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Towards reconstructing a metabolic tree of life

Marina Marcet-Houben¹, Pere Puigbò¹, Antoni Romeu¹ and Santiago Garcia-Vallve^{1, *}

¹Evolutionary Genomics Group, Biochemistry and Biotechnology Department, Rovira i Virgili University, Campus Sescelades, c/ Marcel li Domingo s/n, 43007 TARRAGONA, Spain; Santiago Garcia-Vallve* - E-mail: santi.garcia-vallve@urv.cat; * Corresponding author

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Abstract:

Using information from several metabolic databases, we have built our own metabolic database containing 434 pathways and 1157 different enzymes. We have used this information to construct a dendrogram that demonstrates the metabolic similarities between 282 species. The resulting species distribution and the clusters defined in the tree show a certain taxonomic congruence, especially in recent relationships between species. This dendrogram is another representation of the tree of life, based on metabolism that may complement the trees constructed by other methods. For example, the metabolic dissimilarity we demonstrate between Symbiobacterium thermophilum (previously defined as Actinobacteria) and the other Actinobacteria species, and the metabolic similarity between S. thermophilum and Clostridia, combined with other evidence, suggest that S. thermophilum may be re-classified as Firmicutes, Clostridia.

Keywords: metablic pathways; enzymes; dendogram; taxonomy; species

Background:

For many years phylogenetic trees have been used to study the evolution of organisms. Since Charles Darwin first described the evolution of species as a tree, scientist have attempted to create a tree that could represent a hierarchical classification of all known species based on their evolution and at the same time provide information about extinct species and the common ancestry shared by known species. When sequencing technologies were developed, the use of taxonomic marker molecules such as the small subunit ribosomal RNA seemed sufficient to draw consistent phylogenetic trees. Studies using genes or protein sequences led to a classification of microorganisms and recognised the Archaea as the third domain of life. [1]

When whole genome sequences of prokaryote organisms became available, everyone hoped that this extended information would help them to build more accurate phylogenies but it was then discovered that different genes produced different trees. It was at this point that doubts were raised as to whether a tree structure was the best representation of evolution. [2] Simultaneously, the discovery that horizontal gene transfer events (HGT) between species was more common than previously suspected [3, 4] put a strain on the search for the "true tree". [5] After all, the gene used in a phylogenetic study may very well have been acquired from an organism that was in no way a direct ancestor. [6] In view of the above, some scientists have started to consider that evolution is perhaps better represented by a network than by a tree. [7] Studies have also begun into new ways of creating a universal tree of life. If taking a single gene had become insufficient for consistent tree representation, now that hundreds of whole genomic sequences are available, new phylogenomic methods are being developed. [8] As it is difficult to align the sequences of two genomes, several methods that use traditional sequence alignment tools have been developed to construct genome trees. [8, 9, 10] These methods involve concatenating the homologous sequences from different gene families to construct a single tree [9, 10, 11] or comparing different trees to create a supertree. [12] Another way

to describe the relationships between genomes is to use their gene repertoire. [13] New methods based on gene order or gene content have therefore been developed. [10] The main problem with these methods is the imbalance in the number of genes between small and large genomes. Two large genomes that are not phylogenetically closely related can have more common genes than a large and a small genome that are closely related. Measures to prevent this must be taken so that the phylogenetic tree does not become biased. [10]

Genome trees seem to reveal a phylogenetic signal that supports the three-domain evolutionary scenario and the relationships between some clades of Bacteria. However, deep-level prokaryotic relationships are difficult to infer. [12] We have developed a new method for constructing a genome tree based on the metabolic pathways present in each species. The main structure of the metabolic pathways seems to be largely unaffected by HGT. [14] This enables us to use them as templates for comparing genomes. Using the orthologous groupings of enzymes found in the KEGG database, we have related genomes and metabolic pathways and created a tree-like representation of a fairly large group of organisms based on their metabolism.

Methodology:

Our aim was to create a dendrogram of different eukaryotic and prokaryotic species based on metabolic data. Here we detail the characteristics of the process used:

Database creation

Starting from the metabolic maps available in the KEGG: Kyoto Encyclopedia of Genes and Genomes [15] MetaCyc (http://www.genome.ad.jp/kegg/) and the [16] (http://www.metacyc.org) databases, we defined a representative group of pathways and introduced into our database the enzymes that catalyse each of the reactions that form every pathway by their KO number as defined in KEGG. Since a same pathway can follow slightly different routes in different organisms, we

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added different variants to some of the pathways. For example, we introduced five variants of the glycolysis pathway. At the end, our database contained 434 pathways and 1157 enzymes with different KO numbers.

Percentage matrix

The next step was to relate the data found in our database to a group of organisms. We used the complete genomes found in the KEGG database. For each organism, we created a list of enzymes codified in the genome, listed by their K number. Since the KEGG database is still growing and new genomes are being introduced, some of them still did not have all their KEGG numbers assigned. So, we compared the number of proteins with an assigned KEGG number to the total number of proteins coded in each genome. Those organisms in which the assigned number of proteins in the KEGG database was less than 20 percent were excluded from the list of organisms used to build the dendrogram. Finally we took 282 organisms which are listed in supplementary Table 1 with their abbreviation. Using information from the metabolic database we had previously created, we searched in each genome for the enzymes that completed each pathway. To do so, we made a PERL script that calculated the percentages of enzymes that appeared in a pathway for each organism. The results were presented in a matrix whose rows were the pathways, whose columns were the organisms analysed and in which each element represented the percentage of enzymes of a pathway that one organism contains.

Dendrogram construction

By calculating the Pearson Correlation with the enzyme percentages of all pathways for each pair of organisms, we transformed the percentage matrix into a distance matrix containing the metabolic distance between each pair of organisms. From this distance matrix, and using the PAUP* program version 4.0, we built a dendrogram using the neighbour-joining (NJ) algorithm. This dendrogram graphically represents the relationships between organisms based on their metabolism. We also built the dendrogram with the UPGMA algorithm, but this dendrogram was fairly similar to the one obtained by NJ.

Bootstrap calculation

To verify the dendrogram obtained, we developed a new method based on bootstrap calculations to check how robust each cluster was. From the primary percentage matrix, this method creates a certain number of distance matrices (a thousand in our case) by randomly selecting the metabolic pathways and allowing repetition. Using this group of matrices, we followed the same process as before and obtained a thousand trees. Using the consense program of the Phylip package, we calculated a consensus tree using the majority rule extended option with default parameters. The number of times each node is repeated indicates how reliable that cluster is.

Discussion:

Dendrogram based on metabolism

To ensure that the method developed was suitable for creating a dendrogram that would take into account at least the most basic taxonomic classification, we used it on 282 organisms (9 Eukaryota, 23 Archaea and 250 Bacteria) from the KEGG database. The evolution based on metabolic pathways is represented in the dendrogram in Figure 1. To make comparison easier, we have coloured the branches according to the taxonomic classification of their organism and classified the ISSN 0973-2063

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organisms into fourteen groups. These groups, which differ in size, were defined by taking into account the clusters observed in Figure 1 and their bootstrap values. The result of the groupings and the taxonomic group to which each organism belongs are shown in Table 1 (supplementary material). In general, although this dendrogram does not follow the taxonomic classification perfectly, some large clusters encompass taxonomically related organisms while others appear as mixed clusters. Here we comment two causes that may lead to the grouping of mixed taxonomic clusters.

Reduced genomes

All Archaea are clustered together separately from the bacterial cluster, the only exception is Nanoarchaeum equitans Kin4-M (neq). Unlike the other Archaea we used to construct the dendrogram, this organism is an obligate symbiont. [17] It appears clustered with most of the intracellular or obligate parasites with a small genome found in our dendrogram (groups 4, 5 and 6). Parasitic organisms have reduced genomes, which means that their metabolic capacity has been lowered to a certain degree. This could explain the clustering of several parasite species even though they are phylogenetically distant. In a tree based on metabolic information, therefore, it should not be surprising to find that the only symbiont Archaea clusters with other parasites due to their particular metabolic characteristics.

Metabolic similarity

The firmicutes are grouped in two main groups, Lactobacillales (Group 9) and Bacillales (Group 10). Between these two groups there are smaller groups of other Firmicutes, one of which contains the Clostridia Thermoanaerobacter tengcongensis (tte) and Clostridium tetani (ctc) with two other organisms that do not belong to the Firmicutes phylum: Symbiobacterium thermophilum (sth) and Fusobacterium nucleatum (fnu). The location of F. nucleatum among Firmicutes can be explained by their shared metabolic pathways. [18] Despite being gram negative, F. nucleatum has been found to be more similar to gram positive bacteria than to gram negative ones. This is also true of S. thermophilum. The 16S ribosomal DNA-based phylogeny suggested that this bacterium belongs to an unknown taxon in the gram-positive Actinobacteria [19], even though the traditional Gram-stain result indicates that it is gram negative. [20] Also, the proteins of S. thermophilum show a greater similarity to the proteins found in Firmicutes organisms, in particular to T. tengcongensis, than to those found in Actinobacteria. [20] The metabolic similarity between S. thermophilum and T. tengcongensis shown in figure 1 and the metabolic dissimilarity between S. thermophilum and the other Actinobacteria, combined with previous evidences [20, 21], suggest that S. thermophilum may be re-classified as Firmicutes, Clostridia. [21]

Metabolic influence

Not all kinds of metabolism influence our dendrogram in the same way. In Table 2 (supplementary material) we can see a distribution of the enzymes found in the defined groups in the different metabolic groups. For example, Carbohydrate Metabolism has much more influence on the dendrogram than Energy Metabolism, simply because it has many more enzymes and pathways. Also, some of these enzymes are not very useful for classifying organisms into the different clusters. A clear example is the enzymes that catalyse the reactions that produce the different Aminoacyl-tRNAs as they are present in nearly

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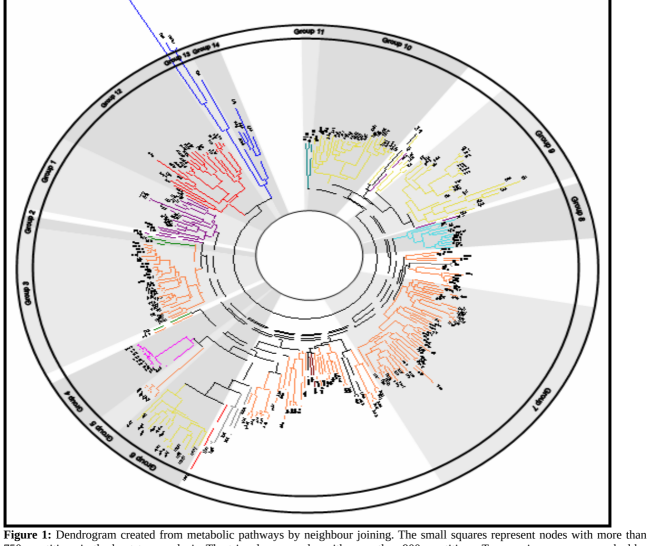
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every group, even those with a reduced genome.

Table 2 (under supplementary material) also shows that for several groups some kinds of metabolisms stand out because of the high number of enzymes they possess compared to the main number of enzymes that the metabolic group has in all organisms. For example, Lipid metabolism in Metazoa (Group 13). This is explained by the presence of pathways such as the synthesis of Lecitin or Cholesterol. The contrary is also true. Some groups have fewer enzymes than most. Examples of this are the three parasitic groups (Group 4, 5 and 6). In their low enzyme values, we can clearly see the effects of genome evolutive reduction due to their parasitic nature.

Limitations of metabolic-based methods

By their nature, metabolic pathways databases are humandefined and may be quite inexact, especially when a metabolic pathway found in one species is generalized to another. Several alternative pathways that have not yet been discovered surely exist in different organisms. Therefore, when only one or a few enzymes from a metabolic pathway are missing in one species, an orthologous gene displacement needs to be considered before we can conclude that the pathway is incomplete. Moreover, when a new sequenced genome is annotated, a high percentage of its proteins are not mapped to any pathway. It may therefore be argued that metabolic databases, while extremely useful for reconstructing metabolic properties of organisms, cannot be used to reconstruct the tree of life. However, we have shown that, assuming that any metabolic prediction of a large group of organisms is still incomplete, the phylogenetic signal that it contains partially agrees with the taxonomic information of the species. A metabolic dendrogram of different species can therefore be used as an additional criterion that may help to correctly re-classify some species, as in the case of the Symbiobacterium thermophilum we described earlier.



750 repetitions in the bootstrap analysis. The triangles are nodes with more than 900 repetitions. Taxonomic groups are marked by the same colouring: Actinobacteria in purple, Archaea in red, Bacteroidetes in green, Chlamydiae in pink, Cyanobacteria in pale blue, Deinococcus-Thermus in cyan, Eukaryota in dark blue, Firmicutes in yellow, Proteobacteria in orange, Spirochaeta in grey, and others in black. ISSN 0973-2063 137

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Conclusion:

We have developed a new method for constructing a dendrogram based on metabolic comparisons between species whose genome has been fully sequenced. Although the evolutionary signal that can be derived from metabolic data is not very strong, it is enough to obtain a rough sketch of the known taxonomic classification. We expect that the reconstruction of metabolic dendrograms may improve as more pathways are discovered and their enzymes are properly situated within those pathways. Until such a time metabolic-based dendrograms may be a useful addition when they are combined with other phylogenetic methods, allowing us to fine-tune dubious classifications that can not be accurately described by other methods.

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Supplementary material

	Eukaryota			Firmicutes (cont)	
Abbr	Organism Name	Group	Abbr	Organism Name	Group
cal	Candida albicans SC5314	14	spn	Streptococcus pneumoniae TIGR4	9
cme	Cyanidioschyzon merolae	14	spy	Streptococcus pyogenes M1 GAS	9
ago	Eremothecium gossypii	14	spa	Streptococcus pyogenes MGAS10394	9
hsa	Homo sapiens	13	spg	Streptococcus pyogenes MGAS315	9
mmu	Mus musculus	13	spz	Streptococcus pyogenes MGAS5005	9
sce	Saccharomyces cerevisiae	14	spb	Streptococcus pyogenes MGAS6180	9
spo	Schizosaccharomyces pombe	14	spm	Streptococcus pyogenes MGAS8232	9
ssc	Sus scrofa	-	sps	Streptococcus pyogenes SSI-1	9
xla	Xenopus laevis	13	stc	Streptococcus thermophilus CNRZ1066	9
	Archaea:		stl	Streptococcus thermophilus LMG 18311	9
Abbr	Organism Name	Group	tte	Thermoanaerobacter tengcongensis MB4	-
ape	Aeropyrum pernix	12	uur	Ureaplasma parvum serovar 3 str. ATCC 700970	6
afu	Archaeoglobus fulgidus DSM 4304	12		Proteobacteria:	
hma	Haloarcula marismortui ATCC 43049	12	Abbr	Organism Name	Group
hal	Halobacterium sp. NRC-1	12	aci	Acinetobacter sp. ADP1	7
mja	Methanocaldococcus jannaschii DSM 2661	12	atc	Agrobacterium tumefaciens str. C58 (Cereon)	7
mmp	Methanococcus maripaludis	12	atu	Agrobacterium tumefaciens str. C58	7
1	Ĩ			(U.Washington/Dupont)	
mka	Methanopyrus kandleri AV19	12	ama	Anaplasma marginale str. St. Maries	-
mac	Methanosarcina acetivorans C2A	12	eba	Azoarcus sp. EbN1	7
mba	Methanosarcina barkeri str. fusaro	12	bhe	Bartonella henselae str. Houston-1	-
mma	Methanosarcina mazei Gol	12	bqu	Bartonella quintana str. Toulouse	-
mth	Methanothermobacter thermautotrophicus str. Delta H	12	bba	Bdellovibrio bacteriovorus HD100	-
neq	Nanoarchaeum equitans Kin4-M	-	bbr	Bordetella bronchiseptica RB50	7
nph	Natronomonas pharaonis DSM 2160	12	bpa	Bordetella parapertussis 12822	7
pto	Picrophilus torridus DSM 9790	12	bpe	Bordetella pertussis Tohama I	7
pai	Pyrobaculum aerophilum	12	bja	Bradyrhizobium japonicum USDA 110	7
pab	Pyrococcus abyssi GE5	12	bmb	Brucella abortus biovar 1 str. 9-941	7
pfu	Pyrococcus furiosus DSM 3638	12	bme	Brucella melitensis 16M	7
pho	Pyrococcus horikoshii OT3	12	bmf	Brucella melitensis biovar Abortus 2308	7
sai	Sulfolobus acidocaldarius DSM 639	12	bms	Brucella suis 1330	7
SSO	Sulfolobus solfataricus P2	12	bab	Buchnera aphidicola (Baizongia pistaciae)	-
sto	Sulfolobus tokodaii str. 7	12	buc	Buchnera aphidicola str. APS (Acyrthosiphon pisum)	-
tko	Thermococcus kodakarensis KOD1	12	bas	Buchnera aphidicola str. Sg (Schizaphis graminum)	-
tac	Thermoplasma acidophilum DSM 1728	12	bma	Burkholderia mallei ATCC 23344	7
tvo	Thermoplasma volcanium	12	bps	Burkholderia pseudomallei K96243	7
	Actinobacteria:		bur	Burkholderia sp. 383	7
Abbr	Organism Name	Group	cjr	Campylobacter jejuni RM1221	-
blo	Bifidobacterium longum NCC2705	-	cje	Campylobacter jejuni subsp. jejuni NCTC 11168	-
cdi	Corynebacterium diphtheriae NCTC 13129	1	bfl	Candidatus Blochmannia floridanus	-
cef	Corynebacterium efficiens YS-314	1	bpn	Candidatus Blochmannia pennsylvanicus str. BPEN	-
cgb	Corynebacterium glutamicum ATCC 13032 (Bielefeld)	1	pub	Candidatus Pelagibacter ubique HTCC1062	7
cgl	Corynebacterium glutamicum ATCC 13032 (Kyowa Hakko)	1	ccr	Caulobacter crescentus CB15	7
cjk	Corynebacterium jeikeium K411	1	cvi	Chromobacterium violaceum ATCC 12472	7
lxx	Leifsonia xyli subsp. xyli str. CTCB07	1	cps	Colwellia psychrerythraea 34H	7

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mpa	Mycobacterium avium subsp. paratuberculosis K-10	1	cbu	Coxiella burnetii RSA 493
mbo	Mycobacterium bovis AF2122/97	1	dar	Dechloromonas aromatica RCB
mle	Mycobacterium leprae TN	1	dps	Desulfotalea psychrophila LSv54
mtc	Mycobacterium tuberculosis CDC1551	1	dvu	Desulfovibrio vulgaris subsp. vulgaris str. Hildenborough
mtu	Mycobacterium tuberculosis H37Rv	1	ecn	Ehrlichia canis str. Jake
nfa	Nocardia farcinica	1	erg	Ehrlichia ruminantium str. Gardel
pac	Propionibacterium acnes KPA171202	-	eru	Ehrlichia ruminantium str. Welgevonden (South Africa)
sma	Streptomyces avermitilis MA-4680	1	erw	Ehrlichia ruminantium str. Welgevonden (France)
sco	Streptomyces coelicolor A3(2)	1	eca	Erwinia carotovora subsp. atroseptica SCRI1043
sth	Symbiobacterium thermophilum IAM 14863	-	ecc	Escherichia coli CFT073
tfu	Thermobifida fusca YX	1	ecj	Escherichia coli K12 W3110
twh	Tropheryma whipplei str. Twist	1	eco	Escherichia coli K12 MG1655
tws	Tropheryma whipplei TW08/27	1	ecs	Escherichia coli O157:H7
	Bacteroidetes:		ece	Escherichia coli O157:H7 EDL933
Abbr	Organism Name	Group	ftu	Francisella tularensis subsp. tularensis
bfs	Bacteroides fragilis NCTC 9343	2	gsu	Geobacter sulfurreducens PCA
bfr	Bacteroides fragilis YCH46	2	gox	Gluconobacter oxydans 621H
bth	Bacteroides thetaiotaomicron VPI-5482	2	hdu	Haemophilus ducreyi 35000HP
pgi	Porphyromonas gingivalis W83	-	hit	Haemophilus influenzae 86-028NP
	Chlamydiae:		hin	Haemophilus influenzae Rd KW20
Abbr	Organism Name	Group	hhe	Helicobacter hepaticus ATCC 51449
cmu	Chlamydia muridarum Nigg	4	hpy	Helicobacter pylori 26695
cta	Chlamydia trachomatis A/HAR-13	4	hpj	Helicobacter pylori J99
ctr	Chlamydia trachomatis D/UW-3/CX	4	ilo	Idiomarina loihiensis L2TR
cab	Chlamydophila abortus S26/3	4	lpf	Legionella pneumophila str. Lens
cca	Chlamydophila caviae GPIC	4	lpp	Legionella pneumophila str. Paris
сра	Chlamydophila pneumoniae AR39	4	lpn	Legionella pneumophila subsp. pneumophila str. Philadelphia 1
cpn	Chlamydophila pneumoniae CWL029	4	msu	Mannheimia succiniciproducens MBEL55E
срј	Chlamydophila pneumoniae J138	4	mlo	Mesorhizobium loti MAFF303099
cpt	Chlamydophila pneumoniae TW-183	4	mca	Methylococcus capsulatus str. Bath
pcu	Parachlamydia sp. UWE25	4	ngo	Neisseria gonorrhoeae FA 1090
•	Cyannobacteria:		nme	Neisseria meningitidis MC58
Abbr	Organism Name	Group	nma	Neisseria meningitidis Z2491
ava	Anabaena variabilis ATCC 29413	8	nwi	Nitrobacter winogradskyi Nb-255
gvi	Gloeobacter violaceus	8	noc	Nitrosococcus oceani ATCC 19707
ana	Nostoc sp. PCC 7120	8	neu	Nitrosomonas europaea ATCC 19718
pmt	Prochlorococcus marinus str. MIT 9313	8	pmu	Pasteurella multocida subsp. multocida str. Pm70
pmn	Prochlorococcus marinus str. NATL2A	8	pca	Pelobacter carbinolicus DSM 2380
pma	Prochlorococcus marinus subsp. marinus str. CCMP1375	8	ppr	Photobacterium profundum
pmm	Prochlorococcus marinus subsp. pastoris str. CCMP1986	8	plu	Photorhabdus luminescens subsp. laumondii TTO1
syc	Synechococcus elongatus PCC 6301	8	pha	Pseudoalteromonas haloplanktis TAC125
syw	Synechococcus sp. WH 8102	8	pae	Pseudomonas aeruginosa PAO1
	Synechocystis sp. PCC 6803	8	pfl	Pseudomonas fluorescens Pf-5
syn	Thermosynechococcus elongatus BP-1	8	pfo	Pseudomonas fluorescens PfO-1
syn tel	Thermosynechococcus elongalus BF-1			Pseudomonas putida KT2440
	Deinococcus-Thermus:		ppu	1 sendomonus pundu K12440
tel		Group	ppu psp	Pseudomonas syringae pv. phaseolicola 1448A
	Deinococcus-Thermus:	Group 11		

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ttj	Thermus thermophilus HB8	11	par	Psychrobacter arcticus 273-4
	Firmicutes:		reu	Ralstonia eutropha JMP134
bbr	Organism Name	Group	rso	Ralstonia solanacearum GMI1000
oaa	Bacillus anthracis str. A2012	10	rsp	Rhodobacter sphaeroides 2.4.1
an	Bacillus anthracis str. Ames	10	rpa	Rhodopseudomonas palustris CGA009
oar	Bacillus anthracis str. 'Ames Ancestor'	10	rco	Rickettsia conorii str. Malish 7
oat	Bacillus anthracis str. Sterne	10	rfe	Rickettsia felis URRWXCal2
ca	Bacillus cereus ATCC 10987	10	rpr	Rickettsia prowazekii str. Madrid E
oce	Bacillus cereus ATCC 14579	10	rty	Rickettsia typhi str. Wilmington
CZ	Bacillus cereus E33L	10	sec	Salmonella enterica subsp. enterica serovar Choleraesuis str. SC-B67
bcl	Bacillus clausii KSM-K16	10	spt	Salmonella enterica subsp. enterica serovar Paratyphi A str. ATCC 9150
oha	Bacillus halodurans	10	sty	Salmonella enterica subsp. enterica serovar Typhi str. CT18
old	Bacillus licheniformis DSM13	10	stt	Salmonella enterica subsp. enterica serovar Typhi Ty2
bli	Bacillus licheniformis ATCC 14580	10	stm	Salmonella typhimurium LT2
su	Bacillus subtilis subsp. subtilis str. 168	10	son	Shewanella oneidensis MR-1
tk	Bacillus thuringiensis serovar konkukian str. 97-27	10	sfx	Shigella flexneri 2a str. 2457T
ac	Clostridium acetobutylicum ATCC 824	-	sfl	Shigella flexneri 2a str. 301
pe	Clostridium perfringens str. 13	9	ssn	Shigella sonnei Ss046
tc	Clostridium tetani E88	-	sil	Silicibacter pomeroyi DSS-3
fa	Enterococcus faecalis V583	9	sme	Sinorhizobium meliloti 1021
ca	Geobacillus kaustophilus HTA426	10	tbd	Thiobacillus denitrificans ATCC 25259
c	Lactobacillus acidophilus NCFM	9	vch	Vibrio cholerae O1 biovar eltor str. N16961
0	Lactobacillus johnsonii NCC 533	9	vfi	Vibrio fischeri ES114
l	Lactobacillus plantarum WCFS1	9	vpa	Vibrio parahaemolyticus RIMD 2210633
a	Lactobacillus sakei subsp. sakei 23K	9	vvu	Vibrio vulnificus CMCP6
a	Lactococcus lactis subsp. lactis Il1403	9	vvy	Vibrio vulnificus YJ016
n	Listeria innocua Clip11262	-	wbr	Wigglesworthia glossinidia endosymbiont of Glossina brevipalpis
no	Listeria monocytogenes EGD-e	-	wol	Wolbachia endosymbiont of Drosophila melanogaster
mf	Listeria monocytogenes str. 4b F2365	-	wbm	Wolbachia endosymbiont strain TRS of Brugia malayi
nfl	Mesoplasma florum L1	6	wsu	Wolinella succinogenes DSM 1740
ga	Mycoplasma gallisepticum R	6	xac	Xanthomonas axonopodis pv. citri str. 306
ge	Mycoplasma genitalium G37	6	xcb	Xanthomonas campestris pv. campestris str. 8004
hy	Mycoplasma hyopneumoniae 232	6	xcc	Xanthomonas campestris pv. campestris str. ATCC 33913
hp	Mycoplasma hyopneumoniae 7448	6	XCV	Xanthomonas campestris pv. vesicatoria str. 85- 10
nhj	Mycoplasma hyopneumoniae J	6	X00	Xanthomonas oryzae pv. oryzae KACC10331
mo	Mycoplasma mobile 163K	6	xfa	Xylella fastidiosa 9a5c
ny	Mycoplasma mycoides subsp. mycoides SC str. PG1	6	xft	Xylella fastidiosa Temecula l
ipe	Mycoplasma penetrans	6	ypm	Yersinia pestis biovar Medievalis str. 91001
pn	Mycoplasma pneumoniae M129	6	ype	Yersinia pestis CO92
pu	Mycoplasma pulmonis UAB CTIP	6	ypk	Yersinia pestis KIM
isy	Mycoplasma synoviae 53	6	yps	Yersinia pseudotuberculosis IP 32953
oih	Oceanobacillus iheyensis HTE831	10	zmo	Zymomonas mobilis subsp. mobilis ZM4
ooy	Onion yellows phytoplasma	6		Spirochaetas:
ab	Staphylococcus aureus RF122	10	Abbr	Organism Name
sac	Staphylococcus aureus subsp. aureus COL	10	lic	Leptospira interrogans serovar Copenhageni str. Fiocruz L1-130

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sar	Staphylococcus aureus subsp. aureus MRSA252	10	lil	Leptospira interrogans serovar Lai str. 56601	-
sas	Staphylococcus aureus subsp. aureus MSSA476	10	bbu	Borrelia burgdorferi B31	-
sav	Staphylococcus aureus subsp. aureus Mu50	10	bga	Borrelia garinii PBi	-
sam	Staphylococcus aureus subsp. aureus MW2	10	tde	Treponema denticola ATCC 35405	-
sau	Staphylococcus aureus subsp. aureus N315	10	tpa	Treponema pallidum subsp. pallidum str. Nichols	-
sep	Staphylococcus epidermidis ATCC 12228	10		Others:	
ser	Staphylococcus epidermidis RP62A	10	Abbr	Organism Name	Group
sha	Staphylococcus haemolyticus JCSC1435	10	aae	Aquifex aeolicus VF5	-
ssp	Staphylococcus saprophyticus subsp. saprophyticus	10	cch	Chlorobium chlorochromatii CaD3	-
sag	Streptococcus agalactiae 2603V/R	9	cte	Chlorobium tepidum TLS	-
sak	Streptococcus agalactiae A909	9	det	Dehalococcoides ethenogenes 195	-
san	Streptococcus agalactiae NEM316	9	deh	Dehalococcoides sp. CBDB1	-
smu	Streptococcus mutans UA159	9	fnu	Fusobacterium nucleatum subsp. nucleatum ATCC 25586	-
spr	Streptococcus pneumoniae R6	9	tma	Thermotoga maritima MSB8	-

 Table 1: Abbreviation and taxonomic classification of the 282 organisms included in the analysis

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	Taxonomic classification (organisms belonging to classification/ total organisms group)		Carbohydrate Metabolism		Energy Metabolism	Lipid Metabolism		Nucleotide Metabolism		Amino Acid Metabolism		Metabolism of Other Amino Acids		Glycan Biosíntesis and Metabolism		Polyketides and Non ribosomal Peptides		Metabolism of Cofactors and Vitamins		Biosíntesis of Secondary Metabolites		Biodegradation of Xenobiotics		tRNAs		Total Enzymes
Group1	Actinobacteria (17/17)	176	(25.9)	28	(4.1)	22	(3.2)	51	(7.5)	207	(30.5)	20	(2.9)	8	(1.2)	4	(0.5)	116	(17.0)	15	(2.3)	14	(2.0)	20	(2.9)	679
Group2	Bacteroidetes (3/3)	201	(29.2)	34	(4.9)	12	(1.8)	48	(7.0)	199	(28.9)	18	(2.7)	23	(3.4)	4	(0.6)	113	(16.4)	13	(1.9)	0	(0.0)	22	(3.2)	688
Group3	Enterobacteria, Vibrionales, Pasteurellales (29/29)	216	(27.8)	38	(4.9)	21	(2.8)	52	(6.6)	224	(28.8)	27	(3.4)	25	(3.2)	2	(0.3)	129	(16.6)	14	(1.8)	7	(0.9)	22	(2.8)	777
Group4	Chlamidiae (10/10)	117	(33.2)	_11	(3.1)	9	(2.6)	_20_	(5.6)	82	(23.3)	5	(1.6)	19	(5.4)	0	(0.1)	63	(17.8)	6	(1.7)	0	(0.0)	20	(5.7)	352
Group5	Rickettsia (4/4)	71	(28.1)	8	(3.4)	6	(2.6)	20	(8.0)	53	(21.0)	7	(2.8)	20	(8.0)	1	(0.4)	38	(15.3)	5	(1.9)	1	(0.6)	20	(8.0)	251
Group6	Mollicutes (14/14)	96	(50.9)	10	(5.3)	4	(1.9)	14	(7.2)	19	(10.0)	4	(2.1)	0	(0.0)	0	(0.0)	21	(10.9)	1	(0.6)	1	(0.3)	21	(10.9)	189
Group7	Proteobacteria (43/43)	188	(24.3)	36	(4.7)	27	(3.5)	52	(6.8)	240	(31.1)	25	(3.3)	21	(2.8)	3	(0.4)	120	(15.6)	16	(2.0)	21	(2.7)	21	(2.7)	771
Group8	Cyanobacteria (11/11)	160	(24.7)	29	(4.4)	17	(2.6)	48	(7.4)	190	(29.4)	18	(2.8)	15	(2.3)	4	(0.5)	123	(19.0)	19	(3.0)	4	(0.6)	20	(3.2)	647
Group9	Lactobacillales (21/22)	152	(31.6)	25	(5.2)	15	(3.1)	47	(9.8)	126	(26.3)	14	(2.9)	7	(1.5)	3	(0.7)	53	(11.1)	13	(2.8)	2	(0.5)	21	(4.4)	479
Group10	Bacillales (26/26)	193	(26.9)	32	(4.4)	27	(3.7)	50	(7.0)	229	(31.9)	27	(3.7)	7	(1.0)	2	(0.3)	108	(15.0)	16	(2.2)	7	(1.0)	21	(2.9)	718
Group11	Deinococcus- Thermus (3/3)	175	(25.7)	34	(5.0)	23	(3.3)	50	(7.3)	227	(33.2)	18	(2.6)	7	(1.0)	2	(0.3)	102	(15.0)	13	(2.0)	9	(1.4)	22	(3.2)	682
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Group12	Archaea (23/23)	122	(25.2)	19	(4.0)	11	(2.4)	44	(9.0)	168	(34.9)	11	(2.2)	2	(0.5)	3	(0.6)	64	(13.2)	13	(2.7)	6	(1.3)	19	(4.0)	483
Group13	Metazoa (3/3)*	202	(28.8)	22	(3.1)	71	(10.1)	52	(7.4)	194	(27.7)	20	(2.9)	14	(2.0)	1	(0.1)	77	(11.1)	17	(2.5)	8	(1.1)	21	(3.0)	699
Group14	Fungi (4/5)	183	(25.9)	29	(4.1)	31	(4.4)	49	(7.0)	237	(33.6)	21	(3.0)	14	(2.0)	1	(0.1)	93	(13.2)	16	(2.3)	9	(1.2)	22	(3.1)	706

Table 2: Metabolic influence over the different clusters. For each group of organisms the mean number of enzymes involved in each kind of metabolism and the percentage of enzymes that belong to a determined metabolism in comparison to the total number of enzymes used to create the dendrogram are shown. Green and red numbers or percentages indicate a group of organisms that has more or lower enzymes of a kind of metabolism than most of the other groups. * Sus scrofa (ssc) was excluded from these data because of the lack of KEGG numbers on important metabolic protein