



Metabolic and evolutionary patterns in the extremely acidophilic archaeon *Ferroplasma acidiphilum* YT

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Supplementary Information to the Manuscript**Metabolic and evolutionary patterns in the extremely acidophilic archaeon *Ferroplasma acidiphilum* Y^T**

Olga V. Golyshina, Hai Tran, Oleg N. Reva, Sofia Lemak, Alexander F. Yakunin, Alexander Goesmann, Taras Y. Nechitaylo, Violetta LaCono, Francesco Smedile, Alexei Slesarev, David Rojo, Coral Barbas, Manuel Ferrer, Michail M. Yakimov and Peter N. Golyshin

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Supplementary Table S1. *F. acidiphilum* Y^T genes examined by real-time reverse-transcription PCR in this study.

Locus tag	Gene	Name	Forward primer / Reverse primer	TaqMan®
FAD_0161	<i>rpl2</i>	50S ribosomal protein L2	F: 5'-ATACAGGAGCCCGAGTCACA-3' R: 5'-TGTTGTGGTAGTGCCGTTGT-3'	5'-CACCGGGTAGAAATGCACCGTT-3'
FAD_0374	<i>gyrB</i>	DNA gyrase B	F: 5'-GCAGAGCTTCAAGGGGAGTT-3' R: 5'-CGCCAACAGGTAATGCAGTA-3'	5'-CTAAACGGTGCTGAAGTAATCA-3'
FAD_0567	<i>porA</i>	2-oxoacid--ferredoxin oxidoreductase, α subunit	F: 5'-CAACCGGGCTTCCAACAAAGA-3' R: 5'-GAGCACTATTCTTGGCATGT-3'	5'-AGATCTCAACCAGGTGCTTGGT-3'
FAD_0703	<i>morME</i>	malate oxidoreductase (malic enzyme)	F: 5'- ATGCCCAGGTGATGCAAAGA-3' R: 5'- TGCATAGGAAGCAGCAACCA-3'	5'- CCAATCAGATAAATAATAGCAT-3'
FAD_0712	<i>korA</i>	2-oxoacid--ferredoxin oxidoreductase, α subunit	F: 5'- AGCAGGGCGTATCAGCTAAC-3' R: 5'- AGGGTCATATGCCTGCCATT-3'	5'- AATGTTTCGAGCCTTTCCTGCA-3'
FAD_0713	<i>korB</i>	2-oxoacid--ferredoxin oxidoreductase, β subunit	F: 5'- AGATTGGTGCCTGGATGTG-3' R: 5'- AGCTGACAAAGCCTGTGTTA-3'	5'- TGACTTTGGTATAGTAAGTGCG-3'
FAD_0714	<i>sdhD</i>	Succinate dehydrogenase subunit D	F: 5'- AGCCCCATTGCCACTGAAA-3' R: 5'-TCAACCATAGTGCCGGAGAA-3'	5'- AATTAACAATGCTATGTACCT-3'
FAD_0717	<i>sdhA</i>	Succinate dehydrogenase subunit A	F: 5'- TTCAACGCGCTGGAAACAAG-3' R: 5'- TGAAGTGTGACCCCTTGT-3'	5'- ATATGCTATGGCTACTGGAGCA-3'
FAD_0718	<i>mdh</i>	Malate dehydrogenase	F: 5'- TTTGCACCCAGGACCCTAA-3' R: 5'- CTGCAATGCGTATGCCATTA-3'	5'- AACAGATTAAGAAATATTCACC-3'
FAD_1044	<i>pepc</i>	phosphoenol pyruvate carboxylase	F: 5'- AGAAGGGAGCGGAAATTGCA-3' R: 5'- GCACGTGGCAAACCTAGCTTT-3'	5'- CTATTTGGATACTCCCGTAGTA-3'

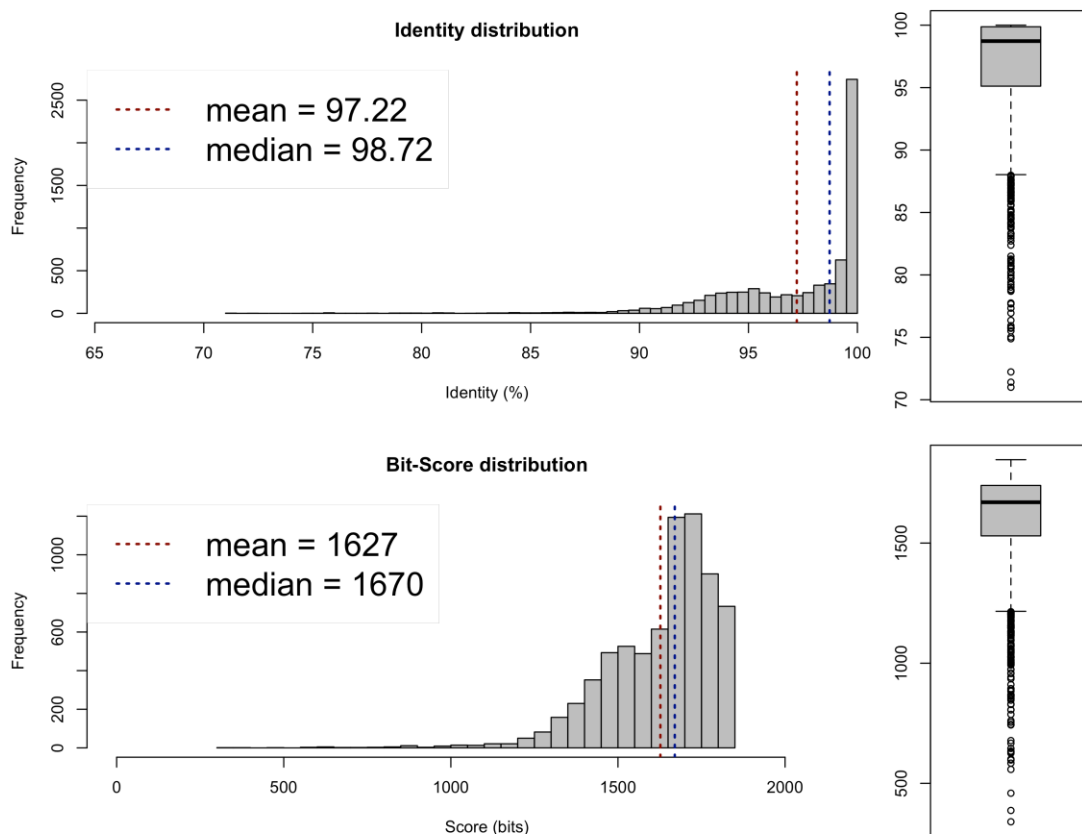
Supplementary Table S2. 116 single-nucleotide substitutions (separate Excel file).

Supplementary Table S3. Transporters in *F. acidiphilum* Y^T

<i>Sugar & polysaccharide transporters</i>	
FAD_1026	ABC-type sugar transporter, ATPase component
FAD_1027	ABC-type sugar transporter, permease component
FAD_1028	ABC-type sugar transporter, permease component
FAD_1029	ABC-type sugar transporter, extracellular component
FAD_1459	sugar ABC transporter 1, permease protein
FAD_1460	sugar ABC transporter 1, ATP binding protein
<i>AA transporters</i>	
FAD_0093	ABC-type peptide transporter, permease component
FAD_0094	oligopeptide ABC transporter Dpp2, permease protein
FAD_0126	6TMS neutral amino acid family transporter
FAD_0228	amino acid transporter
FAD_0377	amino acid transporter related protein
FAD_0539	cationic amino acid transporter
FAD_0636	amino acid transporter
FAD_0655	ABC transporter peptide-binding protein
FAD_0659	ABC transporter permease. dipeptide/oligopeptide/nickel transport system
FAD_0802	amino acid/polyamine/organocation superfamily transporter
FAD_0803	aspartate/glutamate family transporter
FAD_0804	aspartate/glutamate family transporter
FAD_0805	amino acid/polyamine/organocation superfamily transporter

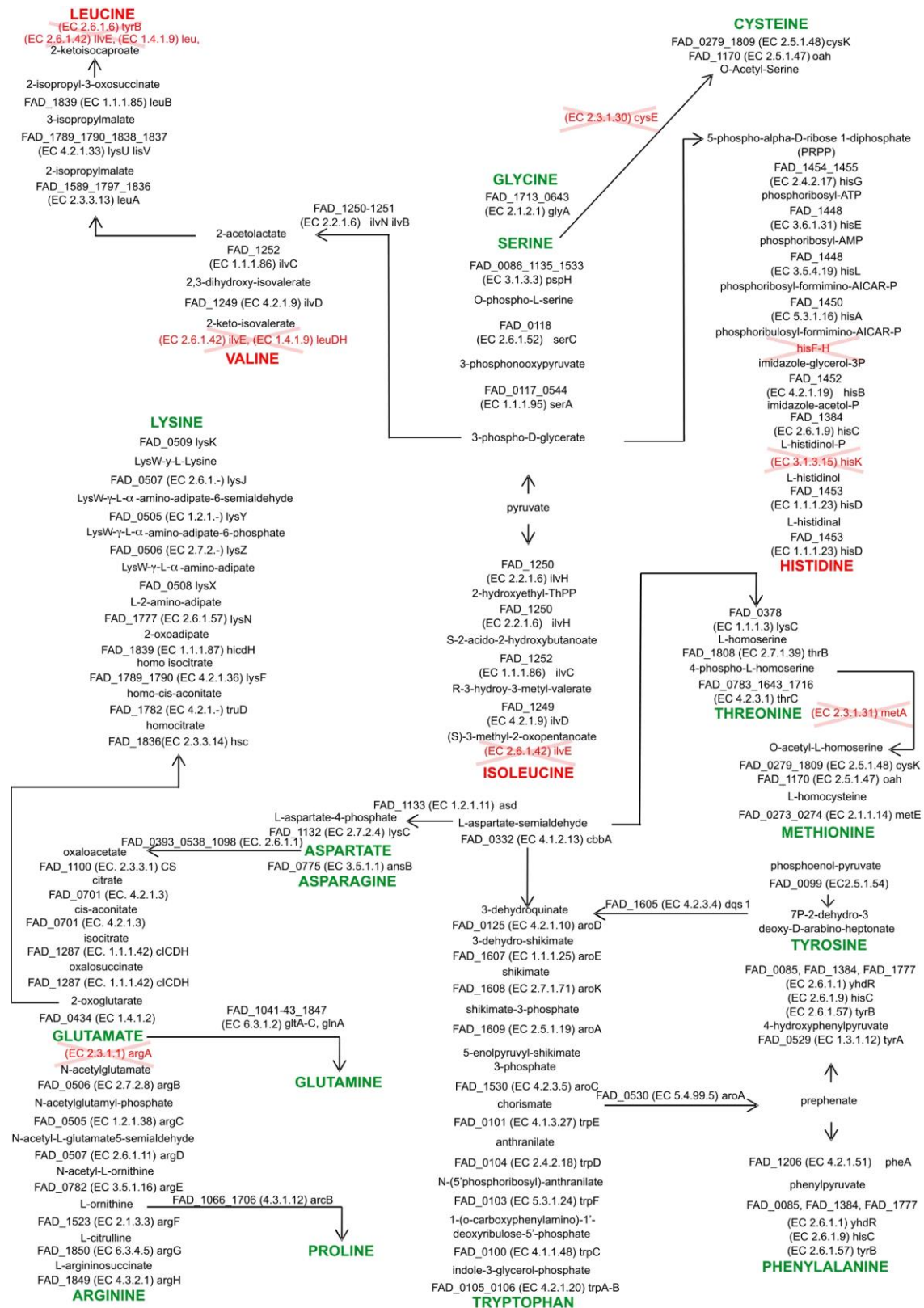
FAD_0853	ABC transporter peptide-binding protein
FAD_0914	amino acid/polyamine/organocation superfamily transporter
FAD_0915	amino acid/polyamine/organocation superfamily transporter
FAD_0919	ABC transporter substrate-binding protein (C-terminal fragment) peptide
FAD_1013	amino acid/polyamine/organocation superfamily transporter
FAD_1014	ABC-type peptide/opine/nickel family transporter ATPase
FAD_1015	ABC-type peptide/opine/nickel family transporter ATPase
FAD_1016	ABC-type peptide/opine/nickel family transporter
FAD_1017	ABC-type peptide/opine/nickel family transporter
FAD_1018	ABC-type peptide/opine/nickel family transporter
FAD_1069	amino acid transporter
FAD_1371	6TMS neutral amino acid family transporter
FAD_1571	amino acid transporter
FAD_1737	transporter probably aa
<p><i>Major facilitator superfamily MFS multidrug efflux pumps and major facilitator superfamily permeases</i></p>	
<p>FAD_0778_0801_0803_0815_0839_1106 FAD_1033_1118_1278_1296_1750_0006_0143_0213_0472_0497_0561_0684_0768 _0796_0822_0844_0849_0979_0994_0996_1000_1011_1018_1038_1040_1047_1048 _1129_1272_1349_1380_1407_1408_1445_1484_1504_1544_1555_1591 _1599_1720_1788_1791</p>	
<p><i>Other transporters</i></p>	
FAD_0049	stomatin family transporter
FAD_0249	cation diffusion facilitator family transporter, that increase tolerance to divalent metal ions such as cadmium, zinc, and cobalt

FAD_0250	cation diffusion facilitator family transporter, that increase tolerance to divalent metal ions such as cadmium, zinc, and cobalt
FAD_1045	ammonium transporter
FAD_1137	C4-dicarboxylate transporter
FAD_1177	ABC-2-type family permease
FAD_1178	ABC-2-type family permease
FAD_0071	VIT family Fe ²⁺ /Mn ²⁺ transporter
FAD_1261	VIT family Fe ²⁺ /Mn ²⁺ transporter
FAD_0016	transporter of sulfur-containing compounds
FAD_0264	transporter of sulfur-containing compounds
FAD_0848	transporter of sulfur-containing compounds
FAD_1341	transporter of sulfur-containing compounds
FAD_1460	nitrate/sulfonate/bicarbonate ABC transporter ATP-binding protein
FAD_1459	nitrate/sulfonate/bicarbonate ABC transporter ATP-binding protein
FAD_1458	nitrate/sulfonate/bicarbonate ABC transporter ATP-binding protein
FAD_1494	monovalent cation:proton antiporter-2 family transporter
FAD_1510	sodium-dependent phosphate transporter
FAD_1593	manganese/divalent cation transporter
FAD_1667	multidrug ABC transporter ATP-binding protein
FAD_1771	daunorubicin resistance ABC transporter
FAD_1800	Kef-type potassium transporter NAD-binding component



Supplementary Figure S1. Average Nucleotide Identity (ANI) analysis (<http://enve-omics.ce.gatech.edu/ani/>) of genomes “*F. acidarmanus* fer1” and *F. acidiphilum* Y^T suggests ANI values above 95 %, which is the accepted cut off for separation of two species based on the whole-genome comparisons, suggesting that on the basis of their genomic data, *F. acidiphilum* Y^T and “*F. acidarmanus*” fer1 belong to the same species.

1. Goris, J., Konstantinidis, K. T., Klappenbach, J. A., Coenye, T., Vandamme, P. & Tiedje, J. M. DNA-DNA hybridization values and their relationship to whole-genome sequence similarities. *Int J Syst Evol Microbiol.* **57**, 81-91 (2007).



Supplementary Figure S2. Overview of amino acid biosynthesis pathways in the *F. acidiphilum* Y^T genome. Missing genes and reaction are indicated by red colour and crossed out. All amino acids with incomplete synthesis pathways are indicated in red. EC numbers are shown in parentheses.

