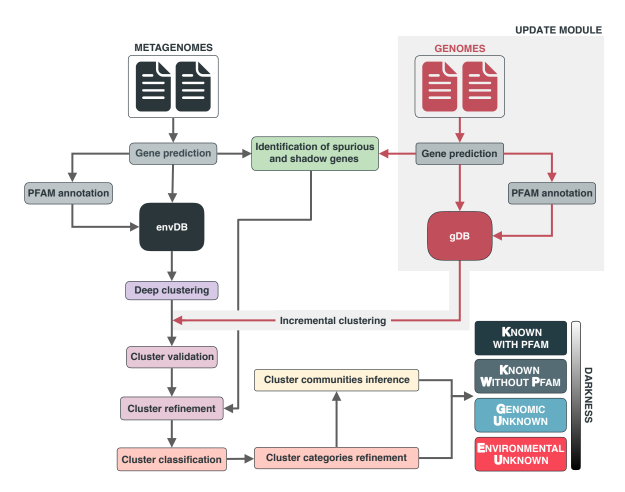
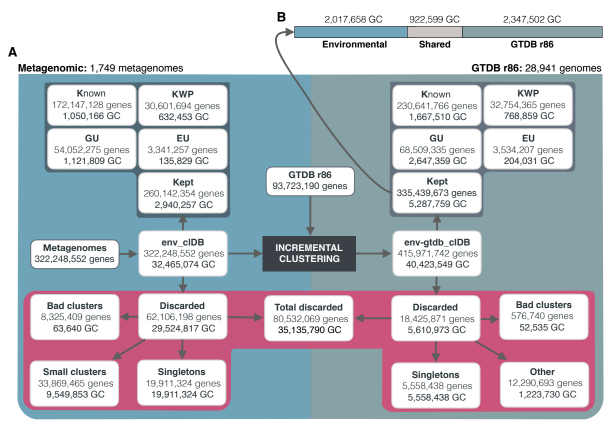
Supplementary figures



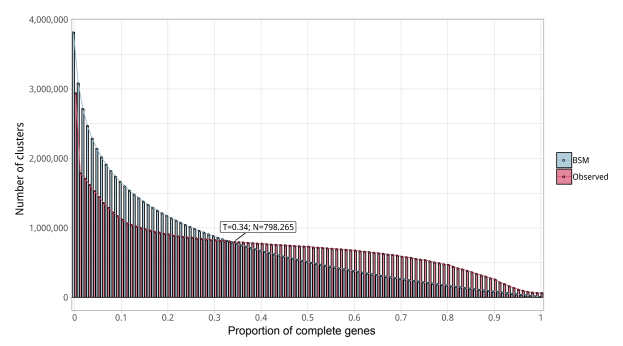


Supplementary Figure 1. Overview of the workflow to partition the genomic and metagenomic coding sequence space between known and unknown. The workflow performs gene prediction, gene clustering, gene clustering validation and refinement, GCC inference, and partitions the coding sequence space in the different known and unknown categories.

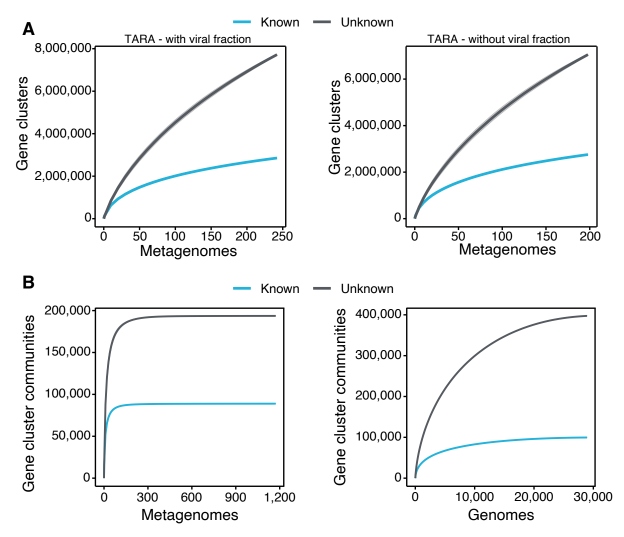


Supplementary Figure 2. The diagram shows a schematic description of the number of genes and GCs that have been kept or discarded. (A) We analyzed a dataset of 1,749 metagenomes from marine and human environments and 28,941 genomes from the GTDB_r86 summing up to 415,971,742 genes. The composition of the genomic box "Other" is described in supplementary Note 5. (B) GC overlap between the environmental and genomic datasets.

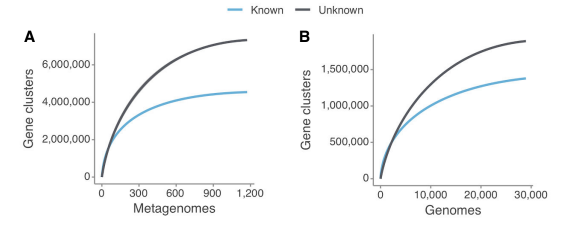




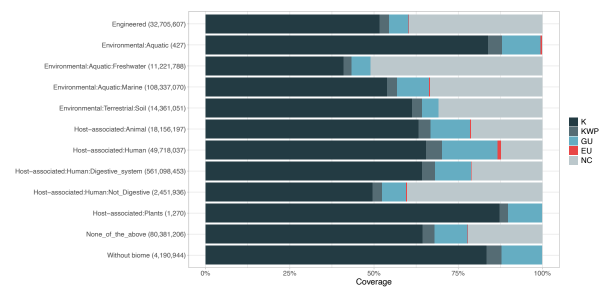
Supplementary Figure 3. Proportion of complete genes per cluster. Distribution of observed values compared with those generated by the Broken-stick model. The cut-off was determined at 34% complete genes per cluster.



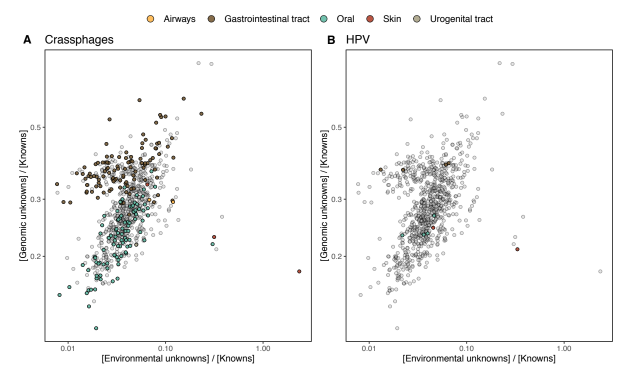
Supplementary Figure 4: Collector curves for the known and unknown coding sequence space. (A) Collector curves at the gene cluster level, for the TARA metagenomes, including the viral fraction (left) and excluding it (right) from the analysis. (B) Collector curves at gene cluster community level for the metagenomes from TARA, MALASPINA, and HMP-I/II projects (left) and the 28,941 GTDB genomes (right).



Supplementary Figure 5: Collector curves for the known and unknown coding sequence space at the gene cluster communities level for (A) the metagenomes from TARA, MALASPINA and HMP-I/II projects, and for (B) the 28,941 GTDB genomes. Singletons were excluded from the calculations.



Supplementary Figure 6. Proportion of gene cluster categories per biome. On the y-axis are reported the 11 main biome categories indicated by MGnify and in parenthesis the total number of genes in each biome. The gray fraction represents the pool of genes from MGnify that were not found in our dataset.



Supplementary Figure 7. HMP outlier samples enriched in (A) crAssphages, and (B) papillomaviruses (HPV).

51 Supplementary Tables

Supplementary Table 1. Number of metagenomic clusters and genes after the validation and refinement steps.

	Good-quality	Bad-quality	Total
Clusters	2,940,257	63,640	32,465,074
Genes	260,142,354	8,325,409	322,248,552

Supplementary Table 2. MG + GTDB high quality (HQ) subset of gene clusters (GCs).

Category	HQ GCs	HQ genes	pHQ GCs	pHQ genes
K	76,718	40,710,936	0.0145	0.120
KWP	16,922	1,733,599	0.00320	0.005132
GU	95,370	9,908,630	0.0180	0.0293
EU	14,207	477,625	0.00269	0.00141
Total	203,217	52,830,790	0.0384	0.1562

Supplementary Table 3. Mean proportion of complete genes per cluster in the four functional categories.

	K	KWP	GU	EU
Mean percentage of complete genes	0.50	0.22	0.68	0.70

Supplementary Table 4. KWP high-quality gene clusters (GCs) distribution in the COG groups. (Full table in Supplementary_tables_1.xlsx)

COG group	Number of GCs	Proportion of GCs
CELLULAR PROCESSES AND SIGNALING	2,292	0.135
INFORMATION STORAGE AND PROCESSING	1,582	0.0935
METABOLISM	1,679	0.0992
POORLY CHARACTERIZED	2,899	0.171
NC	8,470	0.501

Supplementary Table 5. MG + GTDB gene clusters summary statistics.
(Supplementary_tables_2.xlsx)
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Supplementary Table 6. Environmental (metagenomic) dataset description.

(A) Number of samples and sites per metagenomic project.

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Dataset	Reference	Samples	Sites	Contigs
TARA	Sunagawa et al.	242	141	62,404,654
Malaspina	Duarte et al.	116	30	9,330,293
OSD	Kopf et al. ³	145	139	4,127,095
HMP	Lloyd-Price et al.4	1,246	18	80,560,927

Dataset	Reference	Samples	Sites	Reads
GOS	Rush et al.⁵	80	70	12,672,518

(B) Number of predicted genes per completeness category.

Total	"00"	"10"	"01"	"11"
322,248,552	118,717,690	106,031,163	102,966,482	75,694,123

Note: "00"=complete, both start and stop codon identified. "01"=right boundary incomplete. "10"=left boundary incomplete. "11"=both left and right edges incomplete.

Supplementary Table 7. Proportion of genes in each cluster category, and Pfam amino
 acids coverage per cluster category. (Supplementary_tables_1.xlsx)

Supplementary Table 8. List of HMP outlier samples (Supplementary_tables_1.xlsx).

Supplementary Table 9. Summary of the number of EU clusters based on their presence in MAGs and their environmental distribution, obtained with the Levin's Niche Breadth index.

	Total clusters	Broad	Narrow	Non-significant
Total EU	204,031	471	8,421	195,079
EU in MAGs	55,520	88	316	55,116
EU not in MAGs	148,511 (73%)	383 (81%)	8,105 (96%)	140,023 (72%)

Supplementary Table 10. Number of phylogenetic conserved and lineage-specific gene clusters (GCs) in the GTDB bacterial phylogeny. (Supplementary_tables_1.xlsx).

- **Supplementary Table 11.** Clusters in the GU community GU_c_21103
- 94 (Supplementary_tables_1.xlsx).

Supplementary Table 12. Number of lineage-specific gene clusters of unknown function at different taxonomic levels within the *Cand. Patescibacteria* phylum.

Taxonomic level	Number of clusters
Phylum	2
Class	6
Order	104
Family	1,456
Genus	6,987
Species	45,788

Supplementary Table 13. List of filtered samples used for the metagenomic analyses.

(Supplementary_tables_1.xlsx)

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106	Supplementary Table 14. List of terms commonly used to define proteins of unknown
107	function in public databases. (Supplementary_tables_1.xlsx)
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110 Supplementary Notes

111 Supplementary Note 1 - Metagenomic singletons and small

112 gene clusters

Analysis of metagenomic singletons and gene clusters with less than ten genes.

The singletons represent 60% of the gene clusters (GCs) and 6% of the total genes. The GCs with less than ten genes, here referred to as small GCs for simplicity, represent 29% of the GCs and 10.5% of the gene dataset (Supp. Figure 2A). Although we discarded these two sets from the main study, we investigated them to obtain a complete analysis of the initial dataset. Both sets were first searched against the Pfam database of protein domain families⁶, and subsequently classified following the steps described in Supplementary Note 3. For the small GCs classification, we used the cluster consensus sequence, which we extracted using the *hhconsensus* program of the HH-SUITE⁷, from the GC multiple sequence alignments (MSAs), generated with FAMSA⁸.

We could not find any homologous in the Pfam database for the large majority of both singletons and small GCs, 95%, and 89%, respectively (Supp. Table 1-1). After the classification, the large majority of the singletons remained completely uncharacterized, (64% was identified as EU) (Supp. Table 1-2). Similarly, the small GCs were also found dominated by GCs of unknowns, with 38% of the clusters classified as EU and 29% as GU (Supp. Table 1-2).

Supplementary Table 1-1. Singletons and small GCs Pfam annotations.

	Total	Annotated	Not annotated
Singletons	19,911,324	934,548	18,976,776
Small GCs	9,549,853	1,028,076	8,521,777

Supplementary Table 1-2. Number of singletons and small GCs per functional category.

_	К	KWP	GU	EU
Singletons	852,413	3,505,161	2,763,476	12,790,274
Small GCs	946,112	2,213,654	2,744,262	3,645,825

Supplementary Note 2 - Metagenomic gene cluster validation

134 and refinement

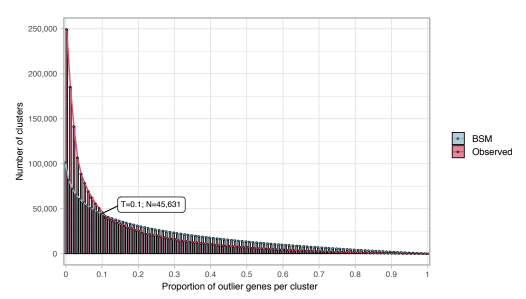
- 135 To obtain a set of gene clusters characterized by a high intra-cluster homogeneity, we
- identified spurious, shadow and outlier genes, and we removed them from the clusters.

- 138 Identification of spurious genes. We identified spurious genes by screening our gene data
- 139 set against the *AntiFam* database ⁹.
- 140 Identification of shadow genes. We identified shadow genes using the procedure described
- in Yooseph et al. ¹⁰. (1) Two genes on the same strand are considered overlapping if their
- 142 intervals overlap by at least 60 bps; (2) genes that are on the opposite strands are
- 143 considered overlapping if their intervals overlap by at least 50 bps, and their 3' ends are
- within each other's intervals, or if their intervals overlap by at least 120 bps and the 5' end of
- one is in the interval of the other.
- 146 Identification of outlier genes. Outlier genes are sequences inside a cluster non-homologous
- to the other cluster genes and were identified during the cluster validation step (see Methods
- 148 Gene cluster validation).
- The number of spurious, shadow and outlier genes identified in the data set is reported in
- 150 Supplementary Table 2-1.
- 151 Cluster refinement. After the validation, we proceeded with the retrieval of the subset of
- 152 "good" clusters. Clusters with ≥ 30% shadow genes were identified as shadow-clusters, as
- proposed in Yooseph et al. ¹⁰. During the cluster validation, we identified a minimum of 10%
- outlier genes as the threshold to classify a cluster as "bad-quality" (Supp. Fig. 2-2; Suppl.
- 155 Table 2-2A). We combined this threshold with a Jaccard similarity index < 1, indicating a low
- 156 intra-cluster Pfam domain architecture (DA) homogeneity, for the Pfam annotated clusters
- 157 (Supp. Table 2-2B). We performed the cluster refinement in three consecutive steps:
- 158 I. Discard the "bad" clusters (≥ 10% outliers & Jaccard similarity index <1)
- 159 II. Discard the "shadow" clusters (≥ 30% shadow genes)
- 160 III. Remove the single shadow, spurious and outlier genes from the remaining clusters.
- 161 The results for each step are shown in Supplementary Table 2-3. From the initial set of ~3M
- 162 clusters with more than ten genes, we identified 57,052 GCs as "bad" and 6,261 as
- 163 "shadow". From the remaining set of 2,940,593 clusters, we removed a total of 2,708,994
- shadow, spurious and outlier genes. During this last step, we discarded 336 more clusters:
- 165 244 resulted being composed only of spurious and outlier genes (one in the Pfam annotated
- set of clusters and 243 in the non-annotated set), and 92 clusters were discarded since they
- were left as singletons after refinement. Besides, we moved 1,190 Pfam annotated clusters

to the non-annotated set since they were left without any annotated gene. In summary, we removed 63,640 GCs and a total of 8,325,409 genes, respectively, 2% and 3% of the initial data set. The refined set contains 2,940,592 GCs and 260,142,354 genes (Supp. Table 3).

Supplementary Table 2-1. Number of spurious, shadow and outlier genes in the metagenomic clusters.

Gene category	Clusters ≥ 10 genes	Clusters < 10 genes	Singletons
Spurious	44,205	6,784	2,335
Shadow	289,258	144,571	177,126
Outliers	3,118,850	-	-



Supplementary Figure 2-1. Proportion of outlier genes detected within each cluster MSA. Distribution of observed values compared with those generated by the Broken-stick model. The cut-off was determined at 10% outlier genes per cluster.

Supplementary Table 2-2. Metagenomic gene cluster validation results.

(A) Evaluation of cluster sequence composition.

	Pre-Compos. validation	good quality	bad quality
Clusters	3,003,897	2,958,266	45,631
Genes	268,467,763	266,268,638	2,199,125

(B) Evaluation of cluster Pfam functional annotations.

Pre-Funct. validation	Funct. good	Funct. bad
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Clusters	1,015,924	1,004,166	11,758
Genes	181,433,541	178,167,583	3,246,002

Supplementary Table 2-3

184 Steps:

185 Step I - Removing of the "bad clusters"

Step II - Removing of the "shadow clusters"

187 Step III - Removing single spurious, shadow or outlier genes

(A) Number of clusters in each step of the cluster refinement.

	Step I	Step II	Step III	Refined
Clusters	3,003,897	2,946,845	2,940,593	2,940,257
Removed	-57,052	-6,252	-336	

(B) Number of genes in each step of the cluster refinement.

	Step I	Step II	Step III	Refined
Genes	268,467,763	263,022,636	262,851,348	260,142,354
Removed	-5,445,127	-171,288	-2,708,994	

Supplementary Note 3 - Metagenomic gene cluster classification and remote homology refinement

197 Classification of the refined subset of gene clusters and remote homology refinement.

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Methods

We searched the gene clusters (GCs) without any Pfam annotated gene against two functional databases, the UniRef90, from UniProt¹¹, and the NCBI *nr* database ¹². We screened the two databases using the cluster consensus sequences, obtained by applying the *hhconsensus* program of the *HH-SUITE*⁷ on the clusters multiple sequence alignments (MSAs) generated with the FAMSA program⁸. We performed two nested searches using the MMSeqs2¹³ program and following a similar workflow as the "2bLCA" described in Hinghamp et al. ¹⁴. The search-workflow consisted of five steps: First, we searched the consensus sequences against the functional database, with -e 1e-05 --cov-mode 2 -c 0.6. Second, we extracted the high scoring pairs (HSP) of the best hits and we searched them again using the same parameters. Third, we merged the top hits from the first with the second search results. Fourth, we filtered out the second search hits with a bigger e-value than the first search top hits. And fifth, we selected the hits that were found in 60% of the log10(best-e-value). We first applied this search-workflow to screen the UniRef90 database (release 2017 11)¹¹. We classified the GCs as GU if their consensus sequences were found annotated to proteins labeled with any of the terms commonly used to define proteins of unknown function in public databases (Supp. Table 14). WE classified, instead, as KWP, the clusters with consensus annotated to functionally characterized proteins. Secondly, we applied the same search-workflow to search the consensus sequences with no homologs in the UniRef90 database, against the NCBI nr database (release 2017 12)12. We used the same criteria to classify a GC as GU or KWP. Ultimately, we classified as EU the GCs whose consensus sequences did not align with any of the NCBI nr entries.

We processed the Pfam annotated GCs to retrieve a GC consensus domain architecture (DA). We classified as GU the GCs with a consensus DA composed only of Pfam domain of unknown function (DUFs) and as K the rest. The methods for this step are described in

Methods - Remote homology classification of gene clusters.

We refined the classified GCs to account for remote homologies. A detailed description of this process can be found in Methods - **Gene cluster remote homology refinement**.

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Results

From the 1,946,737 non-annotated clusters, 1,581,115 were found homologous to UniRef90 entries. Of these hits, more than 50% were found homologous to "hypothetical" proteins and classified as GU, and the other hits were labeled as KWP. The remaining 365,622 clusters, with no homologs to UniRef90, were screened against the NCBI nr database. We found 20,277 clusters in the NCBI nr, of them, 15,998 clusters were homologous to "hypothetical" proteins, and 4,279 clusters to characterized proteins and were classified respectively as GU and KWP. The remaining 345,345 clusters were not found in the NCBI nr database and therefore identified as EU. After the cascaded profile search against UniRef90 and NCBI nr, and the analysis of the GC consensus DAs, we classified the GCs into 912,551 K, 753,718 KWP, 928,643 GU, and 345,345 EU. Detailed results for each search are reported in Supplementary Table 3-1.

Supplementary Table 3-1. Metagenomic gene clusters classification steps.

(A) Results from the search against the UniRef90 database

Search vs UniRef90	Hits		No-hits
Initial clusters:1,946,737	1,581,115		365,622
	Characterized Hypothetical		
	749,439	749,439 831,676	

(B) Results from the search against the and the NCBI nr databases

Search vs NCBI nr	Hits		No-hits
Initial clusters: 365,622	20,277		345,345
	Characterized Hypothetical		
	4,279 15,998		

(C) Classification of the Pfam annotated GCs based on the consensus DAs.

Consensus DA analysis Annotated to DKF DAs		Annotated to DUF DAs
Initial clusters: 993,520	912,551	80,969

Supplementary Table 3-2. Metagenomic GC remote homology refinement steps.

	К	KWP	GU	EU
Initial GCs	912,551	753,718	928,643	345,345
EU refinement	-	+38,333	+171,183	-209,516

Post-EU refinement	912,551	792,051	1,099,826	135,829
KWP refinement	+137,615	-159,598	+21,983	-
Refined GCs	1,050,166	632,453	1,121,809	135,829

Supplementary Note 4 - GTDB integration

Results from the integration of the Genome Taxonomy Database¹⁵ into the metagenomic dataset.

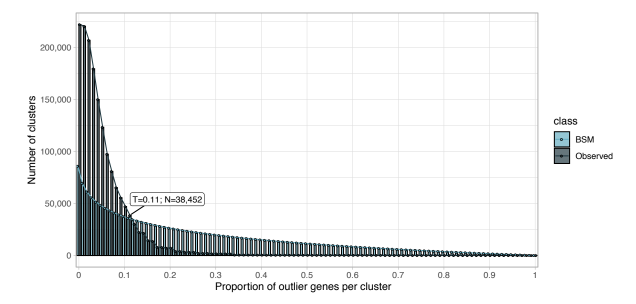
We integrated the metagenomic GCs with the 93,723,190 genes from the archaeal and bacterial GTDB genomes (release 86)¹⁵. A total of 67,446,376 genomic genes, 72% of the whole dataset, were found in the metagenomic GCs. The remaining 26,276,814 (28% of the initial dataset) genes were then clustered separately into 7,958,475 genomic GCs (Supp. Table 4-1). This set of GCs was processed through our workflow steps to be validated, classified and refined.

Within the set of genomic GCs, we identified 5,558,438 singletons and 2,400,037 GCs with more than one gene. We were able to annotate to Pfam protein domain families 41% of the genomic genes. The annotation led to 556,834 annotated GCs and 1,843,203 non-annotated GCs. The validation step determined the minimum proportion of outlier genes per cluster at 11% (Supp. Fig. 4-1). The majority of the genomic GCs showed high intra-cluster homogeneity, both in terms of sequence composition and functional annotations (Supp. Table 4-2).

After the validation, we refined the GCs removing the GCs identified as "bad" and the detected outliers' genes (see Supp. Table 4-3). We classified the refined subset of 2,347,502 GCs into the four functional categories via the same protocol applied for the metagenomic data set. The results of the GC classification are reported in Supplementary Table 4-4. After the classification steps, we refined the EU and KWP GCs searching their HMMs profiles for remote homologies in the Uniclust (release 30_2017_10)¹⁶ and the Pfam (v. 31.0)⁶ databases, respectively, using *HHblits* ¹⁷. An overview of the results step-by-step can be found in Supplementary Table 4-5A. In the end, we obtained 617,344 GCs classified as Known, 136,406 as KWP, 1,525,550 as GU and 68,202 as EU (Supp. Table 4-5B). The genomic dataset appeared highly dominated by the GU, which accounts for 65% of the GCs. In the end, we retrieved a subset of genomic "High Quality" (mostly complete) GCs (Supp. Table 4-6). The numbers of genes and GCs for the integrated (MG+GTDB) dataset are reported in Supplementary Table 4-7.

Supplementary Table 4-1. GTDB integration in the metagenomic dataset.

	Metagenomic	Shared	Genomic	Total
GCs	30,301,693	2,163,381	7,958,475	40,423,549
Genes	199,693,614	190,001,314	26,276,814	415,971,742



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Supplementary Figure 4-1. Proportion of outlier genomic genes identified within each cluster MSA. Distribution of observed values compared with those of the Broken-stick model.

Supplementary Table 4-2. Genomic GC validation results.

(A) Evaluation of cluster sequence composition.

	Pre-Compos. validation	good quality	bad quality
GCs	2,400,037	2,361,585	38,452
Genes	20,718,376	20,364,454	353,922

(B) Evaluation of Pfam functional annotations.

	Pre-Funct. validation	good quality	bad quality
GCs	556,834	542,410	14,424
Genes	10,091,203	9,865,550	225,653

(C) Combined cluster validation results.

	Pre-validation	good quality	bad quality
GCs	2,400,037	2,347,502	52,535
Genes	20,718,376	20,141,636	576,740

Supplementary Table 4-3. Spurious, shadow and outlier genes in the genomic GCs.

Gene category	GCs >= 2 genes	Singletons
Spurious	3,252	1,312
Shadow	223,535	125,262
Outliers	449,080	-

Supplementary Table 4-4. Non-annotated genomic GC classification.

297 (A) Results from the search against the UniRef90 database.

Search vs UniRef90	Hits		No-hits
Initial GCs: 1,816,999	1,570,094		246,905
	Characterized Hypothetical		
	304,004	1,266,090	

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(B) Results from the search against the NCBI nr database.

Search vs NCBI nr	Hits		No-hits
Initial GCs: 246,905	28,704		218,201
	Characterized Hypothetical		
	1,280	27,424	

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(C) Classification of the Pfam annotated GCs based on the consensus DAs.

Consensus DA analysis	DKF DAs	DUF DAs
Initial GCs: 993,520	912,551	65,688

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Supplementary Table 4-5. Genomic GC remote homology refinement and final genomic GC dataset.

304 (A) Remote-homology refinement steps.

	K	KWP	GU	EU
Initial GCs	464,815	305,284	1,359,202	218,201
EU refinement	-	+5,704	+144,295	-149,999
Post-EU refinement	464,815	310,988	1,503,497	68,202
KWP refinement	+152,529	-174,582	+22,053	-
Refined GCs	617,344	136,406	1,525,550	68,202

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(B) Genomic GC refined dataset.

_	К	KWP	GU	EU	Total
Genes	9,997,529	663,107	9,305,621	175,379	20,141,636
GCs	617,344	136,406	1,525,550	68,202	2,347,502

Supplementary Table 4-6. Genomic high quality (HQ) GCs.

Category	HQ GCs	HQ genes	pHQ GCs	pHQ genes
K	12,202	25,105,156	0.0198	0.0096
KWP	4,019	1,349,165	0.0295	0.0214
GU	12,699	8,403,393	0.0083	0.0062
EU	438	471,820	0.0064	0.0074

Supplementary Table 4-7. MG + GTDB seed database. Integrated number of genes and 310 GCs per category.

	К	KWP	GU	EU	Total
Genes	230,641,76	32,754,365	68,509,335	3,534,207	335,439,673
GCs	1,667,510	768,859	2,647,359	204,031	5,287,759

313 Supplementary Note 5 – Summary of the post-genomic

integration dataset

- 315 In-detail description of the integrated metagenomic-genomic dataset.
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- The integration of 93,723,190 genomic genes into the metagenomic dataset (322,248,552
- 318 genes, 32,465,074 GCs) resulted into a dataset of 415,971,742 genes and 40,423,549 GCs
- 319 (Supp. Fig. 2A and Supp. table 4-1). As shown in Supp. Figure 2A, the integrated dataset is
- 320 divided into: (1) "kept" GCs and (2) "discarded" GCs.
- 321 1. The "kept" GCs.
- The "kept" GC dataset contains the 2,940,257 metagenomic "kept" GCs with 260,142,354
- 323 genes (Supp. Fig. 2A), the genomic "kept" 2,347,502 GCs with 20,141,636 genes (Supp.
- Table 4-5B), plus 55,155,683 genomic genes found in the metagenomic set of "kept" GCs
- 325 (Supp. Table 5-1), for a total of 5,287,759 GCs and 335,439,673 genes. A description of the
- 326 integrated "kept" dataset numbers of GCs and genes, and their distribution in the different
- 327 categories can be found in Supp. Figure 2A and Supp. Table 4-7.
- 328 2. The "discarded" GCs.
- 329 The metagenomic "discarded" set includes 8,325,409 genes and 63,640 GCs classified as
- 330 "bad" during the validation and refinement processes (Supp. Note 2), 19,911,324 singletons
- and 33,869,465 genes in 9,549,853 small GCs, i.e. clusters with less than 10 genes (Supp.
- 332 Note 1), for a total of 62,106,198 genes and 29,524,817 GCs.
- 333 The genomic "discarded" dataset consists of 576,740 genes and 52,535 GCs classified as
- 334 "bad", 5,558,438 singletons (Supp. Note 4) and 12,290,693 genomic genes found in
- 335 1,223,730 metagenomic discarded clusters. This last set of genes, labeled as "Other" in
- 336 Supp. Figure 2A, includes 1,578,862 genomic genes found in the set of metagenomic "bad"
- 337 clusters, 7,010,987 genomic genes found in the metagenomic small GCs and 3,700,844
- 338 genomic genes homologous to metagenomic singletons (Supp. Table 5-1).
- The integration of the metagenomic and genomic "discarded" sets resulted in 80,532,069
- 340 genes and 35,135,790 GCs.
- 341 As described above, with the integration of genomic data we enriched metagenomic
- 342 singletons and small GCs. This addition resulted in a set of 52,758 metagenomic singletons
- 343 and 187,953 metagenomic small GCs becoming GCs with more than ten genes. We
- validated and classified the 240,711 GCs in this set. We obtained 223,229 good-quality GCs,
- 345 divided into 17,383 K, 89,205 KWP, 109,636 GU and 7,005 EU.

Supplementary Table 5-1. Overview of genomic genes found homologous to metagenomic genes.

	Total	In MG good- quality GCs	In MG small GCs	In MG singletons	In MG bad- quality GCs
Genes	67,446,376	55,155,683	7,010,987	3,700,844	1,578,862

Supplementary Note 6 - Gene cluster additional information

Additional information on the metagenomic and genomic (MG + GTDB) gene cluster dataset.

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We retrieved a set of statistics for the MG + GTDB GC dataset, including the proportion of complete genes per cluster, the average gene length, the cluster level of darkness and disorder, and a cluster consensus taxonomic affiliation. The methods we applied to obtain these statistics are described in the Methods-Gene cluster characterization paragraph. Overall the K category has the largest average GC size, 139.6 genes (and a max of 168,822 genes). The average GC size is then decreasing from the known to the unknown categories, with the EU presenting the smallest average size, with 17.36 genes per GC. Similarly, the K GCs have, on average, the longest genes (258.55 aa), followed by the GU (177.16 aa), the KWP (133.22 aa) and the EU (130.65 aa). The unknown categories (GU and EU) have the highest level of completion, i.e., the proportion of complete genes per GC. The KWP GCs contain the smallest percentage of complete genes. We evaluated the levels of darkness and disorder of the GCs using the information on the DPD¹⁸ annotations (Supp. Table 6-1). The categories K, KWP and GU showed a degree of darkness inversely proportional to their functional characterization. Interestingly the KWP presented the highest level of disorder (Supp. Table 6, Supp Fig 3), while the proper characterization of these proteins is beyond the scope of this paper, our preliminary analyses suggest that KWP are enriched in intrinsically disordered proteins¹⁹ (Supp. Table 6-1). These proteins, usually involved in signaling and regulatory functions, don't have a well-defined 3-D structure and they can adopt many different conformations.

We used the taxonomy of 214,392,608 genes to evaluate the taxonomic variation within a GC and generated consensus taxonomic annotations for 2,630,338 GCs. The GCs taxonomic variation is low at higher taxonomic levels and it steadily increases towards Genus and Species (Supp. Table 5).

A general overview of the MG + GTDB main properties for the whole GCs dataset can be found in Supplementary Table 5 (Supplementary tables 2.xlsx).

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Supplementary Table 6-1. Number of MG + GTDB GCs annotated to the DPD per functional category.

K	KWP	GU	EU
374,555	8,874	22,135	0

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Supplementary Note 7 - Gene cluster communities

Metagenomic and genomic gene cluster community inference detailed results.

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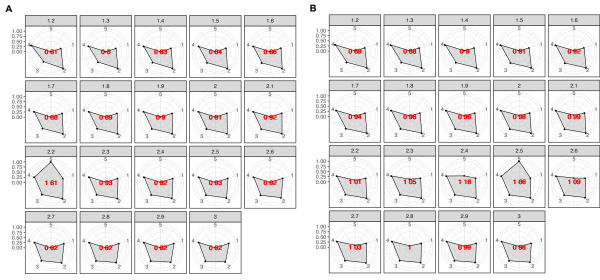
We aggregated the gene clusters (GCs) into gene cluster communities (GCCs) based on their shared distant homologies, which couldn't be detected with the sequence similarity approach. The GCC inference, described in the Methods-Cluster communities inference section, was implemented and tuned on the known coding sequence space (CDS-space), which is constrained by the domain architectures (DAs). Then, we used the information retrieved for the known CDS-space to aggregate the unknown GCs. Since the number of DAs in the known GCs may be inflated due to the fragmented nature of metagenomic genes, a key step for the inference process was the retrieval of a set of non-redundant DAs (Methods - A set of non-redundant domain architectures section).

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397 We reduced the complete set of 29,341 Pfam DAs found in the metagenomic dataset, to 398 23,681 non-redundant DAs, and the 38,765 Pfam DAs found in the genomic dataset to 399 38,060 non-redundant DAs.

To find how the different clusters aggregate at the DA level, we then applied a combination of HMM-HMM searches and community identification using the Markov Cluster Algorithm (MCL) 20 (see Methods - Cluster communities inference). MCL is very sensitive to the inflation value, which determines the granularity of the partitioning. The results of our iterative approach are summarized in the radar plots of Supplementary Figure 7-1. We determined the best inflation value at 2.2 for the metagenomic dataset, value corresponding to the radar plot with the largest area (Supp. Fig. 7-1A). This value is in agreement with the value empirically determined to be the optimal²⁰. The inference led to a set of 283,314 metagenomic GCCs out of ~2.9M GCs, with a reduction rate of 90% (Supp. Table 7-1A).

For the genomic dataset, we first identified the GCs with remote homologies to the metagenomic GCCs. To do this, we searched the genomic GC HMM profiles against the metagenomic ones, using HHblits¹⁷ (-n 2 -Z 10000000 -B 10000000 -e 1). We assigned the genomic GCs sharing a HHblits probability ≥ 50% and a bidirectional coverage > 60% to the respective metagenomic GCCs. We processed the remaining genomic GCs through the GCC inference workflow. We determined the best inflation value at 2.5 (Supp. Fig. 7-1B), which led to the inference of a total of 496,930 GCCs, with a reduction rate of 79% (Supp Table 7-1B). The numbers of identified cluster GCCs for each category are shown in Supplementary Table 7-1.



Supplementary Figure 7-1. Radar plots used to determine the best MCL inflation value for the partitioning of the K into cluster components. The plots were built using a combination of five variables: 1=proportion of clusters with one component and 2=proportion of clusters with more than one member, 3=clan entropy (proportion of clusters with entropy = 0), 4=intra HHblits-Score/Aligned-columns (normalized by the maximum value), and 5=number of clusters (related to the non-redundant set of DAs). (A) Metagenomic dataset. (B) Genomic dataset.

Supplementary Table 7-1. Number of gene clusters, cluster communities and reduction rate shown by functional category.

(A) Metagenomic dataset (MG)

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	К	KWP	GU	EU	Total
Clusters	1,050,166	632,453	1,121,809	135,829	2,940,257
Communities	24,181	64,938	146,100	48,095	283,314
Reduction (%)	97.7	89.73	86.98	64.59	90.36

(B) Genomic dataset (GTDB)

_	K	KWP	GU	EU	Total
Clusters	617,344	136,406	1,525,550	68,202	2,347,502
Communities	52,360	47,203	339,468	57,899	496,930
Reduction (%)	91.52	65.39	77.75	15.11	79.30

Supplementary Note 8 - Gene cluster community validation

The biological significance of the gene cluster communities (GCC) was tested by exploring their distribution within the phylogeny of proteorhodopsin and a set of ribosomal protein families.

Methods

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440 Analysis of the GCC distribution within the proteorhodopsin phylogeny.

We searched the proteorhodopsin (PR) HMM profiles from Olson et al.²¹ against the K and KWP cluster consensus sequences, using the hmmsearch program of the HMMER software (version 3.1b2)²². We filtered the results for alignment coverage > 0.4 and e-value \geq 1e-5. The filtered results were placed in the MicRhoDE PR tree²³ using pplacer²⁴. Then we placed the guery PR sequences into the MicRhode²³ PR tree. We de-duplicated the placed gueries with CD-HIT (v4.6)²⁵ and we cleaned them from sequences with less than 100 amino acids using SEQKIT (v0.10.1) (Shen et al. 2016). Next, we calculated the best substitution model using the EPA-NG modeltest-ng (v0.3.5)²⁶ and we optimized the MicRhoDE PR tree initial parameters and branch lengths using RAxML (v8.2.12)²⁷. Afterward, we incrementally aligned the guery PR sequences against the PR tree reference alignment using the PaPaRA (v2.5) software²⁸. We divided the query alignment and the reference alignment using EPA-NG -split v0.3.5. We combined the PR tree with the related contextual data and the tree alignment, into a phylogenetic reference package using Taxtastic (v0.8.9), and we placed the PR query sequences in the tree using pplacer (v1.1.alpha19-0-g807f6f3)²⁴ with the option -p (-keep-at-most) set to 20. We grafted the PR tree with the query sequences using Guppy, a tool part of pplacer. 3. As the last step, we assigned the PR Supercluster affiliation to the guery sequence, transferring the annotation of its closest relative in the MicRhoDE tree²³ the R packages APE v5.3 and phanghorn v2.5.3²⁹.

Furthermore, we aligned the query sequences annotated as viral to the six viral PRs from Needham et al. 2019³⁰, using Parasail³¹ (-a sg_stats_scan_sse2_128_16 -t 8 -c 1 -x). We then built a sequence similarity network (SSN) using the sequence similarity values to weight the graph edges.

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464 Analysis of standard and high-quality GCCs distribution within ribosomal protein families.

As an additional evaluation, the distributions of standard GCCs and HQ GCCs within ribosomal protein families were investigated and compared. The ribosomal proteins used for the analysis were obtained combining the set of 16 ribosomal proteins from Méheust et al.³² and those contained in the collection of bacterial single-copy genes of Anvi'o³³, that can be

469 downloaded from 470 (https://github.com/merenlab/anvio/blob/master/anvio/data/hmm/Bacteria 71/genes.txt). 471 472 **Results** 473 474 The results of both distribution analyses are shown in Figure 2D and 2C, respectively, and 475 described in the main text. 476 We found 63 of the viral genes placed in the PR tree showing an average similarity of 50% with the viral PR of Needham et al. 30 (Suppl. Table 8-1). Additionally, we found two genes 477 (from two TARA samples: TARA 093 SRF 0.22-3 and TARA 145 SRF 0.22-3) sharing a 478 479 similarity of 100% with one of the Needham et al. PRs (ChoanoV2 VirRyml 1). These 480 genes, however, were not placed in the PR tree. 481 482 Supplementary Table 8-1. Sequence similarity values between viral genes and Needham 483 et al. viral PRs. (Supplementary tables 1.xlsx).

Supplementary Note 9 - HMM-HMM homology network weighting metrics

Validation of the edge weight metrics used for the gene cluster homology network community inference.

Methods

A critical step in the gene cluster community (GCC) inference relies on the determination of the edge weights for the GC HMM-HMM network. We tested two possible metrics to weight the GC homology network resulting from the all-vs-all HMM GC comparison with HHblits¹⁷: (1) the ratio between the HHblits score and the number of aligned columns (*HHblits-Score/Aligned-columns*), metric chosen in this paper; (2) the maximum(HHblits-probability x coverage), weight used in Méheust et al. (2019) ³². In addition, we tested the two different metrics using the ribosomal protein families as reference. For this second test, we filtered the GCCs for those annotated to the 16 ribosomal proteins used in Méheust et al.³², and those contained in the collection of bacterial single-copy genes of Anvi'o³³, which can be downloaded from https://github.com/merenlab/anvio/blob/master/anvio/data/hmm/Bacteria_71/genes.txt. To then compare the two metrics, we used the functions of the R package *aricode* (https://github.com/jchiquet/aricode)³⁴, which allow comparisons between clustering

Results

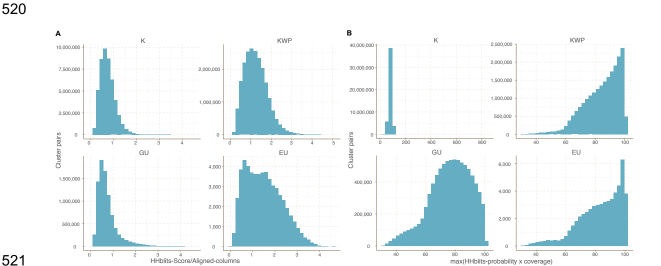
methods.

The results of the test of the different HHblits metrics used to weight the GC homology network are shown separately in Supplementary Figure 9-1 and the comparison in Supplementary Figure 9-2. Both metrics present a very different behavior (Supplementary Figure 9-1), the metric used in Méheust et al. is rescaling the *HHblits-probability* (Supplementary Figure 9-2). While the *HHblits-probability* is useful for deciding if two HMMs are reliable homologs, it is not suitable for measuring similarities due to its dependence on the length of the alignment. On top of this, we can see how the *HHblits-Score/Aligned-columns* values present a similar and more homogenous distribution in all four categories, being more suitable for the MCL clustering.

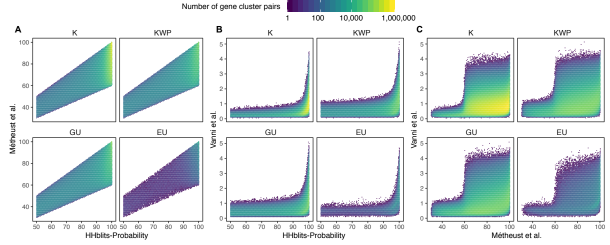
being more suitable for the MCL clustering.

Overall, our approach generated fewer GCCs, as can be observed in Supplementary Figure 9-3. Our clustering was found closer to the "ground truth" represented by the ribosomal protein families compared to the partitioning proposed by Méheust et al. The results from the

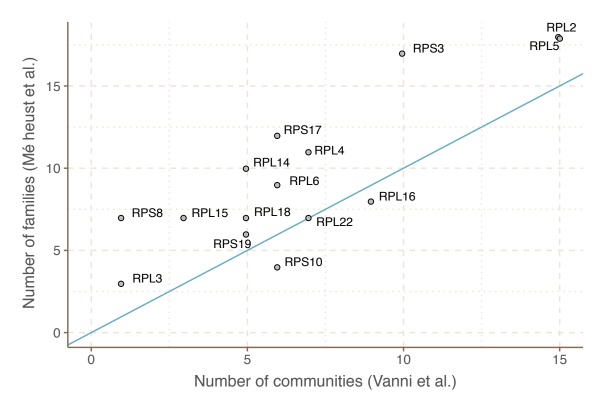
comparison between the two clustering approaches and the ribosomal protein reference are reported in Supplementary Table 9-1.



Supplementary Figure 9-1. Cluster pairs distribution based on the metrics used to weight the gene cluster HMM-HMM homology network. (A) HHblits-Score/Aligned-columns (Vanni et al.). (B) maximum(HHblits-probability x coverage) (Méheust et al.).



Supplementary Figure 9-2. Determination of the edge-weight metrics for the GC HMM-HMM homology network. We tested the metrics used in Méheust et al. and this paper (Vanni et al.). The correlations between metrics are shown per functional category. The metric used by Méheust et al. corresponds to the maximum(HHblits-probability x coverage). The metric applied in this manuscript is *HHblits-Score/Aligned-columns*. (A) Comparison between the metric of Méheust et al. and the HHblits-Probability. (B) Comparison between the metric used in this manuscript and the HHblits-Probability. (C) Comparison between the metric used in this manuscript and the metric of Méheust et al.



Supplementary Figure 9-3. Agreement between the number of communities within ribosomal protein families between our approach and the one described in Méheust et al.

Supplementary Table 9-1. Measures of similarity between the community inference approach proposed in this paper, the one used in Méheust et al. and the "ground truth" represented by the ribosomal protein families.

	Vanni et al. vs Meheust et al.	Vanni et al. vs ribosomal families	Meheust et al. vs ribosomal families
ARI	0.915	0.944	0.906
AMI	0.928	0.916	0.878
NVI	0.101	0.0858	0.124
NID	0.0717	0.0841	0.122
NMI	0.928	0.916	0.878

Note: ARI=Adjusted Rand Index; AMI=Adjusted Mutual Information; NVI=Normalized Variation Information; NID=Normalized Information Distance; NMI=Normalized Mutual Information.

Supplementary Note 10 - EU gene cluster in metagenomeassembled genomes

Metagenome-assembled genomes (MAGs) as a resource to contextualize the environmental unknown gene clusters and cluster communities.

Overall, the MG+GTDB integrated cluster dataset contains 204,031 EU gene clusters (GCs) (grouped in 103,195 cluster communities (GCCs)). The EUs are divided into 127,032 metagenomic, 70,470 genomic, and 9,024, both metagenomic and genomic GCs. The last two subsets contain 52,231 (26%) EU found in GTDB metagenome-assembled genomes (MAGs). To test whether we could also place the subset of metagenomic EU in the context of MAGs, we searched the GCs of this set against the manually curated TARA Ocean MAG collection from Delmont et al. ³⁵.

In addition, we deepened the investigation of the metagenomic EU subset, focusing on the GCCs found broadly distributed in metagenomes according to the results of Levin's niche breadth analysis (Fig. 4). The details of the metagenomic EU analysis are described below.

Methods

We searched the metagenomic EU GCs HMM profiles, obtained from the cluster MSA using the *hhmake* program of the *HH-SUITE*⁷, against the set of 957 high-quality MAGs binned from the TARA Ocean prokaryotic dataset³⁵. We performed the sequence-profile search using the MMSeqs2 *search* program ¹³, using -e 1e-20 --cov-mode 2 -c 0.6. We filtered the results to keep the hits within 90% of the log10(best-e-value). We applied a majority vote function to retrieve the consensus category for each hit. Then, we sorted the results by the smallest e-value and the largest query and target coverage to keep only the best hits. We then filtered the search results focusing on the broadly distributed EU GCs and GCCs. We retrieved MAG contigs containing the EU GCs and GCCs from the Anvi'o MAG profiles using the program *anvi-export-gene-calls* from Anvi'o v4³³. We functionally annotated the contigs searching their genes against the Pfam database (v. 31.0)⁶, using the *hmmsearch* program from the *HMMER* package (version: 3.1b2)²², and complementing the search using *Prokka*³⁶ in metagenomic mode. We then selected the contig with the lowest percentage of hypothetical proteins, and we extracted a region of 1kb surrounding the genes mapping to the EU GCCs.

Results

We found a total of 5,420 EU clusters homologous to 7,661 genes in the 691 TARA MAGs. These EU clusters belong to 4,365 GCCs. We kept only the 71 EU GCCs that showed a broad distribution in TARA samples. These GCCs contained 3,119 clusters and were found in 83 different TARA MAGs. Next, we examined the genomic neighborhood of the broad distributed EU on the MAG contigs. Investigating the genomic neighborhood can lead to the inference of a possible function of the EU. We selected the MAG most enriched with broadly distributed EU, resulted being the Atlantic North-West which in "TARA_ANW_MAG_00076" (Supp. Fig. 10-1A). This MAG contains 23 EU (0.3%) of its genes. It belongs to the bacterial order of Flavobacteriales. Of its 1,283 contigs, 317 include at least one EU. We functionally annotated these contigs with Prokka (and Pfam). Then, we sorted the contigs based on the proportion of genes annotated to hypothetical or characterized proteins, as shown in Supplementary Figure 10-1B. The presence of genes of known function around the EU contributes to prove that these unknown genes are part of a real contig, and possibly an operon. Therefore, we selected for exploration, the contigs with the highest proportion of characterized genes, "TARA ANW MAG 00076 000000000672", with 7 characterized genes out of a total of 13 annotated genes. The contig with the second least amount of hypothetical proteins was "TARA ANW MAG 00076 000000001247", which contained nine characterized genes out of 20. The contig "TARA ANW MAG 00076 000000000672" is shown in Supplementary Figure 10-1C and highlighted in red are the two predicted genes with significant homology to the EU GCs, members of the broadly distributed EU GCCs eu com 769 and eu com 5081. Within their genomic neighborhood, we observe genes relating to nucleotide metabolism, DNA repair and phosphate regulation/sensing, including dUTPase, phoH and protein RecA. Gene placement in prokaryotic genomes is not random. Genes are grouped to increase transcriptional efficiency to respond to stimuli in the environment. Therefore, we can hypothesize that these EU have functions related to their neighboring genes.

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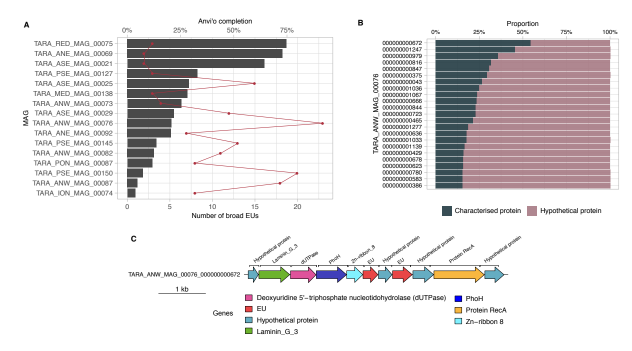
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Supplementary Figure 10-1. (A) EU mapping on TARA MAGs results. Histogram of TARA MAG percent completeness (checkM). The red line represents the number of EU found in the MAGs. (B) Contigs from TARA MAGs TARA_ANW_MAG_00076 in descending order of highest proportion of non-hypothetical gene content. (C) EU communities in the context of a MAG contig. Contig genomic neighborhood around two potential EU communities.

Supplementary Note 11 - Singletons effect on the coding sequence space diversity

Insights into the metagenomic and genomic singletons and their influence on the gene cluster rate of accumulation.

Singletons represent a significant fraction in both the metagenomic (60%) and genomic (55%) datasets. Although we discarded them from the primary analyses presented in this paper, we analyzed their composition in terms of functional categories. The analysis steps are described for the metagenomic singletons in Supplementary Note 1, and, after the integration, we applied the same steps to the genomic singletons (Supp. Table 11-1). As shown in Supp. Note 1, the metagenomic singletons are highly represented by EU genes, while in the genomes we observed the majority of the singletons shared between GU and EU. In general, the singletons are characterized by a high percentage of genes of unknown function.

We tested the singletons role in the rate of accumulation of GCs and GCCs as a function of the number of genomes and metagenomes, as shown in Figure 3C and 3D (to be compared with Supp. Fig. 5A and 5B). For the metagenomic collector curves, we included only the singletons with a sample abundance of 8.36. This value corresponds to the mode sample abundance of the set of metagenomic singletons that became clusters with more than ten genes after the integration of the genomic data.

We observed that, excluding the 19,911,324 singletons from the metagenomic dataset, the accumulation curves of the GCs flatten and approach a plateau. The same effect is observed, excluding the set of 5,558,438 singletons from the genomic dataset (Supp. Fig. 5B; Supp Table 11-2).

Supplementary Table 11-1. Number of genomic singletons per functional category.

	К	KWP	GU	EU
Genes	473,460	896,127	2,528,370	1,660,481

Supplementary Table 11-2. Minimum slope values for the collector curves.

(A) Excluding singletons. In parenthesis, the number of genomes or metagenomes for the first occurrence of slope < 1

Gene Clusters Gene cluster Communities
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_	metaG	GTDB	metaG	GTDB
Known	209.235	6.556	0.1344 (440)	0.07 (15,120)
Unknown	374.5147	5.851	0.1375 (600)	0.621 (27,690)

(B) Including singletons (with a mode abundance in the samples of 8.36).

	Gene Clusters			
	metaG GTDB			
Known	1329.489	66.063		
Unknown	4843.570	158.891		

Supplementary Note 12 - Coverage of external databases

Analysis of the coverage, by our metagenomic dataset, of seven external microbial gene and gene cluster datasets.

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Methods

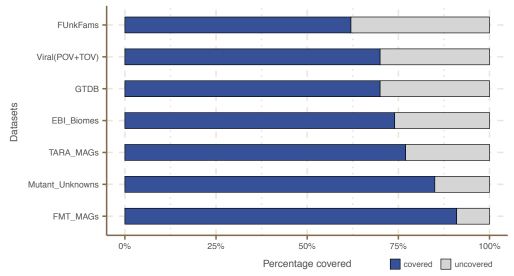
- We searched seven different state-of-the-art databases against our dataset of cluster HMM profiles. The different profile searches were all performed using the MMSeqs2 (version 8.fac81) search program ¹³, setting an e-value threshold of 1e-20 and a query coverage threshold of 60% (-e 1e-20 --cov-mode 2 -c 0.6). We kept the hits within 90% of the log10(best-e-value). Then we applied a majority vote function to retrieve the consensus functional category for each search hit. In the end, the results were sorted by the lowest e-value and the largest query and target coverage to keep only the best hits.
- 661 We applied the described method to the following datasets: the Families of Unknown Functions (FUnkFams) (61,970 genes) ³⁷, the Pacific Ocean Virome (POV) (4,238,638 662 genes) ³⁸ and the Tara Ocean Virome (TOV) (6,642,187 genes) ³⁹. The Genome Taxonomy 663 Database (GTDB) (93,723,190 archaeal and bacterial genes) ¹⁵. The MGnify proteins from 664 the EBI metagenomics database (release 2018 09)⁴⁰ (843,535,611 genes). The manually 665 curated collection of 957 MAGs from TARA metagenomes ³⁵ (TARA MAGs) (2,288,202 666 667 genes), and the one made of 92 MAGs, from the fecal microbiota transplantation study (FMT MAGs) of Lee et al. 41 (188,983 genes). And also the collection of unannotated genes with 668 mutant phenotypes identified in Price et al. 2018 42 (37,684 mutant genes). 669

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Results

- 672 We found our metagenomic GCs in all the main biomes defined by EBI metagenomics (Supp. Fig. 6), with an overall coverage of 74% of the MGnify peptides (Supp. Fig. 12-1). 673 674 Our GCs also covered 62% of the FUnkFam genes of Wyman et al.; 70% of the GTDB 675 genes; and 85% of the gene tested for mutant phenotypes in Price et al.. We also covered 676 50% of the Pacific Ocean Virome proteins, and 77% of the TARA Ocean Virome proteins, for 677 overall coverage of 70% of the selected viral proteins. The majority of genes from both the 678 FMT MAGs of Lee et al. and the TARA MAGs of Delmont et al., were found homologous to 679 genes in our dataset (91% and 77% respectively). With the only exception of the FUnkFams, 680 and the mutant genes, for which we did not find any homology to EU GCs, the other 681 datasets reported homologies to clusters from all four functional categories.
- Moreover, we found that 20% of the Wyman et al FUnkFams and 44% of the unknowns included in the RB-TnSeq experiments by Price et al., 2018 belong to the known CDS-space (Supp. Table 12-1).





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Supplementary Figure 12-1. Coverage of external datasets. The barplot is showing the proportion of covered genes in each of the seven datasets that were screened against the metagenomic set of clusters' HMM profiles.

Supplementary Table 12-1. Re-classification of the unknowns identified in Wyman et al and Price et al.

Study	Original unknown set	Covered fraction	Found as known	Found as unknown
Wyman et al.	61,970	38,174	12,366	25,808
Price et al.	49,736	33,016	21,967	11,049

Supplementary Note 13 - Archaea gene cluster phylogenomic analysis

Gene clusters phylogenetic analysis - results for the archaeal genomes.

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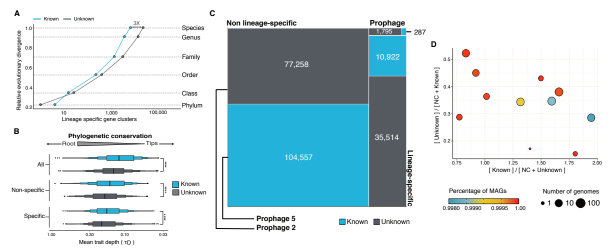
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In the main text are shown the results for the gene clusters (GCs) phylogenetic analyses (clusters phylogenetic conservation and specificity) for the GTDB bacterial genomes. The same methods/analyses were applied for the archaeal genomes, and the results are presented here.

Out of the 230,340 GCs found in GTDB archaeal genomes, we identified 48,518 lineagespecific GCs (precision and sensitivity both ≥95%⁴³). As seen for the Bacteria in Figure 5A, the number of known and unknown archaea lineage-specific GCs increases with the Relative Evolutionary Distance¹⁵, with the differences between the known and the unknown fraction starting to be evident at the Family level (Supp. Fig. 13-1A). The number of unknown lineage-specific GCs for Family, Genus and Species are 2,937, 12,966 and 21,002 respectively (Supp. Tale 13-1). A total of 34,893 GCs were phylogenetically conserved (P < 0.05), where 19,693 were known GCs and 15,200 were unknown GCs. Overall, the unknown GCs are more phylogenetically conserved than the known GCs (Supp. Fig. 13-1B, p < 0.0001). However, considering only the lineage-specific clusters, we observe the opposite, the unknown GCs result in less phylogenetically conserved (Supp. Fig. 13-1B). The GTDB archaeal genomes were also screened for prophages. In total, we identified 2,082 lineagespecific GCs in prophage genomic regions, and 86% of them resulted in clusters of unknown function (Supp. Fig. 13-1C). To identify archaeal phyla enriched in unknown GCs, we partitioned the phyla based on the ratio of known to unknown GCs and vice versa (Supp. Fig. 13-1D). We observed the same pattern found for bacterial phyla in Figure 5D, where the archaeal phyla with a larger number of MAGs are enriched in GCs of unknown function (Supp. Fig. 13-1D).



Supplementary Figure 13-1. Phylogenomic exploration of the unknown coding sequence space in Archaea. (A) Distribution of the lineage-specific gene clusters by taxonomic level. Lineage-specific unknown gene clusters are more abundant at the lower taxonomic levels (genus, species). (B) Phylogenetic conservation of the known and unknown coding sequence space in 1,569 archaeal genomes from GTDB. We calculated the mean trait depth (τ_D) with the consenTRAIT algorithm and the lineage specificity using the F1-score approach from 43 . We observe differences in the conservation between the known and the unknown coding sequence space for lineage- and nonlineage-specific gene clusters (paired Wilcoxon rank-sum test; all p-values < 0.0001). (C) The majority of the lineage-specific clusters are part of the unknown coding sequence space, being a small proportion found in prophages present in the GTDB genomes. (D) Known and unknown coding sequence space of the 1,569 GTDB archaeal genomes grouped by archaeal phyla. Phyla are partitioned based on the ratio of known to unknown gene clusters and vice versa from the set of genomes. Phyla enriched in Metagenomic assembled genomes (MAGs) have a higher proportion in gene clusters of unknown function.

Supplementary Table 13-1. Number of phylogenetic conserved and lineage-specific GCs in the GTDB archaeal phylogeny. (Supplementary tables 1.xlsx).

Supplementary Note 14 - *Cand.* Patescibacteria lineagespecific gene clusters analysis

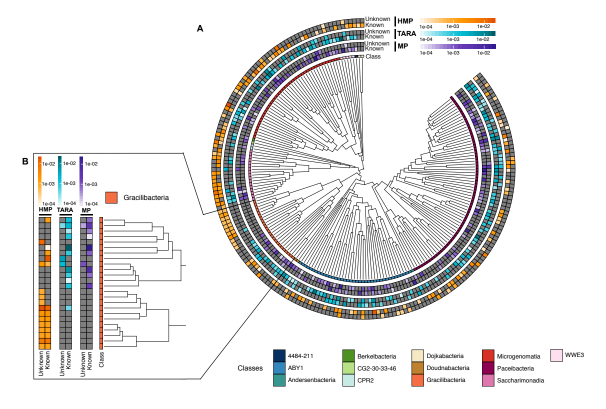
The investigation of the lineage-specific clusters was deepened, focusing on those specific to the Cand. Patescibacteria phylum (former Candidate Phyla Radiation-CPR) and analyzing their cluster distribution in both the Human and marine (TARA and Malaspina) metagenomes.

We found two GU clusters phylum-specific, and a total of 54,343 clusters of unknown function, lineage-specific within the *Cand*. Patescibacteria phylum (Supp. Table 14-1). The majority of this phylum members are particularly poorly understood microorganisms, mostly due to undersampling and the incompleteness of the available genomes. Therefore, we decided to investigate the distribution in the human and marine (TARA and Malaspina) metagenomes of all the clusters lineage-specific inside the *Cand*. Patescibacteria phylum (Supp. Fig. 14-1A).

We chose to have a closer look at the class of *Gracilibacteria*, which shows to be present in both human and marine environments. The first genome for this class was retrieved in a hydrothermal vent environment in the deep sea⁴⁴. The same organisms were then also identified in an oil-degrading community ^{44,45} and as a part of the oral microbiome⁴⁶. As shown in Supplementary Figure 14-1B, we found both known and unknown clusters lineage-specific to this class, distributed in human and marine metagenomes. Among these clusters, we observed cases of environment specificity. For instance, three clusters of unknowns were found exclusive to HMP samples. These clusters could be proposed as novel targets for human-health study since *Gracilibacteria* was found enriched in healthy individuals⁴⁶. We also observed lineage-specific clusters of known and unknown functions specific to the marine environment.

Supplementary Table 14-1. Number of lineage-specific clusters within the *Cand.* Patescibacteria phylum, at different taxonomic levels, subdivided by cluster categories.

Taxonomic level	K	KWP	GU	EU
Phylum	1	0	2	0
Class	11	0	6	0
Order	41	1	104	0
Family	452	9	1,443	13
Genus	625	98	6,649	338
Species	4,116	818	42,710	3,078



Supplementary figure 14-1. Cand. Patescibacteria metagenomic lineage-specific clusters. (A) Phylogenetic tree of Cand. Patescibacteria genera, colored by classes. The heatmaps around the tree show the proportion of lineage-specific gene clusters of knowns and unknowns in the metagenomes from TARA, Malaspina and the HMP. (B) Metagenomic lineage-specific clusters in the class of *Gracilibacteria*.

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