSCHRODER, M.
965 2020. Mutations affecting HVO_1357 or HVO_2248 cause hypermotility in

966 *Haloferax volcanii*, suggesting roles in motility regulation. *Genes (Basel)*, 12,
967 58.

968 COMMICHAU, F. M., HEIDEMANN, J. L., FICNER, R. & STULKE, J. 2019. Making
969 and breaking of an essential poison: the cyclases and phosphodiesterases
970 that produce and degrade the essential second messenger cyclic di-AMP in
971 bacteria. *J Bacteriol*, 201, e00462-18.

972 CONOVER, R. K. & DOOLITTLE, W. F. 1990. Characterization of a gene involved in
973 histidine biosynthesis in *Halobacterium (Haloferax) volcanii*: isolation and
974 rapid mapping by transformation of an auxotroph with cosmid DNA. *J*
975 *Bacteriol*, 172, 3244-9.

976 CORRIGAN, R. M. & GRUNDLING, A. 2013. Cyclic di-AMP: another second
977 messenger enters the fray. *Nat Rev Microbiol*, 11, 513-24.

978 COSTA, M. I., CERLETTI, M., PAGGI, R. A., TROTSCHER, C., DE CASTRO, R. E.,
979 POETSCH, A. & GIMENEZ, M. I. 2018. *Haloferax volcanii* proteome response
980 to deletion of a rhomboid protease gene. *J Proteome Res*, 17, 961-77.

981 CROFTS, A. R., HONG, S., WILSON, C., BURTON, R., VICTORIA, D., HARRISON,
982 C. & SCHULTEN, K. 2013. The mechanism of ubihydroquinone oxidation at
983 the Qo-site of the cytochrome bc1 complex. *Biochim Biophys Acta*, 1827,
984 1362-77.

985 DANCHIN, A., OUZOUNIS, C., TOKUYASU, T. & ZUCKER, J. D. 2018. No wisdom
986 in the crowd: genome annotation in the era of big data - current status and
987 future prospects. *Microb Biotechnol*, 11, 588-605.

988 DAWSON, A., CHEN, M., FYFE, P. K., GUO, Z. & HUNTER, W. N. 2010. Structure
989 and reactivity of *Bacillus subtilis* MenD catalyzing the first committed step in
990 menaquinone biosynthesis. *J Mol Biol*, 401, 253-64.

991 DAWSON, K. S., FREEMAN, K. H. & MACALADY, J. L. 2012. Molecular
992 characterization of core lipids from halophilic archaea grown under different
993 salinity conditions. *Organic Geochemistry*, 48, 1-8.

994 DE CRECY-LAGARD, V. 2014. Variations in metabolic pathways create challenges
995 for automated metabolic reconstructions: Examples from the tetrahydrofolate
996 synthesis pathway. *Comput Struct Biotechnol J*, 10, 41-50.

997 DE CRECY-LAGARD, V., FOROUHAR, F., BROCHIER-ARMANET, C., TONG, L. &
998 HUNT, J. F. 2012. Comparative genomic analysis of the DUF71/COG2102
999 family predicts roles in diphthamide biosynthesis and B12 salvage. *Biol*
1000 *Direct*, 7, 32.

1001 DE, R. M. & GAMBACORTA, A. 1988. The lipids of archaebacteria. *Prog Lipid Res*,
1002 27, 153-75.

1003 DE SILVA, R. T., ABDUL-HALIM, M. F., PITTRICH, D. A., BROWN, H. J.,
1004 POHLSCHRODER, M. & DUGGIN, I. G. 2021. Improved growth and
1005 morphological plasticity of *Haloferax volcanii*. *Microbiology (Reading)*, 167.
1006 DE WIT, L. E. A. & EKER, A. P. M. 1987. 8-Hydroxy-5-deazaflavin-dependent
1007 electron transfer in the extreme halophile *Halobacterium cutirubrum*. *FEMS*
1008 *Microbiology Letters*, 48, 121-25.
1009 DEBUSSCHE, L., COUDER, M., THIBAUT, D., CAMERON, B., CROUZET, J. &
1010 BLANCHE, F. 1992. Assay, purification, and characterization of
1011 cobaltochelatase, a unique complex enzyme catalyzing cobalt insertion in
1012 hydrogenobyrinic acid a,c-diamide during coenzyme B12 biosynthesis in
1013 *Pseudomonas denitrificans*. *J Bacteriol*, 174, 7445-51.
1014 DECAMPS, L., PHILMUS, B., BENJIDIA, A., WHITE, R., BEGLEY, T. P. &
1015 BERTEAU, O. 2012. Biosynthesis of F0, precursor of the F420 cofactor,
1016 requires a unique two radical-SAM domain enzyme and tyrosine as substrate.
1017 *J Am Chem Soc*, 134, 18173-6.
1018 DENDA, K., FUJIWARA, T., SEKI, M., YOSHIDA, M., FUKUMORI, Y. &
1019 YAMANAKA, T. 1991. Molecular cloning of the cytochrome aa3 gene from the
1020 archaeon (Archaeobacterium) *Halobacterium halobium*. *Biochem Biophys Res*
1021 *Commun*, 181, 316-22.
1022 DENG, H., BOTTING, C. H., HAMILTON, J. T., RUSSELL, R. J. & O'HAGAN, D.
1023 2008. S-adenosyl-L-methionine:hydroxide adenosyltransferase: a SAM
1024 enzyme. *Angew Chem Int Ed Engl*, 47, 5357-61.
1025 DESGUIN, B., SOUMILLION, P., HOLS, P. & HAUSINGER, R. P. 2016. Nickel-
1026 pincer cofactor biosynthesis involves LarB-catalyzed pyridinium carboxylation
1027 and LarE-dependent sacrificial sulfur insertion. *Proc Natl Acad Sci U S A*,
1028 113, 5598-603.
1029 DESMARAIS, J. J., FLAMHOLZ, A. I., BLIKSTAD, C., DUGAN, E. J., LAUGHLIN, T.
1030 G., OLTROGGE, L. M., CHEN, A. W., WETMORE, K., DIAMOND, S., WANG,
1031 J. Y. & SAVAGE, D. F. 2019. DABs are inorganic carbon pumps found
1032 throughout prokaryotic phyla. *Nat Microbiol*, 4, 2204-15.
1033 DRESIOS, J., CHAN, Y. L. & WOOL, I. G. 2002. The role of the zinc finger motif and
1034 of the residues at the amino terminus in the function of yeast ribosomal
1035 protein YL37a. *J Mol Biol*, 316, 475-88.
1036 DUGGIN, I. G., AYLETT, C. H., WALSH, J. C., MICHIE, K. A., WANG, Q.,
1037 TURNBULL, L., DAWSON, E. M., HARRY, E. J., WHITCHURCH, C. B.,
1038 AMOS, L. A. & LOWE, J. 2015. CetZ tubulin-like proteins control archaeal cell
1039 shape. *Nature*, 519, 362-5.

1040 DUMITRU, R. V. & RAGSDALE, S. W. 2004. Mechanism of 4-(beta-D-
1041 ribofuranosyl)aminobenzene 5'-phosphate synthase, a key enzyme in the
1042 methanopterin biosynthetic pathway. *J Biol Chem*, 279, 39389-95.

1043 DUMMER, A. M., BONSALL, J. C., CIHLA, J. B., LAWRY, S. M., JOHNSON, G. C. &
1044 PECK, R. F. 2011. Bacterioopsin-mediated regulation of bacterioruberin
1045 biosynthesis in *Halobacterium salinarum*. *J Bacteriol*, 193, 5658-67.

1046 EL YACOUBI, B., PHILLIPS, G., BLABY, I. K., HAAS, C. E., CRUZ, Y.,
1047 GREENBERG, J. & DE CRECY-LAGARD, V. 2009. A Gateway platform for
1048 functional genomics in *Haloferax volcanii*: deletion of three tRNA modification
1049 genes. *Archaea*, 2, 211-9.

1050 ELLING, F. J., BECKER, K. W., KONNEKE, M., SCHRODER, J. M., KELLERMANN,
1051 M. Y., THOMM, M. & HINRICHS, K. U. 2016. Respiratory quinones in
1052 Archaea: phylogenetic distribution and application as biomarkers in the
1053 marine environment. *Environ Microbiol*, 18, 692-707.

1054 EUSTAQUIO, A. S., HARLE, J., NOEL, J. P. & MOORE, B. S. 2008. S-Adenosyl-L-
1055 methionine hydrolase (adenosine-forming), a conserved bacterial and
1056 archaeal protein related to SAM-dependent halogenases. *ChemBiochem*, 9,
1057 2215-9.

1058 FAEHNLE, C. R., OHREN, J. F. & VIOLA, R. E. 2005. A new branch in the family:
1059 structure of aspartate-beta-semialdehyde dehydrogenase from
1060 *Methanococcus jannaschii*. *J Mol Biol*, 353, 1055-68.

1061 FALB, M., MULLER, K., KONIGSMAIER, L., OBERWINKLER, T., HORN, P., VON
1062 GRONAU, S., GONZALEZ, O., PFEIFFER, F., BORNBERG-BAUER, E. &
1063 OESTERHELT, D. 2008. Metabolism of halophilic archaea. *Extremophiles*,
1064 12, 177-96.

1065 FALB, M., PFEIFFER, F., PALM, P., RODEWALD, K., HICKMANN, V., TITTOR, J. &
1066 OESTERHELT, D. 2005. Living with two extremes: conclusions from the
1067 genome sequence of *Natronomonas pharaonis*. *Genome Res*, 15, 1336-43.

1068 FISCHER, M., ROMISCH, W., SCHIFFMANN, S., KELLY, M., OSCHKINAT, H.,
1069 STEINBACHER, S., HUBER, R., EISENREICH, W., RICHTER, G. &
1070 BACHER, A. 2002. Biosynthesis of riboflavin in archaea studies on the
1071 mechanism of 3,4-dihydroxy-2-butanone-4-phosphate synthase of
1072 *Methanococcus jannaschii*. *J Biol Chem*, 277, 41410-6.

1073 FISCHER, S., JOHN VON FREYEND, S., SABAG-DAIGLE, A., DANIELS, C. J.,
1074 ALLERS, T. & MARCHFELDER, A. 2012. Assigning a function to a conserved
1075 archaeal metallo-beta-lactamase from *Haloferax volcanii*. *Extremophiles*, 16,
1076 333-43.

1077 FONSECA, M. V., BUAN, N. R., HORSWILL, A. R., RAYMENT, I. & ESCALANTE-
1078 SEMERENA, J. C. 2002. The ATP:Co(I)rrinoid adenosyltransferase (CobA)
1079 enzyme of *Salmonella enterica* requires the 2'-OH group of ATP for function
1080 and yields inorganic triphosphate as its reaction byproduct. *J Biol Chem*, 277,
1081 33127-31.

1082 FOROUHAR, F., ABASHIDZE, M., XU, H., GROCHOWSKI, L. L., SEETHARAMAN,
1083 J., HUSSAIN, M., KUZIN, A., CHEN, Y., ZHOU, W., XIAO, R., ACTON, T. B.,
1084 MONTELIONE, G. T., GALINIER, A., WHITE, R. H. & TONG, L. 2008.
1085 Molecular insights into the biosynthesis of the F420 coenzyme. *J Biol Chem*,
1086 283, 11832-40.

1087 FRESQUET, V., WILLIAMS, L. & RAUSHEL, F. M. 2004. Mechanism of cobyrinic
1088 acid a,c-diamide synthetase from *Salmonella typhimurium* LT2. *Biochemistry*,
1089 43, 10619-27.

1090 FUJITA, S., CHO, S. H., YOSHIDA, A., HASEBE, F., TOMITA, T., KUZUYAMA, T. &
1091 NISHIYAMA, M. 2017. Crystal structure of LysK, an enzyme catalyzing the
1092 last step of lysine biosynthesis in *Thermus thermophilus*, in complex with
1093 lysine: Insight into the mechanism for recognition of the amino-group carrier
1094 protein, LysW. *Biochem Biophys Res Commun*, 491, 409-15.

1095 FUJIWARA, T., FUKUMORI, Y. & YAMANAKA, T. 1989. Purification and properties
1096 of *Halobacterium halobium* "cytochrome aa3" which lacks CuA and CuB. *J*
1097 *Biochem (Tokyo)*, 105, 287-92.

1098 GAO, C., WANG, Y., ZHANG, Y., LV, M., DOU, P., XU, P. & MA, C. 2015. NAD-
1099 independent L-lactate dehydrogenase required for L-lactate utilization in
1100 *Pseudomonas stutzeri* A1501. *J Bacteriol*, 197, 2239-47.

1101 GIANI, M., MIRALLES-ROBLEDILLO, J. M., PEIRO, G., PIRE, C. & MARTINEZ-
1102 ESPINOSA, R. M. 2020. Deciphering pathways for carotenogenesis in
1103 haloarchaea. *Molecules*, 25, 1197.

1104 GRADIN, C. H., HEDERSTEDT, L. & BALTSCHIEFFSKY, H. 1985. Soluble succinate
1105 dehydrogenase from the halophilic archaebacterium, *Halobacterium*
1106 *halobium*. *Arch Biochem Biophys*, 239, 200-5.

1107 GRAHAM, D. E., XU, H. & WHITE, R. H. 2002. A member of a new class of GTP
1108 cyclohydrolases produces formylaminopyrimidine nucleotide
1109 monophosphates. *Biochemistry*, 41, 15074-84.

1110 GRAHAM, D. E., XU, H. & WHITE, R. H. 2003. Identification of the 7,8-didemethyl-8-
1111 hydroxy-5-deazariboflavin synthase required for coenzyme F(420)
1112 biosynthesis. *Arch Microbiol*, 180, 455-64.

1113 GRAUPNER, M., XU, H. & WHITE, R. H. 2002a. Characterization of the 2-phospho-
1114 L-lactate transferase enzyme involved in coenzyme F(420) biosynthesis in
1115 *Methanococcus jannaschii*. *Biochemistry*, 41, 3754-61.

1116 GRAUPNER, M., XU, H. & WHITE, R. H. 2002b. The pyrimidine nucleotide
1117 reductase step in riboflavin and F(420) biosynthesis in archaea proceeds by
1118 the eukaryotic route to riboflavin. *J Bacteriol*, 184, 1952-7.

1119 GRISOLIA, V., CARLOMAGNO, M. S., NAPPO, A. G. & BRUNI, C. B. 1985. Cloning,
1120 structure, and expression of the *Escherichia coli* K-12 *hisC* gene. *J Bacteriol*,
1121 164, 1317-23.

1122 GROCHOWSKI, L. L., XU, H., LEUNG, K. & WHITE, R. H. 2007. Characterization of
1123 an Fe(2+)-dependent archaeal-specific GTP cyclohydrolase, MptA, from
1124 *Methanocaldococcus jannaschii*. *Biochemistry*, 46, 6658-67.

1125 GROCHOWSKI, L. L., XU, H. & WHITE, R. H. 2008. Identification and
1126 characterization of the 2-phospho-L-lactate guanylyltransferase involved in
1127 coenzyme F420 biosynthesis. *Biochemistry*, 47, 3033-7.

1128 GULKO, M. K., DYALL-SMITH, M., GONZALEZ, O. & OESTERHELT, D. 2014. How
1129 do haloarchaea synthesize aromatic amino acids? *PLoS One*, 9, e107475.

1130 GUNDLACH, J., MEHNE, F. M., HERZBERG, C., KAMPF, J., VALERIUS, O.,
1131 KAEVER, V. & STULKE, J. 2015. An essential poison: synthesis and
1132 degradation of cyclic di-AMP in *Bacillus subtilis*. *J Bacteriol*, 197, 3265-74.

1133 GUO, R., GU, J., ZONG, S., WU, M. & YANG, M. 2018. Structure and mechanism of
1134 mitochondrial electron transport chain. *Biomed J*, 41, 9-20.

1135 HAASE, I., MORTL, S., KOHLER, P., BACHER, A. & FISCHER, M. 2003.
1136 Biosynthesis of riboflavin in archaea. 6,7-dimethyl-8-ribityllumazine synthase
1137 of *Methanococcus jannaschii*. *Eur J Biochem*, 270, 1025-32.

1138 HAASE, P., DEPPENMEIER, U., BLAUT, M. & GOTTSCHALK, G. 1992. Purification
1139 and characterization of F420H₂-dehydrogenase from *Methanlobus tindarius*.
1140 *Eur J Biochem*, 203, 527-31.

1141 HAQUE, R. U., PARADISI, F. & ALLERS, T. 2020. *Haloferax volcanii* for
1142 biotechnology applications: challenges, current state and perspectives. *Appl*
1143 *Microbiol Biotechnol*, 104, 1371-82.

1144 HARTMAN, A. L., NORRIS, C., BADGER, J. H., DELMAS, S., HALDENBY, S.,
1145 MADUPU, R., ROBINSON, J., KHOURI, H., REN, Q., LOWE, T. M.,
1146 MAUPIN-FURLOW, J., POHLSCHRODER, M., DANIELS, C., PFEIFFER, F.,
1147 ALLERS, T. & EISEN, J. A. 2010. The complete genome sequence of
1148 *Haloferax volcanii* DS2, a model archaeon. *PLoS One*, 5, e9605.

1149 HARTWELL, L. H. & MAGASANIK, B. 1963. The molecular basis of histidase
1150 induction in *Bacillus subtilis*. *J Mol Biol*, 7, 401-20.

1151 HATTORI, T., SHIBA, H., ASHIKI, K., ARAKI, T., NAGASHIMA, Y. K.,
1152 YOSHIMATSU, K. & FUJIWARA, T. 2016. Anaerobic growth of haloarchaeon
1153 *Haloferax volcanii* by denitrification is controlled by the transcription regulator
1154 NarO. *J Bacteriol*, 198, 1077-86.

1155 HE, J., YIN, W., GALPERIN, M. Y. & CHOU, S. H. 2020. Cyclic di-AMP, a second
1156 messenger of primary importance: tertiary structures and binding
1157 mechanisms. *Nucleic Acids Res*, 48, 2807-29.

1158 HEATH, C., POSNER, M. G., AASS, H. C., UPADHYAY, A., SCOTT, D. J., HOUGH,
1159 D. W. & DANSON, M. J. 2007. The 2-oxoacid dehydrogenase multi-enzyme
1160 complex of the archaeon *Thermoplasma acidophilum* - recombinant
1161 expression, assembly and characterization. *FEBS J*, 274, 5406-15.

1162 HEYER, R., DORR, M., JELLEN-RITTER, A., SPATH, B., BABSKI, J., JASCHINSKI,
1163 K., SOPPA, J. & MARCHFELDER, A. 2012. High throughput sequencing
1164 reveals a plethora of small RNAs including tRNA derived fragments in
1165 *Haloferax volcanii*. *RNA biology*, 9, 1011-8.

1166 HILDEBRANDT, P., MATYSIK, J., SCHRADER, B., SCHARF, B. & ENGELHARD,
1167 M. 1994. Raman spectroscopic study of the blue copper protein halocyanin
1168 from *Natronobacterium pharaonis*. *Biochemistry*, 33, 11426-31.

1169 HO, Y. Y., HUANG, Y. H. & HUANG, C. Y. 2013. Chemical rescue of the post-
1170 translationally carboxylated lysine mutant of allantoinase and dihydroorotase
1171 by metal ions and short-chain carboxylic acids. *Amino Acids*, 44, 1181-91.

1172 HOCHULI, M., PATZELT, H., OESTERHELT, D., WUTHRICH, K. & SZYPERSKI, T.
1173 1999. Amino acid biosynthesis in the halophilic archaeon *Haloarcula*
1174 *hispanica*. *J Bacteriol*, 181, 3226-37.

1175 HORIE, A., TOMITA, T., SAIKI, A., KONO, H., TAKA, H., MINEKI, R., FUJIMURA, T.,
1176 NISHIYAMA, C., KUZUYAMA, T. & NISHIYAMA, M. 2009. Discovery of
1177 proteinaceous N-modification in lysine biosynthesis of *Thermus thermophilus*.
1178 *Nat Chem Biol*, 5, 673-9.

1179 HOWELL, D. M., HARICH, K., XU, H. M. & WHITE, R. H. 1998. Alpha-keto acid
1180 chain elongation reactions involved in the biosynthesis of coenzyme b (7-
1181 mercaptoheptanoyl threonine phosphate) in methanogenic archaea.
1182 *Biochemistry*, 37, 10108-17.

1183 HOWELL, D. M., XU, H. & WHITE, R. H. 1999. (R)-citramalate synthase in
1184 methanogenic archaea. *J Bacteriol*, 181, 331-3.

1185 HUANG, M., OPPERMANN-SANIO, F. B. & STEINBUCHER, A. 1999. Biochemical
1186 and molecular characterization of the *Bacillus subtilis* acetoin catabolic
1187 pathway. *J Bacteriol*, 181, 3837-41.

1188 HUNTER, S., APWEILER, R., ATTWOOD, T. K., BAIROCH, A., BATEMAN, A.,
1189 BINNS, D., BORK, P., DAS, U., DAUGHERTY, L., DUQUENNE, L., FINN, R.
1190 D., GOUGH, J., HAFT, D., HULO, N., KAHN, D., KELLY, E., LAUGRAUD, A.,
1191 LETUNIC, I., LONSDALE, D., LOPEZ, R., MADERA, M., MASLEN, J.,
1192 MCANULLA, C., MCDOWALL, J., MISTRY, J., MITCHELL, A., MULDER, N.,
1193 NATALE, D., ORENGO, C., QUINN, A. F., SELENGUT, J. D., SIGRIST, C. J.,
1194 THIMMA, M., THOMAS, P. D., VALENTIN, F., WILSON, D., WU, C. H. &
1195 YEATS, C. 2009. InterPro: the integrative protein signature database. *Nucleic
1196 Acids Res*, 37, D211-5.

1197 HWANG, S., CORDOVA, B., ABDO, M., PFEIFFER, F. & MAUPIN-FURLOW, J. A.
1198 2017. ThiN as a versatile domain of transcriptional repressors and catalytic
1199 enzymes of thiamine biosynthesis. *J Bacteriol*, 199, e00810-16.

1200 ISHIKAWA, R., ISHIDO, Y., TACHIKAWA, A., KAWASAKI, H., MATSUZAWA, H. &
1201 WAKAGI, T. 2002. *Aeropyrum pernix* K1, a strictly aerobic and
1202 hyperthermophilic archaeon, has two terminal oxidases, cytochrome ba3 and
1203 cytochrome aa3. *Arch Microbiol*, 179, 42-9.

1204 ISSALY, I. M. & ISSALY, A. S. 1974. Control of ornithine carbamoyltransferase
1205 activity by arginase in *Bacillus subtilis*. *Eur J Biochem*, 49, 485-95.

1206 ISUPOV, M. N., BOYKO, K. M., SUTTER, J. M., JAMES, P., SAYER, C., SCHMIDT,
1207 M., SCHONHEIT, P., NIKOLAEVA, A. Y., STEKHANOVA, T. N.,
1208 MARDANOV, A. V., RAVIN, N. V., BEZSUDNOVA, E. Y., POPOV, V. O. &
1209 LITTLECHILD, J. A. 2019. Thermostable branched-chain amino acid
1210 transaminases from the archaea *Geoglobus acetivorans* and *Archaeoglobus
1211 fulgidus*: biochemical and structural characterization. *Front Bioeng Biotechnol*,
1212 7, 7.

1213 JAIN, S., CAFORIO, A., FODRAN, P., LOLKEMA, J. S., MINNAARD, A. J. &
1214 DRIESSEN, A. J. M. 2014. Identification of CDP-archaeol synthase, a missing
1215 link of ether lipid biosynthesis in Archaea. *Chem Biol*, 21, 1392-401.

1216 JENSEN, P. E., GIBSON, L. C., HENNINGSEN, K. W. & HUNTER, C. N. 1996.
1217 Expression of the *chlI*, *chlD*, and *chlH* genes from the Cyanobacterium
1218 *Synechocystis* PCC6803 in *Escherichia coli* and demonstration that the three
1219 cognate proteins are required for magnesium-protoporphyrin chelatase
1220 activity. *J Biol Chem*, 271, 16662-7.

1221 JENSEN, P. E., GIBSON, L. C. & HUNTER, C. N. 1998. Determinants of catalytic
1222 activity with the use of purified I, D and H subunits of the magnesium
1223 protoporphyrin IX chelatase from *Synechocystis* PCC6803. *Biochem J*, 334 (Pt 2), 335-44.
1224

1225 JIANG, M., CHEN, M., GUO, Z. F. & GUO, Z. 2010. A bicarbonate cofactor
1226 modulates 1,4-dihydroxy-2-naphthoyl-coenzyme A synthase in menaquinone
1227 biosynthesis of *Escherichia coli*. *J Biol Chem*, 285, 30159-69.

1228 JOHNSEN, U., DAMBECK, M., ZAISS, H., FUHRER, T., SOPPA, J., SAUER, U. &
1229 SCHONHEIT, P. 2009. D-xylose degradation pathway in the halophilic
1230 archaeon *Haloferax volcanii*. *J Biol Chem*, 284, 27290-303.

1231 JOHNSEN, U., SUTTER, J. M., SCHULZ, A. C., TASTENSEN, J. B. & SCHONHEIT, P. 2015. XacR - a novel transcriptional regulator of D-xylose and L-arabinose
1232 catabolism in the haloarchaeon *Haloferax volcanii*. *Environ Microbiol*, 17, 1663-76.
1233
1234

1235 JOHNSON, C. L., PECHONICK, E., PARK, S. D., HAVEMANN, G. D., LEAL, N. A. &
1236 BOBIK, T. A. 2001. Functional genomic, biochemical, and genetic
1237 characterization of the *Salmonella pduO* gene, an ATP:cob(I)alamin
1238 adenosyltransferase gene. *J Bacteriol*, 183, 1577-84.

1239 JOHNSON, E. F. & MUKHOPADHYAY, B. 2005. A new type of sulfite reductase, a
1240 novel coenzyme F420-dependent enzyme, from the methanarchaeon
1241 *Methanocaldococcus jannaschii*. *J Biol Chem*, 280, 38776-86.

1242 JOHNSON, M., ZARETSKAYA, I., RAYTSELIS, Y., MERZHUH, Y., MCGINNIS, S. & MADDEN, T. L. 2008. NCBI BLAST: a better web interface. *Nucleic Acids Res*, 36, W5-9.
1243
1244

1245 JOLLEY, K. A., MADDOCKS, D. G., GYLES, S. L., MULLAN, Z., TANG, S. L., DYALL-SMITH, M. L., HOUGH, D. W. & DANSON, M. J. 2000. 2-Oxoacid
1246 dehydrogenase multienzyme complexes in the halophilic Archaea? Gene
1247 sequences and protein structural predictions. *Microbiology (Reading)*, 146 (Pt 5), 1061-69.
1248
1249

1250 KAILA, V. R. I. & WIKSTROM, M. 2021. Architecture of bacterial respiratory chains. *Nat Rev Microbiol*, 19, 319-30.
1251

1252 KAJIWARA, Y., SANTANDER, P. J., ROESSNER, C. A., PEREZ, L. M. & SCOTT, A. I. 2006. Genetically engineered synthesis and structural characterization of
1253 cobalt-precorrin 5A and -5B, two new intermediates on the anaerobic pathway
1254 to vitamin B12: definition of the roles of the CbiF and CbiG enzymes. *J Am Chem Soc*, 128, 9971-8.
1255
1256

1257 KAMINSKAS, E., KIMHI, Y. & MAGASANIK, B. 1970. Urocanase and N-formimino-L-
1258 glutamate formiminohydrolase of *Bacillus subtilis*, two enzymes of the
1259 histidine degradation pathway. *J Biol Chem*, 245, 3536-44.

1260 KAMINSKI, L. & EICHLER, J. 2014. *Haloferax volcanii* N-glycosylation: delineating
1261 the pathway of dTDP-rhamnose biosynthesis. *PLoS One*, 9, e97441.

1262 KANDIBA, L., LIN, C. W., AEBI, M., EICHLER, J. & GUERARDEL, Y. 2016.
1263 Structural characterization of the N-linked pentasaccharide decorating
1264 glycoproteins of the halophilic archaeon *Haloferax volcanii*. *Glycobiology*, 26,
1265 745-56.

1266 KANEHISA, M., SATO, Y., FURUMICHI, M., MORISHIMA, K. & TANABE, M. 2019.
1267 New approach for understanding genome variations in KEGG. *Nucleic Acids*
1268 *Res*, 47, D590-D95.

1269 KANG, B. S. & KIM, Y. M. 1999. Cloning and molecular characterization of the genes
1270 for carbon monoxide dehydrogenase and localization of molybdopterin, flavin
1271 adenine dinucleotide, and iron-sulfur centers in the enzyme of
1272 *Hydrogenophaga pseudoflava*. *J Bacteriol*, 181, 5581-90.

1273 KARATZA, P. & FRILLINGOS, S. 2005. Cloning and functional characterization of
1274 two bacterial members of the NAT/NCS2 family in *Escherichia coli*. *Mol*
1275 *Membr Biol*, 22, 251-61.

1276 KARDINAH, S., SCHMIDT, C. L., HANSEN, T., ANEMULLER, S., PETERSEN, A. &
1277 SCHAFFER, G. 1999. The strict molybdate-dependence of glucose-
1278 degradation by the thermoacidophile *Sulfolobus acidocaldarius* reveals the
1279 first crenarchaeotic molybdenum containing enzyme - an aldehyde
1280 oxidoreductase. *European Journal of Biochemistry*, 260, 540-48.

1281 KATES, M. 1993. Biology of halophilic bacteria, Part II. Membrane lipids of extreme
1282 halophiles: biosynthesis, function and evolutionary significance. *Experientia*,
1283 49, 1027-36.

1284 KATES, M., MOLDOVEANU, N. & STEWART, L. C. 1993. On the revised structure
1285 of the major phospholipid of *Halobacterium salinarum*. *Biochim Biophys Acta*,
1286 1169, 46-53.

1287 KATO, C., KURIHARA, T., KOBASHI, N., YAMANE, H. & NISHIYAMA, M. 2004.
1288 Conversion of feedback regulation in aspartate kinase by domain exchange.
1289 *Biochem Biophys Res Commun*, 316, 802-8.

1290 KAUR, N., GAUTAM, A., KUMAR, S., SINGH, A., SINGH, N., SHARMA, S.,
1291 SHARMA, R., TEWARI, R. & SINGH, T. P. 2011. Biochemical studies and
1292 crystal structure determination of dihydrodipicolinate synthase from
1293 *Pseudomonas aeruginosa*. *Int J Biol Macromol*, 48, 779-87.

1294 KAWAI, S., MORI, S., MUKAI, T., SUZUKI, S., YAMADA, T., HASHIMOTO, W. &
1295 MURATA, K. 2000. Inorganic Polyphosphate/ATP-NAD kinase of
1296 *Micrococcus flavus* and *Mycobacterium tuberculosis* H37Rv. *Biochem*
1297 *Biophys Res Commun*, 276, 57-63.

1298 KEIGHTLEY, J. A., ZIMMERMANN, B. H., MATHER, M. W., SPRINGER, P.,
1299 PASTUSZYN, A., LAWRENCE, D. M. & FEE, J. A. 1995. Molecular genetic
1300 and protein chemical characterization of the cytochrome ba3 from *Thermus*
1301 *thermophilus* HB8. *J Biol Chem*, 270, 20345-58.

1302 KELLERMANN, M. Y., YOSHINAGA, M. Y., VALENTINE, R. C., WORMER, L. &
1303 VALENTINE, D. L. 2016. Important roles for membrane lipids in haloarchaeal
1304 bioenergetics. *Biochim Biophys Acta*, 1858, 2940-56.

1305 KERSCHER, L. & OESTERHELT, D. 1976. A ferredoxin from halobacteria. *FEBS*
1306 *Letters*, 67, 320-22.

1307 KERSCHER, L. & OESTERHELT, D. 1977. Ferredoxin is the coenzyme of α -
1308 ketoacid oxidoreductases in *Halobacterium halobium*. *FEBS Letters*, 83, 197-
1309 201.

1310 KERSCHER, L. & OESTERHELT, D. 1981a. The catalytic mechanism of 2-
1311 oxoacid:ferredoxin oxidoreductases from *Halobacterium halobium*. One-
1312 electron transfer at two distinct steps of the catalytic cycle. *Eur J Biochem*,
1313 116, 595-600.

1314 KERSCHER, L. & OESTERHELT, D. 1981b. Purification and properties of two 2-
1315 oxoacid:ferredoxin oxidoreductases from *Halobacterium halobium*. *Eur J*
1316 *Biochem*, 116, 587-94.

1317 KERSCHER, L., OESTERHELT, D., CAMMACK, R. & HALL, D. O. 1976. A new
1318 plant-type ferredoxin from halobacteria. *Eur J Biochem*, 71, 101-7.

1319 KILJUNEN, S., PAJUNEN, M. I., DILKS, K., STORF, S., POHLSCHRODER, M. &
1320 SAVILAHTI, H. 2014. Generation of comprehensive transposon insertion
1321 mutant library for the model archaeon, *Haloferax volcanii*, and its use for gene
1322 discovery. *BMC Biol*, 12, 103.

1323 KIM, K., PARK, J. & RHEE, S. 2007. Structural and functional basis for (S)-allantoin
1324 formation in the ureide pathway. *J Biol Chem*, 282, 23457-64.

1325 KLEIN, A. R., BERK, H., PURWANTINI, E., DANIELS, L. & THAUER, R. K. 1996. Si-
1326 face stereospecificity at C5 of coenzyme F420 for F420-dependent glucose-6-
1327 phosphate dehydrogenase from *Mycobacterium smegmatis* and F420-
1328 dependent alcohol dehydrogenase from *Methanoculleus thermophilicus*. *Eur*
1329 *J Biochem*, 239, 93-7.

1330 KLEINE, T., LOCKHART, P. & BATSCHAUER, A. 2003. An *Arabidopsis* protein
1331 closely related to *Synechocystis* cryptochrome is targeted to organelles. *Plant*
1332 *J*, 35, 93-103.

1333 KLETZIN, A., HEIMERL, T., FLECHSLER, J., VAN NIFTRIK, L., RACHEL, R. &
1334 KLINGL, A. 2015. Cytochromes c in Archaea: distribution, maturation, cell
1335 architecture, and the special case of *Ignicoccus hospitalis*. *Front Microbiol*, 6,
1336 439.

1337 KOIKE-TAKESHITA, A., KOYAMA, T. & OGURA, K. 1997. Identification of a novel
1338 gene cluster participating in menaquinone (vitamin K2) biosynthesis. Cloning
1339 and sequence determination of the 2-heptaprenyl-1,4-naphthoquinone
1340 methyltransferase gene of *Bacillus stearothermophilus*. *J Biol Chem*, 272,
1341 12380-3.

1342 KRIVENTSEVA, E. V., KUZNETSOV, D., TEGENFELDT, F., MANNI, M., DIAS, R.,
1343 SIMAO, F. A. & ZDOBNOV, E. M. 2019. OrthoDB v10: sampling the diversity
1344 of animal, plant, fungal, protist, bacterial and viral genomes for evolutionary
1345 and functional annotations of orthologs. *Nucleic Acids Res*, 47, D807-D11.

1346 KUBATOVA, N., JONKER, H. R. A., SAXENA, K., RICHTER, C., VOGEL, V.,
1347 SCHREIBER, S., MARCHFELDER, A. & SCHWALBE, H. 2020. Solution
1348 Structure and Dynamics of the Small Protein HVO_2922 from *Haloferax*
1349 *volcanii*. *Chembiochem*, 21, 149-56.

1350 KUHNER, M., HAUFSCILD, K., NEUMANN, A., STORBECK, S., STREIF, J. &
1351 LAYER, G. 2014. The alternative route to heme in the methanogenic
1352 archaeon *Methanosarcina barkeri*. *Archaea*, 2014, Article ID 327637.

1353 KUNOW, J., SCHWOERER, B., STETTER, K. O. & THAUER, R. K. 1993. A F420-
1354 dependent NADP reductase in the extremely thermophilic sulfate-reducing
1355 *Archaeoglobus fulgidus*. *Archives of Microbiology*, 160, 199-205.

1356 KUPRAT, T., JOHNSEN, U., ORTJOHANN, M. & SCHONHEIT, P. 2020. Acetate
1357 metabolism in Archaea: characterization of an acetate transporter and of
1358 enzymes involved in acetate activation and gluconeogenesis in *Haloferax*
1359 *volcanii*. *Front Microbiol*, 11, 604926.

1360 KUPRAT, T., ORTJOHANN, M., JOHNSEN, U. & SCHONHEIT, P. 2021. Glucose
1361 metabolism and acetate switch in Archaea: the enzymes in *Haloferax volcanii*.
1362 *J Bacteriol*, 203, e00690-20.

1363 KUSHWAHA, S. C., KRAMER, J. K. G. & KATES, M. 1975. Isolation and
1364 characterization of C50-carotenoid pigments and other polar isoprenoids from
1365 *Halobacterium cutirubrum*. *Biochimica et Biophysica Acta (BBA) - Lipids and*
1366 *Lipid Metabolism*, 398, 303-14.

1367 LARGE, A., STAMME, C., LANGE, C., DUAN, Z., ALLERS, T., SOPPA, J. & LUND,
1368 P. A. 2007. Characterization of a tightly controlled promoter of the halophilic
1369 archaeon *Haloferax volcanii* and its use in the analysis of the essential *cct1*
1370 gene. *Mol Microbiol*, 66, 1092-106.

1371 LEE, T.-X., METZGER, S. U., CHO, Y. S., WHITMARSH, J. & KALLAS, T. 2001.
1372 Modification of inhibitor binding sites in the cytochrome bf complex by
1373 directed mutagenesis of cytochrome b6 in *Synechococcus* sp. PCC 7002.
1374 *Biochimica et Biophysica Acta (BBA) - Bioenergetics*, 1504, 235-47.

1375 LEE, Y., LEE, D. H., KHO, C. W., LEE, A. Y., JANG, M., CHO, S., LEE, C. H., LEE,
1376 J. S., MYUNG, P. K., PARK, B. C. & PARK, S. G. 2005. Transthyretin-related
1377 proteins function to facilitate the hydrolysis of 5-hydroxyisourate, the end
1378 product of the uricase reaction. *FEBS Lett*, 579, 4769-74.

1379 LEFFERS, H., GROPP, F., LOTTSPEICH, F., ZILLIG, W. & GARRETT, R. A. 1989.
1380 Sequence, organization, transcription and evolution of RNA polymerase
1381 subunit genes from the archaebacterial extreme halophiles *Halobacterium*
1382 *halobium* and *Halococcus morrhuae*. *J Mol Biol*, 206, 1-17.

1383 LEIF, H., SLED, V. D., OHNISHI, T., WEISS, H. & FRIEDRICH, T. 1995. Isolation
1384 and characterization of the proton-translocating NADH: ubiquinone
1385 oxidoreductase from *Escherichia coli*. *Eur J Biochem*, 230, 538-48.

1386 LEIGH, J. A., ALBERS, S. V., ATOMI, H. & ALLERS, T. 2011. Model organisms for
1387 genetics in the domain Archaea: methanogens, halophiles, *Thermococcales*
1388 and *Sulfolobales*. *FEMS Microbiol Rev*, 35, 577-608.

1389 LEMKE, C., YEUNG, M. & HOWELL, P. L. 1999. Expression, purification,
1390 crystallization and preliminary X-ray analysis of *Escherichia coli*
1391 argininosuccinate synthetase. *Acta Crystallogr D Biol Crystallogr*, 55, 2028-
1392 30.

1393 LI, H., GRAUPNER, M., XU, H. & WHITE, R. H. 2003. CofE catalyzes the addition of
1394 two glutamates to F420-0 in F420 coenzyme biosynthesis in *Methanococcus*
1395 *jannaschii*. *Biochemistry*, 42, 9771-8.

1396 LI, Z., RODRIGUEZ-FRANCO, M., ALBERS, S. V. & QUAX, T. E. F. 2020. The
1397 switch complex ArlCDE connects the chemotaxis system and the archaeellum.
1398 *Mol Microbiol*, 114, 468-79.

1399 LIAO, Y., ITHURBIDE, S., EVENHUIS, C., LOWE, J. & DUGGIN, I. G. 2021. Cell
1400 division in the archaeon *Haloferax volcanii* relies on two FtsZ proteins with
1401 distinct functions in division ring assembly and constriction. *Nat Microbiol*, 6,
1402 594-605.

1403 LIN, X. L. & WHITE, R. H. 1986. Occurrence of coenzyme F420 and its gamma-
1404 monoglutamyl derivative in nonmethanogenic archaeobacteria. *J Bacteriol*,
1405 168, 444-8.

1406 LIN, Y. K., MYHRMAN, R., SCHRAG, M. L. & GELB, M. H. 1988. Bacterial N-
1407 succinyl-L-diaminopimelic acid desuccinylase. Purification, partial
1408 characterization, and substrate specificity. *J Biol Chem*, 263, 1622-7.

1409 LIU, J., KRULWICH, T. A. & HICKS, D. B. 2008. Purification of two putative type II
1410 NADH dehydrogenases with different substrate specificities from alkaliphilic
1411 *Bacillus pseudofirmus* OF4. *Biochim Biophys Acta*, 1777, 453-61.

1412 LOC'H, J., BLAUD, M., RETY, S., LEBARON, S., DESCHAMPS, P., BAREILLE, J.,
1413 JOMBART, J., ROBERT-PAGANIN, J., DELBOS, L., CHARDON, F., ZHANG,
1414 E., CHARENTON, C., TOLLERVEY, D. & LEULLIOT, N. 2014. RNA mimicry
1415 by the fap7 adenylate kinase in ribosome biogenesis. *PLoS Biol*, 12,
1416 e1001860.

1417 MADEN, B. E. 2000. Tetrahydrofolate and tetrahydromethanopterin compared:
1418 functionally distinct carriers in C1 metabolism. *Biochem J*, 350 Pt 3, 609-29.

1419 MADON, J. & ZILLIG, W. 1983. A form of the DNA-dependent RNA polymerase of
1420 *Halobacterium halobium*, containing an additional component, is able to
1421 transcribe native DNA. *Eur J Biochem*, 133, 471-4.

1422 MAIER, L. K., STACHLER, A. E., BRENDDEL, J., STOLL, B., FISCHER, S., HAAS, K.
1423 A., SCHWARZ, T. S., ALKHNBASHI, O. S., SHARMA, K., URLAUB, H.,
1424 BACKOFEN, R., GOPHNA, U. & MARCHFELDER, A. 2019. The nuts and
1425 bolts of the Haloferax CRISPR-Cas system I-B. *RNA Biol*, 16, 469-80.

1426 MARTINEZ-RODRIGUEZ, S., GARCIA-PINO, A., LAS HERAS-VAZQUEZ, F. J.,
1427 CLEMENTE-JIMENEZ, J. M., RODRIGUEZ-VICO, F., GARCIA-RUIZ, J. M.,
1428 LORIS, R. & GAVIRA, J. A. 2012. Mutational and structural analysis of L-N-
1429 carbamoylase reveals new insights into a peptidase M20/M25/M40 family
1430 member. *J Bacteriol*, 194, 5759-68.

1431 MASHHADI, Z., XU, H., GROCHOWSKI, L. L. & WHITE, R. H. 2010. Archaeal RibL:
1432 a new FAD synthetase that is air sensitive. *Biochemistry*, 49, 8748-55.

1433 MASHHADI, Z., XU, H. & WHITE, R. H. 2009. An Fe²⁺-dependent cyclic
1434 phosphodiesterase catalyzes the hydrolysis of 7,8-dihydro-D-neopterin 2',3'-
1435 cyclic phosphate in methanopterin biosynthesis. *Biochemistry*, 48, 9384-92.

1436 MATTAR, S. 1996. *Molekularbiologische und Biochemische Charakterisierung*
1437 *zweier Komplexe der Atmungskette von Natronobacterium pharaonis*. Ph.D.,
1438 Ruhr-Universität Bochum.

1439 MATTAR, S. & ENGELHARD, M. 1997. Cytochrome ba₃ from *Natronobacterium*
1440 *pharaonis*--an archaeal four-subunit cytochrome-c-type oxidase. *Eur J*
1441 *Biochem*, 250, 332-41.

1442 MATTAR, S., SCHARF, B., KENT, S. B., RODEWALD, K., OESTERHELT, D. &
1443 ENGELHARD, M. 1994. The primary structure of halocyanin, an archaeal
1444 blue copper protein, predicts a lipid anchor for membrane fixation. *Journal of*
1445 *Biological Chemistry*, 269, 14939-45.

1446 MAURER, S., LUDT, K. & SOPPA, J. 2018. Characterization of copy number control
1447 of two *Haloferax volcanii* replication origins using deletion mutants and
1448 haloarchaeal artificial chromosomes. *J Bacteriol*, 200.

1449 MCCREADY, S. & MARCELLO, L. 2003. Repair of UV damage in *Halobacterium*
1450 *salinarum*. *Biochem Soc Trans*, 31, 694-8.

1451 MILLER, M. J. & GENNIS, R. B. 1983. The purification and characterization of the
1452 cytochrome d terminal oxidase complex of the *Escherichia coli* aerobic
1453 respiratory chain. *J Biol Chem*, 258, 9159-65.

1454 MIYAZAKI, J., KOBASHI, N., NISHIYAMA, M. & YAMANE, H. 2001. Functional and
1455 evolutionary relationship between arginine biosynthesis and prokaryotic lysine
1456 biosynthesis through alpha-aminoadipate. *J Bacteriol*, 183, 5067-73.

1457 MOON, S., KIM, J., KOO, J. & BAE, E. 2019. Structural and mutational analyses of
1458 psychrophilic and mesophilic adenylate kinases highlight the role of
1459 hydrophobic interactions in protein thermal stability. *Struct Dyn*, 6, 024702.

1460 MOORE, S. J., LAWRENCE, A. D., BIEDENDIECK, R., DEERY, E., FRANK, S.,
1461 HOWARD, M. J., RIGBY, S. E. & WARREN, M. J. 2013. Elucidation of the
1462 anaerobic pathway for the corrin component of cobalamin (vitamin B12). *Proc*
1463 *Natl Acad Sci U S A*, 110, 14906-11.

1464 MORII, H., KIYONARI, S., ISHINO, Y. & KOGA, Y. 2009. A novel biosynthetic
1465 pathway of archaetidyl-myo-inositol via archaetidyl-myo-inositol phosphate
1466 from CDP-archaeol and D-glucose 6-phosphate in methanoarchaeon
1467 *Methanothermobacter thermautotrophicus* cells. *J Biol Chem*, 284, 30766-74.

1468 MORII, H. & KOGA, Y. 2003. CDP-2,3-Di-O-geranylgeranyl-sn-glycerol:L-serine O-
1469 archaetidyltransferase (archaetidylserine synthase) in the methanogenic
1470 archaeon *Methanothermobacter thermautotrophicus*. *J Bacteriol*, 185, 1181-9.

1471 MOSHIRI, F., CHAWLA, A. & MAIER, R. J. 1991. Cloning, characterization, and
1472 expression in *Escherichia coli* of the genes encoding the cytochrome d
1473 oxidase complex from *Azotobacter vinelandii*. *J Bacteriol*, 173, 6230-41.

1474 NAGEL, C., MACHULLA, A., ZAHN, S. & SOPPA, J. 2019. Several one-domain zinc
1475 finger micro-proteins of *Haloferax volcanii* are important for stress adaptation,
1476 biofilm formation, and swarming. *Genes (Basel)*, 10, 361.

1477 NAM, S. E., KIM, A. C. & PAETZEL, M. 2012. Crystal structure of *Bacillus subtilis*
1478 signal peptide peptidase A. *J Mol Biol*, 419, 347-58.

1479 NANBA, T. & MUKOHATA, Y. 1987. A membrane-bound ATPase from
1480 *Halobacterium halobium*: purification and characterization. *J Biochem*
1481 (Tokyo), 102, 591-8.

1482 NEELON, K., ROBERTS, M. F. & STEC, B. 2011. Crystal structure of a trapped
1483 catalytic intermediate suggests that forced atomic proximity drives the
1484 catalysis of mIPS. *Biophys J*, 101, 2816-24.

1485 NEWTON, W. A., MORINO, Y. & SNELL, E. E. 1965. Properties of crystalline
1486 tryptophanase. *J Biol Chem*, 240, 1211-8.

1487 NG, S. Y., CHABAN, B., VANDYKE, D. J. & JARRELL, K. F. 2007. Archaeal signal
1488 peptidases. *Microbiology (Reading)*, 153, 305-14.

1489 NOCEK, B., EVDOKIMOVA, E., PROUDFOOT, M., KUDRITSKA, M.,
1490 GROCHOWSKI, L. L., WHITE, R. H., SAVCHENKO, A., YAKUNIN, A. F.,
1491 EDWARDS, A. & JOACHIMIAK, A. 2007. Structure of an amide bond forming
1492 F(420):gamma-glutamyl ligase from *Archaeoglobus fulgidus* -- a member of a
1493 new family of non-ribosomal peptide synthases. *J Mol Biol*, 372, 456-69.

1494 NUSSBAUM, P., ITHURBIDE, S., WALSH, J. C., PATRO, M., DELPECH, F.,
1495 RODRIGUEZ-FRANCO, M., CURMI, P. M. G., DUGGIN, I. G., QUAX, T. E.
1496 F. & ALBERS, S. V. 2020. An oscillating MinD protein determines the cellular
1497 positioning of the motility machinery in Archaea. *Curr Biol*, 30, 4956-72 e4.

1498 O'TOOLE, G. A., TRZEBIATOWSKI, J. R. & ESCALANTE-SEMERENA, J. C. 1994.
1499 The cobC gene of *Salmonella typhimurium* codes for a novel phosphatase
1500 involved in the assembly of the nucleotide loop of cobalamin. *Journal of*
1501 *Biological Chemistry*, 269, 26503-11.

1502 OBERTO, J. 2013. SyntTax: a web server linking synteny to prokaryotic taxonomy.
1503 *BMC Bioinformatics*, 14, 4.

1504 ODA, M., SUGISHITA, A. & FURUKAWA, K. 1988. Cloning and nucleotide
1505 sequences of histidase and regulatory genes in the *Bacillus subtilis hut*
1506 operon and positive regulation of the operon. *J Bacteriol*, 170, 3199-205.

1507 OHNISHI, H., KITAMURA, H., MINOWA, T., SAKAI, H. & OHTA, T. 1992. Molecular
1508 cloning of a glucoamylase gene from a thermophilic *Clostridium* and kinetics
1509 of the cloned enzyme. *Eur J Biochem*, 207, 413-8.

1510 OPPERMAN, F. B., SCHMIDT, B. & STEINBUHEL, A. 1991. Purification and
1511 characterization of acetoin:2,6-dichlorophenolindophenol oxidoreductase,
1512 dihydrolipoamide dehydrogenase, and dihydrolipoamide acetyltransferase of
1513 the *Pelobacter carbinolicus* acetoin dehydrogenase enzyme system. *J*
1514 *Bacteriol*, 173, 757-67.

1515 OREN, A. 2002. Molecular ecology of extremely halophilic Archaea and Bacteria.
1516 *FEMS Microbiol. Ecol.*, 39, 1-7.

1517 OREN, A., VENTOSA, A. & GRANT, W. D. 1997. Proposed minimal standards for
1518 description of new taxa in the order *Halobacteriales*. *International Journal of*
1519 *Systematic Bacteriology*, 47, 233-38.

1520 OTAKA, E., HIGO, K. & ITOH, T. 1984. Yeast ribosomal proteins. VIII. Isolation of
1521 two proteins and sequence characterization of twenty-four proteins from
1522 cytoplasmic ribosomes. *Mol Gen Genet*, 195, 544-6.

1523 OUCHI, T., TOMITA, T., HORIE, A., YOSHIDA, A., TAKAHASHI, K., NISHIDA, H.,
1524 LASSAK, K., TAKA, H., MINEKI, R., FUJIMURA, T., KOSONO, S.,
1525 NISHIYAMA, C., MASUI, R., KURAMITSU, S., ALBERS, S. V., KUZUYAMA,
1526 T. & NISHIYAMA, M. 2013. Lysine and arginine biosyntheses mediated by a
1527 common carrier protein in *Sulfolobus*. *Nat Chem Biol*, 9, 277-83.

1528 PALMER, D. R., GARRETT, J. B., SHARMA, V., MEGANATHAN, R., BABBITT, P.
1529 C. & GERLT, J. A. 1999. Unexpected divergence of enzyme function and
1530 sequence: "N-acylamino acid racemase" is o-succinylbenzoate synthase.
1531 *Biochemistry*, 38, 4252-8.

1532 PAN, X., CAO, D., XIE, F., XU, F., SU, X., MI, H., ZHANG, X. & LI, M. 2020.
1533 Structural basis for electron transport mechanism of complex I-like
1534 photosynthetic NAD(P)H dehydrogenase. *Nat Commun*, 11, 610.

1535 PENNINGTON, J. M., KEMP, M., MCGARRY, L., CHEN, Y. & STROUPE, M. E.
1536 2020. Siroheme synthase orients substrates for dehydrogenase and
1537 chelatase activities in a common active site. *Nat Commun*, 11, 864.

1538 PEREZ-ARNAIZ, P., DATTANI, A., SMITH, V. & ALLERS, T. 2020. *Haloferax*
1539 *volcanii*-a model archaeon for studying DNA replication and repair. *Open Biol*,
1540 10, 200293.

1541 PETROVA, T. E., BOYKO, K. M., NIKOLAEVA, A. Y., STEKHANOVA, T. N.,
1542 GRUZDEV, E. V., MARDANOV, A. V., STROILOV, V. S., LITTLECHILD, J.
1543 A., POPOV, V. O. & BEZSUDNOVA, E. Y. 2018. Structural characterization
1544 of geranylgeranyl pyrophosphate synthase GACE1337 from the
1545 hyperthermophilic archaeon *Geoglobus acetivorans*. *Extremophiles*, 22, 877-
1546 88.

1547 PFEIFFER, F., BROICHER, A., GILLICH, T., KLEE, K., MEJIA, J., RAMPP, M. &
1548 OESTERHELT, D. 2008a. Genome information management and integrated
1549 data analysis with HaloLex. *Arch Microbiol*, 190, 281-99.

1550 PFEIFFER, F., LOSENSKY, G., MARCHFELDER, A., HABERMANN, B. & DYALL-
1551 SMITH, M. 2020. Whole-genome comparison between the type strain of
1552 *Halobacterium salinarum* (DSM 3754(T)) and the laboratory strains R1 and
1553 NRC-1. *Microbiologyopen*, 9, e974.

1554 PFEIFFER, F. & OESTERHELT, D. 2015. A manual curation strategy to improve
1555 genome annotation: application to a set of haloarchael genomes. *Life (Basel)*,
1556 5, 1427-44.

1557 PFEIFFER, F., SCHUSTER, S. C., BROICHER, A., FALB, M., PALM, P.,
1558 RODEWALD, K., RUEPP, A., SOPPA, J., TITTOR, J. & OESTERHELT, D.
1559 2008b. Evolution in the laboratory: the genome of *Halobacterium salinarum*
1560 strain R1 compared to that of strain NRC-1. *Genomics*, 91, 335-46.

1561 PFRIMER, P., DE MORAES, L. M., GALDINO, A. S., SALLES, L. P., REIS, V. C., DE
1562 MARCO, J. L., PRATES, M. V., BLOCH, C., JR. & TORRES, F. A. 2010.
1563 Cloning, purification, and partial characterization of *Bacillus subtilis* urate
1564 oxidase expressed in *Escherichia coli*. *J Biomed Biotechnol*, 2010, 674908.

1565 PHILLIPS, G., GROCHOWSKI, L. L., BONNETT, S., XU, H., BAILLY, M., BLABY-
1566 HAAS, C., EL YACOUBI, B., IWATA-REUYL, D., WHITE, R. H. & DE
1567 CRECY-LAGARD, V. 2012. Functional promiscuity of the COG0720 family.
1568 *ACS Chem Biol*, 7, 197-209.

1569 PHILLIPS, R. S. & GOLLNICK, P. D. 1989. Evidence that cysteine 298 is in the
1570 active site of tryptophan indole-lyase. *Journal of Biological Chemistry*, 264,
1571 10627-32.

1572 PHILMUS, B., DECAMPS, L., BERTEAU, O. & BEGLEY, T. P. 2015. Biosynthetic
1573 versatility and coordinated action of 5'-deoxyadenosyl radicals in deazaflavin
1574 biosynthesis. *J Am Chem Soc*, 137, 5406-13.

1575 PICKL, A., JOHNSEN, U. & SCHONHEIT, P. 2012. Fructose degradation in the
1576 haloarchaeon *Haloferax volcanii* involves a bacterial type
1577 phosphoenolpyruvate-dependent phosphotransferase system, fructose-1-
1578 phosphate kinase, and class II fructose-1,6-bisphosphate aldolase. *Journal of*
1579 *bacteriology*, 194, 3088-97.

1580 PLAGA, W., LOTTSPEICH, F. & OESTERHELT, D. 1992. Improved purification,
1581 crystallization and primary structure of pyruvate:ferredoxin oxidoreductase
1582 from *Halobacterium halobium*. *Eur J Biochem*, 205, 391-7.

1583 POHLSCHRODER, M. & ESQUIVEL, R. N. 2015. Archaeal type IV pili and their
1584 involvement in biofilm formation. *Front Microbiol*, 6, 190.

1585 PORAT, I., SIEPRAWKA-LUPA, M., TENG, Q., BOHANON, F. J., WHITE, R. H. &
1586 WHITMAN, W. B. 2006. Biochemical and genetic characterization of an early
1587 step in a novel pathway for the biosynthesis of aromatic amino acids and p-
1588 aminobenzoic acid in the archaeon *Methanococcus maripaludis*. *Mol*
1589 *Microbiol*, 62, 1117-31.

1590 PORAT, I., WATERS, B. W., TENG, Q. & WHITMAN, W. B. 2004. Two biosynthetic
1591 pathways for aromatic amino acids in the archaeon *Methanococcus*
1592 *maripaludis*. *J Bacteriol*, 186, 4940-50.

1593 PROMPONAS, V. J., ILIOPOULOS, I. & OUZOUNIS, C. A. 2015. Annotation
1594 inconsistencies beyond sequence similarity-based function prediction -
1595 phylogeny and genome structure. *Stand Genomic Sci*, 10, 108.

1596 PRUNETTI, L., REUTER, C. J., HEPOWIT, N. L., WU, Y., BARRUETO, L.,
1597 MIRANDA, H. V., KELLY, K. & MAUPIN-FURLOW, J. A. 2014. Structural and
1598 biochemical properties of an extreme 'salt-loving' proteasome activating
1599 nucleotidase from the archaeon *Haloferax volcanii*. *Extremophiles*, 18, 283-
1600 93.

1601 PURWANTINI, E. & MUKHOPADHYAY, B. 2013. Rv0132c of *Mycobacterium*
1602 *tuberculosis* encodes a coenzyme F420-dependent hydroxymycolic acid
1603 dehydrogenase. *PLoS One*, 8, e81985.

1604 QI, Q., ITO, Y., YOSHIMATSU, K. & FUJIWARA, T. 2016. Transcriptional regulation
1605 of dimethyl sulfoxide respiration in a haloarchaeon, *Haloferax volcanii*.
1606 *Extremophiles*, 20, 27-36.

1607 QU, Q., LEE, S. J. & BOOS, W. 2004. Molecular and biochemical characterization of
1608 a fructose-6-phosphate-forming and ATP-dependent fructokinase of the
1609 hyperthermophilic archaeon *Thermococcus litoralis*. *Extremophiles*, 8, 301-8.

1610 QUAX, T. E. F., ALTEGOER, F., ROSSI, F., LI, Z., RODRIGUEZ-FRANCO, M.,
1611 KRAUS, F., BANGE, G. & ALBERS, S. V. 2018. Structure and function of the
1612 archaeal response regulator CheY. *Proc Natl Acad Sci U S A*, 115, E1259-
1613 E68.

1614 RAFFAELLI, N., EMANUELLI, M., PISANI, F. M., AMICI, A., LORENZI, T.,
1615 RUGGIERI, S. & MAGNI, G. 1999. Identification of the archaeal NMN
1616 adenyltransferase gene. *Molecular & Cellular Biochemistry*, 193, 99-102.

1617 RAFFAELLI, N., PISANI, F. M., LORENZI, T., EMANUELLI, M., AMICI, A.,
1618 RUGGIERI, S. & MAGNI, G. 1997. Characterization of nicotinamide

1619 mononucleotide adenylyltransferase from thermophilic archaea. *J Bacteriol*,
1620 179, 7718-23.

1621 RANGASWAMY, V. & ALTEKAR, W. 1994. Ketohexokinase (ATP:D-fructose 1-
1622 phosphotransferase) from a halophilic archaeobacterium, *Haloarcula*
1623 *vallismortis*: purification and properties. *J Bacteriol*, 176, 5505-12.

1624 RAUT, P., GLASS, J. B. & LIEBERMAN, R. L. 2021. Archaeal roots of
1625 intramembrane aspartyl protease siblings signal peptide peptidase and
1626 presenilin. *Proteins*, 89, 232-41.

1627 RAUX, E., LEECH, H. K., BECK, R., SCHUBERT, H. L., SANTANDER, P. J.,
1628 ROESSNER, C. A., SCOTT, A. I., MARTENS, J. H., JAHN, D., THERMES,
1629 C., RAMBACH, A. & WARREN, M. J. 2003. Identification and functional
1630 analysis of enzymes required for precorrin-2 dehydrogenation and metal ion
1631 insertion in the biosynthesis of sirohaem and cobalamin in *Bacillus*
1632 *megaterium*. *Biochem J*, 370, 505-16.

1633 RAWLS, K. S., YACOVONE, S. K. & MAUPIN-FURLOW, J. A. 2010. GlpR represses
1634 fructose and glucose metabolic enzymes at the level of transcription in the
1635 haloarchaeon *Haloferax volcanii*. *J Bacteriol*, 192, 6251-60.

1636 REDDY, S. G., SACCHETTINI, J. C. & BLANCHARD, J. S. 1995. Expression,
1637 purification, and characterization of *Escherichia coli* dihydrodipicolinate
1638 reductase. *Biochemistry*, 34, 3492-501.

1639 REINHARDT, A., JOHNSEN, U. & SCHONHEIT, P. 2019. L-Rhamnose catabolism in
1640 archaea. *Mol Microbiol*, 111, 1093-108.

1641 REN, H., WANG, L., BENNETT, M., LIANG, Y., ZHENG, X., LU, F., LI, L., NAN, J.,
1642 LUO, M., ERIKSSON, S., ZHANG, C. & SU, X. D. 2005. The crystal structure
1643 of human adenylylase kinase 6: An adenylylase kinase localized to the cell
1644 nucleus. *Proc Natl Acad Sci U S A*, 102, 303-8.

1645 REUTER, C. J. & MAUPIN-FURLOW, J. A. 2004. Analysis of proteasome-dependent
1646 proteolysis in *Haloferax volcanii* cells, using short-lived green fluorescent
1647 proteins. *Appl. Environ. Microbiol.*, 70, 7530-38.

1648 REUTER, C. J., UTHANDI, S., PUENTES, J. A. & MAUPIN-FURLOW, J. A. 2010.
1649 Hydrophobic carboxy-terminal residues dramatically reduce protein levels in
1650 the haloarchaeon *Haloferax volcanii*. *Microbiology (Reading)*, 156, 248-55.

1651 RICH, P. R. & MARECHAL, A. 2010. The mitochondrial respiratory chain. *Essays*
1652 *Biochem*, 47, 1-23.

1653 RIVLIN, A. A., CHAN, Y. L. & WOOL, I. G. 1999. The contribution of a zinc finger
1654 motif to the function of yeast ribosomal protein YL37a. *J Mol Biol*, 294, 909-
1655 19.

1656 RODIONOV, D. A., VITRESCHAK, A. G., MIRONOV, A. A. & GELFAND, M. S. 2003.
1657 Comparative genomics of the vitamin B12 metabolism and regulation in
1658 prokaryotes. *J Biol Chem*, 278, 41148-59.

1659 RODIONOVA, I. A., VETTING, M. W., LI, X., ALMO, S. C., OSTERMAN, A. L. &
1660 RODIONOV, D. A. 2017. A novel bifunctional transcriptional regulator of
1661 riboflavin metabolism in Archaea. *Nucleic Acids Res*, 45, 3785-99.

1662 ROESSNER, C. A., WARREN, M. J., SANTANDER, P. J., ATSHAVES, B. P.,
1663 OZAKI, S.-I., STOLOWICH, N. J., IIDA, K. & SCOTT, A. I. 1992. Expression
1664 of 9 *Salmonella typhimurium* enzymes for cobinamide synthesis. Identification
1665 of the 11-methyl and 20-methyl transferases of corrin biosynthesis. *FEBS*
1666 *Letters*, 301, 73-78.

1667 ROMISCH-MARGL, W., EISENREICH, W., HAASE, I., BACHER, A. & FISCHER, M.
1668 2008. 2,5-diamino-6-ribitylamino-4(3H)-pyrimidinone 5'-phosphate synthases
1669 of fungi and archaea. *FEBS J*, 275, 4403-14.

1670 RUDOLPH, B., HANSEN, T. & SCHONHEIT, P. 2004. Glucose-6-phosphate
1671 isomerase from the hyperthermophilic archaeon *Methanococcus jannaschii*:
1672 characterization of the first archaeal member of the phosphoglucose
1673 isomerase superfamily. *Arch Microbiol*, 181, 82-7.

1674 RUEPP, A., MULLER, H. N., LOTTSPEICH, F. & SOPPA, J. 1995. Catabolic
1675 ornithine transcarbamylase of halobacterium halobium (salinarium) -
1676 purification, characterization, sequence determination, and evolution. *Journal*
1677 *of Bacteriology*, 177, 1129-36.

1678 SANTANDER, P. J., KAJIWARA, Y., WILLIAMS, H. J. & SCOTT, A. I. 2006.
1679 Structural characterization of novel cobalt corrinoids synthesized by enzymes
1680 of the vitamin B12 anaerobic pathway. *Bioorg Med Chem*, 14, 724-31.

1681 SANTANDER, P. J., ROESSNER, C. A., STOLOWICH, N. J., HOLDERMAN, M. T. &
1682 SCOTT, A. I. 1997. How corrinoids are synthesized without oxygen: nature's
1683 first pathway to vitamin B12. *Chemistry & Biology*, 4, 659-66.

1684 SATO, S., NAKADA, Y., KANAYA, S. & TANAKA, T. 1988. Molecular cloning and
1685 nucleotide sequence of *Thermus thermophilus* HB8 *trpE* and *trpG*. *Biochimica*
1686 *et Biophysica Acta (BBA) - Gene Structure and Expression*, 950, 303-12.

1687 SCHADT, H. S., SCHADT, S., OLDACH, F. & SUSSMUTH, R. D. 2009. 2-Amino-2-
1688 deoxyisochorismate is a key intermediate in *Bacillus subtilis* p-aminobenzoic
1689 acid biosynthesis. *J Am Chem Soc*, 131, 3481-3.

1690 SCHAFER, G., ENGELHARD, M. & MULLER, V. 1999. Bioenergetics of the
1691 Archaea. *Microbiol Mol Biol Rev*, 63, 570-620.

1692 SCHARF, B. & ENGELHARD, M. 1993. Halocyanin, an archaebacterial blue copper
1693 protein (type I) from *Natronobacterium pharaonis*. *Biochemistry*, 32, 12894-
1694 900.

1695 SCHARF, B., WITTENBERG, R. & ENGELHARD, M. 1997. Electron transfer proteins
1696 from the haloalkaliphilic archaeon *Natronobacterium pharaonis*: possible
1697 components of the respiratory chain include cytochrome bc and a terminal
1698 oxidase cytochrome ba3. *Biochemistry*, 36, 4471-9.

1699 SCHILLER, H., SCHULZE, S., MUTAN, Z., DE VAULX, C., RUNCIE, C.,
1700 SCHWARTZ, J., RADOS, T., BISSON FILHO, A. W. & POHLSCHRODER, M.
1701 2020. *Haloferax volcanii* immersed liquid biofilms develop independently of
1702 known biofilm machineries and exhibit rapid honeycomb pattern formation.
1703 *mSphere*, 5, e00976-20.

1704 SCHMIDT, D. M., HUBBARD, B. K. & GERLT, J. A. 2001. Evolution of enzymatic
1705 activities in the enolase superfamily: functional assignment of unknown
1706 proteins in *Bacillus subtilis* and *Escherichia coli* as L-Ala-D/L-Glu epimerases.
1707 *Biochemistry*, 40, 15707-15.

1708 SCHNOES, A. M., BROWN, S. D., DODEVSKI, I. & BABBITT, P. C. 2009.
1709 Annotation error in public databases: misannotation of molecular function in
1710 enzyme superfamilies. *PLoS Comput Biol*, 5, e1000605.

1711 SCHOLZEN, T. & ARNDT, E. 1991. Organization and nucleotide sequence of ten
1712 ribosomal protein genes from the region equivalent to the spectinomycin
1713 operon in the archaebacterium *Halobacterium marismortui*. *Mol Gen Genet*,
1714 228, 70-80.

1715 SCHUBERT, H. L., ROSE, R. S., LEECH, H. K., BRINDLEY, A. A., HILL, C. P.,
1716 RIGBY, S. E. & WARREN, M. J. 2008. Structure and function of SirC from
1717 *Bacillus megaterium*: a metal-binding precorrin-2 dehydrogenase. *Biochem J*,
1718 415, 257-63.

1719 SCHULLER, J. M., BIRRELL, J. A., TANAKA, H., KONUMA, T., WULFHORST, H.,
1720 COX, N., SCHULLER, S. K., THIEMANN, J., LUBITZ, W., SETIF, P.,
1721 IKEGAMI, T., ENGEL, B. D., KURISU, G. & NOWACZYK, M. M. 2019.
1722 Structural adaptations of photosynthetic complex I enable ferredoxin-
1723 dependent electron transfer. *Science*, 363, 257-60.

1724 SCHULTZ, A. C., NYGAARD, P. & SAXILD, H. H. 2001. Functional analysis of 14
1725 genes that constitute the purine catabolic pathway in *Bacillus subtilis* and
1726 evidence for a novel regulon controlled by the PucR transcription activator. *J*
1727 *Bacteriol*, 183, 3293-302.

1728 SCHULZE, S., ADAMS, Z., CERLETTI, M., DE CASTRO, R., FERREIRA-CERCA,
1729 S., FUFUZAN, C., GIMENEZ, M. I., HIPPLER, M., JEVTIC, Z., KNUPPEL, R.,
1730 LEGERME, G., LENZ, C., MARCHFELDER, A., MAUPIN-FURLOW, J.,
1731 PAGGI, R. A., PFEIFFER, F., POETSCH, A., URLAUB, H. &
1732 POHLSCHRODER, M. 2020. The Archaeal Proteome Project advances
1733 knowledge about archaeal cell biology through comprehensive proteomics.
1734 *Nat Commun*, 11, 3145.

1735 SCHULZE, S., PFEIFFER, F., GARCIA, B. A. & POHLSCHRODER, M. 2021.
1736 Glycoproteomics of *Haloferax volcanii* reveals an extensive glycoproteome
1737 and concurrence of different N-glycosylation pathways. Available:
1738 [https://www.biorxiv.org/content/biorxiv/early/2021/01/21/2021.01.21.427637.f](https://www.biorxiv.org/content/biorxiv/early/2021/01/21/2021.01.21.427637.full.pdf)
1739 [ull.pdf](https://www.biorxiv.org/content/biorxiv/early/2021/01/21/2021.01.21.427637.full.pdf) [Accessed January 21, 2021].

1740 SCOTT, J. W. & RASCHE, M. E. 2002. Purification, overproduction, and partial
1741 characterization of beta-RFAP synthase, a key enzyme in the methanopterin
1742 biosynthesis pathway. *J Bacteriol*, 184, 4442-8.

1743 SELBY, C. P. & SANCAR, A. 2006. A cryptochrome/photolyase class of enzymes
1744 with single-stranded DNA-specific photolyase activity. *Proc Natl Acad Sci U S*
1745 *A*, 103, 17696-700.

1746 SELENGUT, J. D. & HAFT, D. H. 2010. Unexpected abundance of coenzyme
1747 F(420)-dependent enzymes in *Mycobacterium tuberculosis* and other
1748 actinobacteria. *J Bacteriol*, 192, 5788-98.

1749 SERINO, L., REIMMANN, C., BAUR, H., BEYELER, M., VISCA, P. & HAAS, D.
1750 1995. Structural genes for salicylate biosynthesis from chorismate in
1751 *Pseudomonas aeruginosa*. *Mol Gen Genet*, 249, 217-28.

1752 SHAHEEN, N., KOBAYASHI, K., TERAONO, H., FUKUSHIGE, T., HORIUCHI, M.
1753 & SAHEKI, T. 1994. Characterization of human wild-type and mutant
1754 argininosuccinate synthetase proteins expressed in bacterial cells. *Enzyme*
1755 *Protein*, 48, 251-64.

1756 SHALEV, Y., SOUCY, S. M., PAPKE, R. T., GOGARTEN, J. P., EICHLER, J. &
1757 GOPHNA, U. 2018. Comparative analysis of surface layer glycoproteins and
1758 genes involved in protein glycosylation in the genus *Haloferax*. *Genes*
1759 *(Basel)*, 9, 172.

1760 SHALEV, Y., TURGEMAN-GROTT, I., TAMIR, A., EICHLER, J. & GOPHNA, U.
1761 2017. Cell surface glycosylation is required for efficient mating of *Haloferax*
1762 *volcanii*. *Front Microbiol*, 8, 1253.

1763 SHIMA, S., WARKENTIN, E., GRABARSE, W., SORDEL, M., WICKE, M., THAUER,
1764 R. K. & ERMLER, U. 2000. Structure of coenzyme F(420) dependent

1765 methylenetetrahydromethanopterin reductase from two methanogenic
1766 archaea. *J Mol Biol*, 300, 935-50.

1767 SHIMIZU, T., TOMITA, T., KUZUYAMA, T. & NISHIYAMA, M. 2016. Crystal structure
1768 of the LysY.LysW complex from *Thermus thermophilus*. *J Biol Chem*, 291,
1769 9948-59.

1770 SIDDARAMAPPA, S., CHALLACOMBE, J. F., DECASTRO, R. E., PFEIFFER, F.,
1771 SASTRE, D. E., GIMENEZ, M. I., PAGGI, R. A., DETTER, J. C.,
1772 DAVENPORT, K. W., GOODWIN, L. A., KYRPIDES, N., TAPIA, R.,
1773 PITLUCK, S., LUCAS, S., WOYKE, T. & MAUPIN-FURLOW, J. A. 2012. A
1774 comparative genomics perspective on the genetic content of the alkaliphilic
1775 haloarchaeon *Natrialba magadii* ATCC 43099^T. *BMC Genomics*, 13, 165.

1776 SIMMS, S. A., VOIGE, W. H. & GILVARG, C. 1984. Purification and characterization
1777 of succinyl-CoA: tetrahydrodipicolinate N-succinyltransferase from
1778 *Escherichia coli*. *J Biol Chem*, 259, 2734-41.

1779 SISIGNANO, M., MORBITZER, D., GATGENS, J., OLDIGES, M. & SOPPA, J. 2010.
1780 A 2-oxoacid dehydrogenase complex of *Haloferax volcanii* is essential for
1781 growth on isoleucine but not on other branched-chain amino acids.
1782 *Microbiology*, 156, 521-9.

1783 SLOCK, J., STAHLY, D. P., HAN, C. Y., SIX, E. W. & CRAWFORD, I. P. 1990. An
1784 apparent *Bacillus subtilis* folic acid biosynthetic operon containing *pab*, an
1785 amphibolic *trpG* gene, a third gene required for synthesis of para-
1786 aminobenzoic acid, and the dihydropteroate synthase gene. *J Bacteriol*, 172,
1787 7211-26.

1788 SOPPA, J. 2011. Functional genomic and advanced genetic studies reveal novel
1789 insights into the metabolism, regulation, and biology of *Haloferax volcanii*.
1790 *Archaea*, 2011, 602408.

1791 SPATH, B., SCHUBERT, S., LIEBEROTH, A., SETTELE, F., SCHUTZ, S.,
1792 FISCHER, S. & MARCHFELDER, A. 2008. Two archaeal tRNase Z enzymes:
1793 similar but different. *Arch Microbiol*, 190, 301-8.

1794 SREERAMULU, K. 2003. Purification and partial characterization of cytochrome c552
1795 from *Halobacterium salinarum*. *Indian Journal of Biochemistry and*
1796 *Biophysics*, 40, 274-77.

1797 SREERAMULU, K., SCHMIDT, C. L., SCHAFER, G. & ANEMULLER, S. 1998.
1798 Studies of the electron transport chain of the euryarchaeon *Halobacterium*
1799 *salinarum*: indications for a type II NADH dehydrogenase and a complex III
1800 analog. *J Bioenerg Biomembr*, 30, 443-53.

1801 STEINERT, K., WAGNER, V., KROTH PANCIC, P. G. & BICKEL SANDKOETTER,
1802 S. 1997. Characterization and subunit structure of the ATP synthase of the
1803 halophilic archaeon *Haloferax volcanii* and organization of the ATP synthase
1804 genes. *Journal of Biological Chemistry*, 272, 6261-69.

1805 STORBECK, S., ROLFES, S., RAUX-DEERY, E., WARREN, M. J., JAHN, D. &
1806 LAYER, G. 2010. A novel pathway for the biosynthesis of heme in Archaea:
1807 genome-based bioinformatic predictions and experimental evidence.
1808 *Archaea*, 2010, 175050.

1809 STORF, S., PFEIFFER, F., DILKS, K., CHEN, Z. Q., IMAM, S. & POHLSCHRODER,
1810 M. 2010. Mutational and bioinformatic analysis of haloarchaeal lipobox-
1811 containing proteins. *Archaea*, 2010, Article ID 410975.

1812 STRAUB, J., BRENNEIS, M., JELLEN-RITTER, A., HEYER, R., SOPPA, J. &
1813 MARCHFELDER, A. 2009. Small RNAs in haloarchaea: Identification,
1814 differential expression and biological function. *RNA Biol*, 6.

1815 STROUPE, M. E., LEECH, H. K., DANIELS, D. S., WARREN, M. J. & GETZOFF, E.
1816 D. 2003. CysG structure reveals tetrapyrrole-binding features and novel
1817 regulation of siroheme biosynthesis. *Nat Struct Biol*, 10, 1064-73.

1818 SU, X., LIN, Z., CHEN, W., JIANG, H., ZHANG, S. & LIN, H. 2012. Chemogenomic
1819 approach identified yeast YLR143W as diphthamide synthetase. *Proc Natl*
1820 *Acad Sci U S A*, 109, 19983-7.

1821 SUTTER, J. M., JOHNSEN, U., REINHARDT, A. & SCHONHEIT, P. 2020. Pentose
1822 degradation in archaea: *Halorhabdus* species degrade D-xylose, L-arabinose
1823 and D-ribose via bacterial-type pathways. *Extremophiles*, 24, 759-72.

1824 SUTTER, J. M., TASTENSEN, J. B., JOHNSEN, U., SOPPA, J. & SCHONHEIT, P.
1825 2016. Key enzymes of the semiphosphorylative Entner-Doudoroff pathway in
1826 the haloarchaeon *Haloferax volcanii*: Characterization of glucose
1827 dehydrogenase, gluconate dehydratase and 2-keto-3-deoxy-6-
1828 phosphogluconate aldolase. *J Bacteriol*, 198, 2251-62.

1829 SUVARNA, K., STEVENSON, D., MEGANATHAN, R. & HUDSPETH, M. E. 1998.
1830 Menaquinone (vitamin K2) biosynthesis: localization and characterization of
1831 the menA gene from *Escherichia coli*. *J Bacteriol*, 180, 2782-7.

1832 TACHIBANA, A. 1994. A novel prenyltransferase, farnesylgeranyl diphosphate
1833 synthase, from the haloalkaliphilic archaeon, *Natronobacterium pharaonis*.
1834 *FEBS Letters*, 341, 291-94.

1835 TACHIBANA, A., YANO, Y., OTANI, S., NOMURA, N., SAKO, Y. & TANIGUCHI, M.
1836 2000. Novel prenyltransferase gene encoding farnesylgeranyl diphosphate
1837 synthase from a hyperthermophilic archaeon, *Aeropyrum pernix* - Molecular

1838 evolution with alteration in product specificity. *European Journal of*
1839 *Biochemistry*, 267, 321-28.

1840 TAKAO, M., KOBAYASHI, T., OIKAWA, A. & YASUI, A. 1989. Tandem arrangement
1841 of photolyase and superoxide dismutase genes in *Halobacterium halobium*. *J*
1842 *Bacteriol*, 171, 6323-9.

1843 TANAKA, M., OGAWA, N., IHARA, K., SUGIYAMA, Y. & MUKOHATA, Y. 2002.
1844 Cytochrome aa(3) in *Haloferax volcanii*. *J Bacteriol*, 184, 840-5.

1845 TÄSTENSEN, J. B., JOHNSEN, U., REINHARDT, A., ORTJOHANN, M. &
1846 SCHONHEIT, P. 2020. D-galactose catabolism in archaea: operation of the
1847 DeLey-Doudoroff pathway in *Haloferax volcanii*. *FEMS Microbiol Lett*, 367,
1848 fnaa029.

1849 TE BRÖMMELSTROET, B. W., HENSGENS, C. M., KELTJENS, J. T., VAN DER
1850 DRIFT, C. & VOGELS, G. D. 1990. Purification and properties of 5,10-
1851 methylenetetrahydromethanopterin reductase, a coenzyme F420-dependent
1852 enzyme, from *Methanobacterium thermoautotrophicum* strain delta H. *Journal*
1853 *of Biological Chemistry*, 265, 1852-57.

1854 TITTES, C., SCHWARZER, S., PFEIFFER, F., DYALL-SMITH, M., RODRIGUEZ-
1855 FRANCO, M., OKSANEN, H. M. & QUAX, T. E. F. 2021. Cellular and
1856 genomic properties of *Haloferax gibbonsii* LR2-5, the host of euryarchaeal
1857 virus HFTV1. *Front Microbiol*, 12, 625599.

1858 TRIPEPI, M., YOU, J., TEMEL, S., ONDER, O., BRISSON, D. & POHLSCHRODER,
1859 M. 2012. N-Glycosylation of *Haloferax volcanii* flagellins requires known Agl
1860 proteins and Is essential for biosynthesis of stable flagella. *Journal of*
1861 *bacteriology*, 194, 4876-87.

1862 TRZEBIATOWSKI, J. R., O'TOOLE, G. A. & ESCALANTE-SEMERENA, J. C. 1994.
1863 The cobT gene of *Salmonella typhimurium* encodes the NaMN: 5,6-
1864 dimethylbenzimidazole phosphoribosyltransferase responsible for the
1865 synthesis of N1-(5-phospho-alpha-D-ribosyl)-5,6-dimethylbenzimidazole, an
1866 intermediate in the synthesis of the nucleotide loop of cobalamin. *J Bacteriol*,
1867 176, 3568-75.

1868 TURKOWYD, B., SCHREIBER, S., WORTZ, J., SEGAL, E. S., MEVARECH, M.,
1869 DUGGIN, I. G., MARCHFELDER, A. & ENDESFELDER, U. 2020.
1870 Establishing live-cell single-molecule localization microscopy imaging and
1871 single-particle tracking in the archaeon *Haloferax volcanii*. *Front Microbiol*, 11,
1872 583010.

1873 UNIPROT, C. 2021. UniProt: the universal protein knowledgebase in 2021. *Nucleic*
1874 *Acids Res*, 49, D480-D89.

1875 UTHMAN, S., BAR, C., SCHEIDT, V., LIU, S., TEN HAVE, S., GIORGINI, F.,
1876 STARK, M. J. & SCHAFFRATH, R. 2013. The amidation step of diphthamide
1877 biosynthesis in yeast requires DPH6, a gene identified through mining the
1878 DPH1-DPH5 interaction network. *PLoS Genet*, 9, e1003334.

1879 VAN OUYEN, J. & SOPPA, J. 2007. Three 2-oxoacid dehydrogenase operons in
1880 *Haloferax volcanii*: expression, deletion mutants and evolution. *Microbiology*,
1881 153, 3303-13.

1882 VANNICE, J. C., SKAFF, D. A., KEIGHTLEY, A., ADDO, J. K., WYCKOFF, G. J. &
1883 MIZIORKO, H. M. 2014. Identification in *Haloferax volcanii* of
1884 phosphomevalonate decarboxylase and isopentenyl phosphate kinase as
1885 catalysts of the terminal enzyme reactions in an archaeal alternate
1886 mevalonate pathway. *J Bacteriol*, 196, 1055-63.

1887 VAUPEL, M. & THAUER, R. K. 1995. Coenzyme F420-dependent N5,N10-
1888 methylenetetrahydromethanopterin reductase (Mer) from *Methanobacterium*
1889 *thermoautotrophicum* strain Marburg. Cloning, sequencing, transcriptional
1890 analysis, and functional expression in *Escherichia coli* of the *mer* gene. *Eur J*
1891 *Biochem*, 231, 773-8.

1892 VAUPEL, M., VORHOLT, J. A. & THAUER, R. K. 1998. Overproduction and one-step
1893 purification of the N5,N10-methenyltetrahydromethanopterin cyclohydrolase
1894 (Mch) from the hyperthermophilic *Methanopyrus kandleri*. *Extremophiles*, 2,
1895 15-22.

1896 VENCES-GUZMAN, M. A., GEIGER, O. & SOHLENKAMP, C. 2008. *Sinorhizobium*
1897 *meliloti* mutants deficient in phosphatidylserine decarboxylase accumulate
1898 phosphatidylserine and are strongly affected during symbiosis with alfalfa. *J*
1899 *Bacteriol*, 190, 6846-56.

1900 WALSH, J. C., ANGSTMANN, C. N., BISSON-FILHO, A. W., GARNER, E. C.,
1901 DUGGIN, I. G. & CURMI, P. M. G. 2019. Division plane placement in
1902 pleomorphic archaea is dynamically coupled to cell shape. *Mol Microbiol*.

1903 WANNER, C. & SOPPA, J. 2002. Functional role for a 2-oxo acid dehydrogenase in
1904 the halophilic archaeon *Haloferax volcanii*. *J Bacteriol*, 184, 3114-21.

1905 WERNER, A. K., MEDINA-ESCOBAR, N., ZULAWSKI, M., SPARKES, I. A., CAO, F.
1906 Q. & WITTE, C. P. 2013. The ureide-degrading reactions of purine ring
1907 catabolism employ three amidohydrolases and one aminohydrolase in
1908 *Arabidopsis*, soybean, and rice. *Plant Physiol*, 163, 672-81.

1909 WERNER, A. K., ROMEIS, T. & WITTE, C. P. 2010. Ureide catabolism in
1910 *Arabidopsis thaliana* and *Escherichia coli*. *Nat Chem Biol*, 6, 19-21.

1911 WESTENBERG, D. J., BRAUNE, A., RUPPERT, C., MULLER, V., HERZBERG, C.,
1912 GOTTSCHALK, G. & BLAUT, M. 1999. The F420H₂-dehydrogenase from
1913 *Methanobolbus tindarius*: cloning of the *ffd* operon and expression of the genes
1914 in *Escherichia coli*. *FEMS Microbiol Lett*, 170, 389-98.

1915 WHITE, P. J. & KELLY, B. 1965. Purification and properties of diaminopimelate
1916 decarboxylase from *Escherichia coli*. *Biochem J*, 96, 75-84.

1917 WHITE, R. H. 1988. Analysis and characterization of the folates in the
1918 nonmethanogenic archaeobacteria. *J Bacteriol*, 170, 4608-12.

1919 WHITE, R. H. 2004. L-Aspartate semialdehyde and a 6-deoxy-5-ketohexose 1-
1920 phosphate are the precursors to the aromatic amino acids in
1921 *Methanocaldococcus jannaschii*. *Biochemistry*, 43, 7618-27.

1922 WILLIAMS, T. J., ALLEN, M. A., LIAO, Y., RAFTERY, M. J. & CAVICCHIOLI, R.
1923 2019. Sucrose metabolism in haloarchaea: reassessment using genomics,
1924 proteomics, and metagenomics. *Appl Environ Microbiol*, 85, e02935-18.

1925 WISEMAN, J. S. & NICHOLS, J. S. 1984. Purification and properties of
1926 diaminopimelic acid epimerase from *Escherichia coli*. *J Biol Chem*, 259, 8907-
1927 14.

1928 WOODSON, J. D. & ESCALANTE-SEMERENA, J. C. 2004. CbiZ, an
1929 amidohydrolase enzyme required for salvaging the coenzyme B12 precursor
1930 cobinamide in archaea. *Proc Natl Acad Sci U S A*, 101, 3591-6.

1931 WOODSON, J. D., PECK, R. F., KREBS, M. P. & ESCALANTE-SEMERENA, J. C.
1932 2003a. The *cobY* gene of the archaeon *Halobacterium* sp. strain NRC-1 is
1933 required for de novo cobamide synthesis. *J Bacteriol*, 185, 311-6.

1934 WOODSON, J. D., ZAYAS, C. L. & ESCALANTE-SEMERENA, J. C. 2003b. A new
1935 pathway for salvaging the coenzyme B12 precursor cobinamide in archaea
1936 requires cobinamide-phosphate synthase (CbiB) enzyme activity. *J Bacteriol*,
1937 185, 7193-201.

1938 WYSS, L., WASER, M., GEBETSBERGER, J., ZYWICKI, M. & POLACEK, N. 2018.
1939 mRNA-specific translation regulation by a ribosome-associated ncRNA in
1940 *Haloferax volcanii*. *Sci Rep*, 8, 12502.

1941 XI, H., SCHNEIDER, B. L. & REITZER, L. 2000. Purine catabolism in *Escherichia coli*
1942 and function of xanthine dehydrogenase in purine salvage. *J Bacteriol*, 182,
1943 5332-41.

1944 XU, Z., JIANG, W. H., JIAO, R. S. & YANG, Y. L. 2002. [Cloning, sequencing and
1945 high expression in *Escherichia coli* of D-hydantoinase gene from *Burkholderia*
1946 *pickettii*]. *Sheng Wu Gong Cheng Xue Bao*, 18, 149-54.

1947 YANG, Y., YATSUNAMI, R., ANDO, A., MIYOKO, N., FUKUI, T., TAKAICHI, S. &
1948 NAKAMURA, S. 2015. Complete biosynthetic pathway of the C50 carotenoid
1949 bacterioruberin from lycopene in the extremely halophilic archaeon
1950 *Haloarcula japonica*. *J Bacteriol*, 197, 1614-23.

1951 YIN, J., XU, L. X., CHERNEY, M. M., RAUX-DEERY, E., BINDLEY, A. A.,
1952 SAVCHENKO, A., WALKER, J. R., CUFF, M. E., WARREN, M. J. & JAMES,
1953 M. N. 2006. Crystal structure of the vitamin B12 biosynthetic
1954 cobaltochelatase, CbiXS, from *Archaeoglobus fulgidus*. *J Struct Funct*
1955 *Genomics*, 7, 37-50.

1956 YIN, W., CAI, X., MA, H., ZHU, L., ZHANG, Y., CHOU, S. H., GALPERIN, M. Y. &
1957 HE, J. 2020. A decade of research on the second messenger c-di-AMP.
1958 *FEMS Microbiol Rev*, 44, 701-24.

1959 YOSHIDA, A., TOMITA, T., ATOMI, H., KUZUYAMA, T. & NISHIYAMA, M. 2016.
1960 Lysine biosynthesis of *Thermococcus kodakarensis* with the capacity to
1961 function as an ornithine biosynthetic system. *J Biol Chem*, 291, 21630-43.

1962 YOSHIDA, A., TOMITA, T., FUJIMURA, T., NISHIYAMA, C., KUZUYAMA, T. &
1963 NISHIYAMA, M. 2015. Structural insight into amino group-carrier protein-
1964 mediated lysine biosynthesis: crystal structure of the LysZ.LysW complex
1965 from *Thermus thermophilus*. *J Biol Chem*, 290, 435-47.

1966 YU, Y., LIANG, Y. H., BROSTROMER, E., QUAN, J. M., PANJIKAR, S., DONG, Y.
1967 H. & SU, X. D. 2006. A catalytic mechanism revealed by the crystal structures
1968 of the imidazolonepropionase from *Bacillus subtilis*. *J Biol Chem*, 281, 36929-
1969 36.

1970 ZAFRILLA, B., MARTINEZ-ESPINOSA, R. M., BONETE, M. J., BUTT, J. N.,
1971 RICHARDSON, D. J. & GATES, A. J. 2011. A haloarchaeal ferredoxin
1972 electron donor that plays an essential role in nitrate assimilation. *Biochem*
1973 *Soc Trans*, 39, 1844-8.

1974 ZAHN, S., KUBATOVA, N., PYPPE, D. J., CASSIDY, L., SAXENA, K., THOLEY, A.,
1975 SCHWALBE, H. & SOPPA, J. 2021. Biological functions, genetic and
1976 biochemical characterization, and NMR structure determination of the small
1977 zinc finger protein HVO_2753 from *Haloferax volcanii*. *FEBS J*, 288, 2042-62.

1978 ZAYAS, C. L. & ESCALANTE-SEMERENA, J. C. 2007. Reassessment of the late
1979 steps of coenzyme B12 synthesis in *Salmonella enterica*: evidence that
1980 dephosphorylation of adenosylcobalamin-5'-phosphate by the CobC
1981 phosphatase is the last step of the pathway. *J Bacteriol*, 189, 2210-8.

1982 ZERULLA, K., CHIMILESCHI, S., NATHER, D., GOPHNA, U., PAPKE, R. T. &
1983 SOPPA, J. 2014. DNA as a phosphate storage polymer and the alternative
1984 advantages of polyploidy for growth or survival. *PLoS One*, 9, e94819.
1985 ZHANG, F., SCHEERER, P., OBERPICHLER, I., LAMPARTER, T. & KRAUSS, N.
1986 2013. Crystal structure of a prokaryotic (6-4) photolyase with an Fe-S cluster
1987 and a 6,7-dimethyl-8-ribityllumazine antenna chromophore. *Proc Natl Acad*
1988 *Sci U S A*, 110, 7217-22.
1989 ZHANG, P., BATTCHIKOVA, N., PAAKKARINEN, V., KATOH, H., IWAI, M.,
1990 IKEUCHI, M., PAKRASI, H. B., OGAWA, T. & ARO, E. M. 2005. Isolation,
1991 subunit composition and interaction of the NDH-1 complexes from
1992 *Thermosynechococcus elongatus* BP-1. *Biochem J*, 390, 513-20.
1993 ZHU, X., DZIKOVSKI, B., SU, X., TORELLI, A. T., ZHANG, Y., EALICK, S. E.,
1994 FREED, J. H. & LIN, H. 2011. Mechanistic understanding of *Pyrococcus*
1995 *horikoshii* Dph2, a [4Fe-4S] enzyme required for diphthamide biosynthesis.
1996 *Mol Biosyst*, 7, 74-81.
1997 ZHU, X., KIM, J., SU, X. & LIN, H. 2010. Reconstitution of diphthine synthase activity
1998 in vitro. *Biochemistry*, 49, 9649-57.
1999 ZIMMERMANN, B. H., NITSCHKE, C. I., FEE, J. A., RUSNAK, F. & MUNCK, E. 1988.
2000 Properties of a copper-containing cytochrome ba3: a second terminal oxidase
2001 from the extreme thermophile *Thermus thermophilus*. *Proc Natl Acad Sci U S*
2002 *A*, 85, 5779-83.
2003