# SCIENTIFIC DATA



ANALYSIS

## **OPEN** Unravelling the diversity of magnetotactic bacteria through analysis of open genomic databases

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Magnetotactic bacteria (MTB) are prokaryotes that possess genes for the synthesis of membranebounded crystals of magnetite or greigite, called magnetosomes. Despite over half a century of studying MTB, only about 60 genomes have been sequenced. Most belong to Proteobacteria, with a minority affiliated with the Nitrospirae, Omnitrophica, Planctomycetes, and Latescibacteria. Due to the scanty information available regarding MTB phylogenetic diversity, little is known about their ecology, evolution and about the magnetosome biomineralization process. This study presents a large-scale search of magnetosome biomineralization genes and reveals 38 new MTB genomes. Several of these genomes were detected in the phyla Elusimicrobia, Candidatus Hydrogenedentes, and Nitrospinae, where magnetotactic representatives have not previously been reported. Analysis of the obtained putative magnetosome biomineralization genes revealed a monophyletic origin capable of putative greigite magnetosome synthesis. The ecological distributions of the reconstructed MTB genomes were also analyzed and several patterns were identified. These data suggest that open databases are an excellent source for obtaining new information of interest.

#### Introduction

The amount of data obtained from genome and metagenome sequencing has been sharply increasing for the last several years<sup>1</sup>. These data are kept in open databases, such as the widely used NCBI<sup>2</sup> and IMG<sup>3</sup> databases. In the case of IMG, the number of entries for metagenomic data greatly exceeds that for genomic ones<sup>3</sup>. In most cases, scientists use only a part of the sequencing information uploaded to the databases, leaving large quantities of information essentially unanalyzed. This gives the possibility that the obtained data may contribute to other studies and shorten the time and efforts of other scientists. In the present study, data stored in open genomic and metagenomic databases were used to search for magnetosome biomineralization genes related to magnetotactic bacteria (MTB).

The MTB are a group of organisms characterized by the ability to synthesize magnetosomes, which are crystals of magnetite (Fe<sub>3</sub>O<sub>4</sub>) or greigite (Fe<sub>3</sub>S<sub>4</sub>) enveloped by a lipid membrane<sup>4</sup>. These crystals can be applied in medicine as contrast agents for MRI<sup>5</sup> and for treating tumors using magnetic hyperthermia<sup>6</sup>, and they are also of great interest in geology<sup>7-9</sup> and astrobiology<sup>10</sup>. The synthesis of magnetosomes is controlled by the magnetosome gene cluster (MGC), previously called the magnetosome island or MAI. The MGC comprises genes that control magnetosome biosynthesis and that determine magnetosome morphology and chemical composition. The MGCs are unique and are associated only with MTB. The genes essential to the biomineralization process are called mam (magnetosome membrane) genes. Nine of them (mamA, -B, -M, -K, -P, -Q, -E, -O, and -I), are present in all MGCs<sup>11,12</sup>. In addition to the mam genes, genes specific to certain groups may also occur; for instance, mad genes are found in MTB from the Deltaproteobacteria and Nitrospirae, while man genes are present only in the Nitrospirae<sup>13</sup>.

At present, only about 60 MTB genomes are known, and most are affiliated with the phyla Proteobacteria, Nitrospirae, and Ca. Omnitrophica. Recently, MTB genomes associated with Latescibacteria<sup>14</sup> and Planctomycetes<sup>12</sup> have been found in open databases, implying that these databases could contain substantial amounts of new information about MTB.

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Organism	Phylum/Class	Accession in NCBI/IMG	Size (bp)	Scaffolds (no.)	GC (%)	N50 (bp)	CheckM completeness (%)	CheckM contamination (%)
Magnetovibrio sp. ARS8 <sup>51,83</sup>	Alphaproteobacteria	GCA_002686765.1	2019305	197	59.64	10605	62.87	1.00
Elusimicrobia bacterium NORP122 <sup>64,84</sup>	Elusimicrobia	GCA_002401485.1	2913226	191	54.93	19622	74.06	1.82
Unclassified Nitrospina Bin 25 <sup>45,114</sup>	Nitrospinae	2651870060	4158979	431	37.69	11956	92.31	4.27
Planctomycetes bacterium SCGC_JGI090-P21 <sup>115</sup>	Planctomycetes	2264265205	1230646	242	49.20	12722	38.87	2.19

Table 1. Characteristics of genomes with MGCs obtained from the NCBI and IMG database genomic data.

To date, due to the lack of sufficient amounts of genomic data, little is known about the origin and evolution of MGCs<sup>15</sup>. Thus, additional investigations are needed to determine the mono- or polyphyletic origin of the MGCs, their evolutionary history, and whether the original MGCs were responsible for magnetite or greigite biomineralization.

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This article describes the first large-scale search of magnetosome biomineralization genes in open genomic and metagenomic databases. Bioinformatics analysis of the search results allowed new MTB genomes to be obtained. Taxonomic assignments for the studied genomes provided the first evidence of their affiliation to new for MTB taxonomic ranks, including three new phyla. These results significantly expanded the knowledge of MTB diversity. The analysis of the ecological distribution of the reconstructed MTB genomes helped to identify several new patterns. Further comparative analysis of MGCs and marker genes of studied genomes allowed new data to be obtained concerning the origin and evolution of magnetosome biomineralization genes.

#### Results

The search for magnetosome biomineralization genes in open databases. The search for MTB genomes in open databases was guided by detecting MGCs unique to magnetotactic bacteria. Unfortunately, MGC sequences are not annotated as magnetosomal in open databases. This necessitated the use of previously known sequences of MGCs as search targets. The search was further complicated by the low identity values between the sequences of the same MGC gene in different MTB taxonomic groups. To cover the maximum number of new MTB representatives, MGC protein sequences were drawn from all known taxonomic groups where MTB were found previously. For this purpose, a database was created of known MGC protein sequences [2-43] (Supplementary Table S1). The database included 67 MGCs from *Proteobacteria*, *Nitrospirae*, *Ca*. Omnitrophica, *Latescibacteria*, and *Planctomycetes*. The sequences of nine Mam proteins present in all MGCs were used to conduct BLASTp with genomic data from the NCBI and IMG databases. This resulted in the detection of four new genomes containing magnetosome biomineralization genes (Table 1, Supplementary Table S2).

The use of all nine Mam proteins in metagenomic databases is complicated by the fact that much more data is kept in metagenomic than in genomic ones. To hasten the search process, one Mam protein out of nine common ones that met the required parameters was chosen for further BLAST analysis. The first chosen parameter was the identity between sequences from different taxonomic groups in each protein. The low values of these identities allowed exclusion of MamE, MamO, and MamP proteins from the analysis. The remaining MamA, -B, -M, -K, -I, and -Q proteins were assessed for sequences with the highest -ln of e-values, in addition to high identities (Fig. 1a). MamI was the least consistent with these requirements and was not used in further analyses. By contrast, MamK was the most consistent.

Each Mam protein has its homologs in non-MTB that are not involved in the magnetosomes biomineralization process. These homologs should be avoided when searching for MGCs. For this, Mam protein was chosen whose identities and -ln of e-values were significantly varied from these parameters in homologs (Fig. 1b). MamK showed the best result in this case, and its minimum identity and -ln e-value between sequences were 30 and 135, respectively. However, part of homologs had identities and -ln of e-values similar to the values found between Mam protein sequences. These homologs were confirmed not to be Mam sequences by verifying their phylogenetic separation (Fig. 1c). The sequences of each Mam protein formed monophyletic clades, while MamK formed two clades. Despite this, no homologs were observed inside the MamK clades. Based on all the investigated parameter results, the MamK protein sequences were chosen for the MGC gene search in the open databases.

The MamK protein sequences were used for BLAST for 10587 metagenomes from water, terrestrial, engineered, and host-associated ecosystems. The analysis revealed 2798 sequences potentially affiliated with the MamK protein (Supplementary Fig. S1a). Their scaffolds were checked for the presence of other Mam protein sequences. After that, 227 MamK sequences referring to 135 metagenomes were obtained (Supplementary Tables S3 and S4). These and previously known MamK sequences were used to construct a phylogenetic tree (Supplementary Fig. S1b), which revealed that the identified MamK sequences were not closely related to previously known sequences. This assumes that they could refer to taxonomic groups in which MTB were not found before.

**Metagenome binning, phylogenomic inferences, and MGC reconstruction.** The phylogenetic position of genomes to which the MamK sequences belonged was assessed by conducting metagenome binning, and it yielded 14688 metagenome-assembled genomes (MAGs) (Supplementary Table S3). Two metagenomes were also determined to be single-cell amplified genomes (SAGs), so no binning procedures were required for

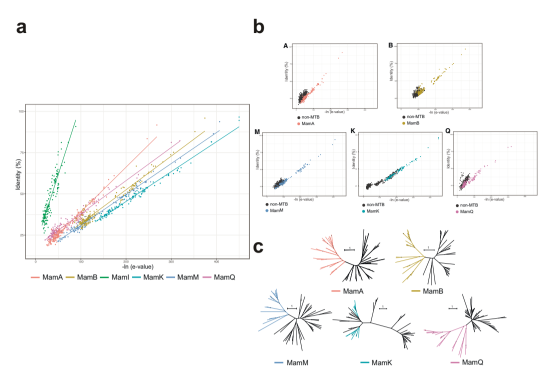


Fig. 1 The choice of Mam protein for further searching for MGCs in open databases. (a) Correlations between  $-\ln$  of e-values (x axis) and identities (y axis) among MamA, -B, -M, -K, -I, and -Q proteins sequences. (b) Correlations between identities and  $-\ln$  of e-values among Mam protein sequences with their homologs. (c) Phylogenetic trees based on investigated sequences. Trees were reconstructed by the maximum-likelihood method with LG+F+I+G4 substitution model. Bootstrap values were calculated based on 1000 resamplings. Bar represents one substitution per 100 amino acid positions.

them. Of all the MAGs obtained in this study, only 140 contained previously detected MamK sequences. For those of the 140 whose completeness was >45% decontamination was conducted. This left 32 MAGs with completeness >45% and contamination <10% that contained MGCs (Table 2, Supplementary Table S6). The phylogenomic affiliations of the obtained MAGs, SAGs, and genomes were then determined, the MGCs genes were reconstructed, and the ecological distributions were studied.

The identification of the phylogenomic position of the studied genomes revealed, for the first time, their affiliation to the phyla *Elusimicrobia*, *Ca*. Hydrogenedentes, and *Nitrospinae* (Supplementary Fig. S2, Supplementary Tables S2 and S5). One genome was affiliated with the phylum *Elusimicrobia* and referred to order UBA1565 in the *Elusimicrobia* class. After MGC reconstruction, the *mamI*, *-B*, *-M*, and *-N* genes were revealed in the investigated genome (Fig. 2). Two MAGs from *Ca*. Hydrogenedentes belonged to the same species (98.70% average nucleotide identity), but they were obtained independently from different metagenomes. These MAGs referred to the GCA-2746185 family in the order *Hydrogenedentiales*. The 16S rRNA gene from the *Ca*. Hydrogenedentes bacterium MAG\_17971\_hgd\_130<sup>44</sup> had 90% similarity with the closest non-MTB *Ca*. Hydrogenedentes bacterium YC-ZSS-LKJ63. All these data confirmed that the obtained binning results were regular and did not represent a computational error. Only *mam* genes were found in the MGCs of the studied genomes.

In the *Nitrospinae* phylum, two MAGs were affiliated with different genera of the order *Nitrospinales*. Their MGCs revealed the presence of *mam* and *mms* (magnetic particle-membrane specific) genes. Samples for the metagenomes of the obtained MAGs were collected from the Gulf of Mexico<sup>45</sup> and Arctic Ocean waters. Non-MTB representatives of this phylum were also detected only in marine habitats<sup>46,47</sup>, indicating that bacteria from the *Nitrospinae* could prefer to inhabit marine environments.

The 14 reconstructed MAGs belonged to different families of *Deltaproteobacteria*. Of the 14, three MAGs were affiliated with the UBA8499 genus in the *Pelobacteraceae* family. In their MGCs, apart from the *mam* and *mad* genes, which are typical for *Deltaproteobacteria*, the *man* genes were detected for the first time. Previously, the *man* genes were associated only with MTB from the *Nitrospirae*. Another two MAGs were affiliated with the *Syntrophobacteraceae* family, where MTB were discovered previously<sup>41</sup>. This is further evidence that binning was conducted correctly and that MTB representatives are indeed present in this family.

Three genomes also belonged to the *Desulfobulbales* order. Of these, the *Deltaproteobacteria* bacterium MAG\_22309\_dsfv\_022<sup>48</sup> contained *man3* gene in addition to the *mam* and *mad* genes, thereby confirming the routine presence of *man* genes in *Deltaproteobacteria*. A further four MAGs were related to the NaphS2 family in the *Desulfatiglanales* order. Analysis of their MGCs revealed genes responsible for putative greigite magnetosome synthesis. Metagenomic samples of the studied genomes were obtained from marine sediments, as well as all other known non-MTB genomes of this family<sup>49,50</sup>.

Ca. Hydrogenedentes bacterium   AG. Hydrogenedentes   3300017963   3018788   288   60.18   11662   71.11   1.46   1.46   Ca. Hydrogenedentes bacterium   AG. Hydrogenedentes   3300017971   2683901   240   60.43   12541   60.01   1.16   1.16   Deltaproteobacteria bacterium   AG. Hydrogenedentes   3300017971   2683901   240   60.43   12541   60.01   1.16   1.16   Deltaproteobacteria bacterium   AG. 0134_naph_006 <sup>86,136</sup>   Deltaproteobacteria   3300000134   1498667   692   49.54   2676   60.69   3.87   MAG. 0021_aph_0106 <sup>86,136</sup>   Deltaproteobacteria bacterium   Deltaproteobacteria bacterium   AG. 0021_aph_0106 <sup>86,136</sup>   Deltaproteobacteria bacterium   AG. 00792_naph_016 <sup>86,136</sup>   Deltaproteobacteria bacterium   AG. 09788_naph_3 <sup>709</sup>   Deltaproteobacteria bacterium   MAG. 09788_naph_3 <sup>709</sup>   Deltaproteobacteria bacterium   MAG. 1978_0978_naph_13 <sup>709</sup>   Deltaproteobacteria bacterium   MAG. 1978_0978_naph_3 <sup>709</sup>   Deltaproteobacteria bacterium   AG. 1978_0978_naph_3 <sup>709</sup>   Deltaproteobacteria bacterium   AG. 1979_0978_naph_26 <sup>86</sup>   Deltaproteobacteria   3300015370   3868622   334   48.42   14397   89.68   5.59   Deltaproteobacteria bacterium   MAG. 1979_098_nab_26 <sup>87</sup>   Deltaproteobacteria   3300017999   2777907   276   53.10   17193   62.13   5.10   Deltaproteobacteria bacterium   AG. 1979_096_snb_20 <sup>88</sup>   Deltaproteobacteria   3300017996   169108   454   53.11   4033   50.53   2.33   Deltaproteobacteria bacterium   AG. 2016_096_096_096_096_096_096_096_096_096_09
Deltaproteobacteria bacterium   Deltaproteobacteria   330000792   3032840   499   49.74   11269   89.28   5.86
Deltaproteobacteria bacterium   MAG_00134_naph_006 <sup>Ma159</sup>   Deltaproteobacteria   3300000134   1498667   692   49.54   2676   60.69   3.87
Deltaproteobacteria bacterium   MAG_00241_naph_010 <sup>07,139</sup>   Deltaproteobacteria   3300000241   1547003   324   49.45   6761   55.59   2.41   Deltaproteobacteria bacterium   MAG_00792_naph_016 <sup>08,19</sup>   Deltaproteobacteria bacterium   MAG_00792_naph_016 <sup>08,19</sup>   Deltaproteobacteria bacterium   Deltaproteobacteria bacterium   MAG_07978_naph_0797   Deltaproteobacteria bacterium   Deltaproteobacteria bacterium   Deltaproteobacteria   330000788   899797   137   47.24   7579   49.08   0.97   MAG_07978_naph_0797   Deltaproteobacteria bacterium   Deltaproteobacteria   3300015370   3868622   334   48.42   14397   89.68   5.59   Deltaproteobacteria bacterium   Deltaproteobacteria   3300017929   2777907   276   53.10   17193   62.13   5.10   Deltaproteobacteria bacterium   Deltaproteobacteria   3300017996   1691080   454   53.11   4033   50.53   2.33   Deltaproteobacteria bacterium   Deltaproteobacteria   3300022204   2675335   75   52.74   60141   89.52   0.36   Deltaproteobacteria bacterium   Deltaproteobacteria   3300022309   2902378   66   55.15   78905   91.60   1.79   MAG_22309_ds/v_022 <sup>39</sup>   Deltaproteobacteria   330000150   2847655   486   49.07   8986   98.17   3.96   Gammaproteobacteria bacterium   MAG_00160_gam_00998   Gammaproteobacteria   330000172   2866084   274   48.97   18904   96.95   3.05   Gammaproteobacteria bacterium   MAG_00172_gam_018 <sup>39</sup>   Gammaproteobacteria   330000172   2866084   274   48.97   18904   96.95   3.05   Gammaproteobacteria bacterium   MAG_00182_gam_00998   Gammaproteobacteria   330000188   2672010   567   48.83   6818   95.12   4.19   Gammaproteobacteria bacterium   MAG_00112_gam_108 <sup>39</sup>   Gammaproteobacteria   330000150   293128   507   49.02   8845   95.73   5.34   MAG_20155_gam_000998   Gammaproteobacteria   330000155   2931288   507   49.02   8845   95.73   5.34   MAG_20155_gam_000998   Gammaproteobacteria   330000155   358593   300   52.41   5203   84.82   3.65   350000000000000000000000000000000000
Deltaproteobacteria bacterium   MAG_00792_naph_0168_hiiiiiiiii   Deltaproteobacteria   330000792   3032840   409   49.74   11269   89.28   5.86
Deltaproteobacteria bacterium   MAG_0978B_naph_3798   Deltaproteobacteria   3300009788   899797   137   47.24   7579   49.08   0.97
Deltaproteobacteria bacterium   MAG_15370_dsf6_81 %0,120   Deltaproteobacteria   3300015370   3868622   334   48.42   14397   89.68   5.59   Deltaproteobacteria bacterium   MAG_17929_sntb_263
Deltaproteobacteria bacterium   Deltaproteobacteria   3300017929   277907   276   33.10   17193   62.13   5.10     Deltaproteobacteria bacterium   MAG_17996_sntb_20°2   Deltaproteobacteria   3300017996   1691080   454   53.11   4033   50.53   2.33     Deltaproteobacteria bacterium   Deltaproteobacteria   3300022204   2675335   75   52.74   60141   89.52   0.36     Deltaproteobacteria bacterium   Deltaproteobacteria   3300022309   2902378   66   55.15   78905   91.60   1.79     Deltaproteobacteria bacterium   MAG_22309_dsfv_022*8   Deltaproteobacteria   3300002309   2847655   486   49.07   8986   98.17   3.96     Gammaproteobacteria bacterium   MAG_00150_gam_010°3   Gammaproteobacteria   330000150   2847655   486   49.07   8986   98.17   3.96     Gammaproteobacteria bacterium   MAG_00160_gam_009°3   Gammaproteobacteria   330000172   2866084   274   48.97   18904   96.95   3.05     Gammaproteobacteria bacterium   MAG_00172_gam_018*6   Gammaproteobacteria   330000172   2866084   274   48.97   18904   96.95   3.05     Gammaproteobacteria bacterium   MAG_00184_gam_006°7   Gammaproteobacteria   330000188   2672010   567   48.83   6818   95.12   4.19     Gammaproteobacteria bacterium   MAG_00212_gam_12*8   Gammaproteobacteria   330000215   2931288   507   49.02   8845   95.73   5.34     Magnetococcales bacterium   MAG_00215_gam_020°9   Ca. Etaproteobacteria   330002105   3585593   930   52.41   5203   84.82   3.65   Nitrospinae bacterium   Nitrospinae
MAG_17996_sntb_20 <sup>92</sup>   Deltaproteobacteria   3300017996   1691080   454   53.11   4035   50.53   2.35     Deltaproteobacteria bacterium   MAG_22204_dsfv_001 <sup>93</sup>   Deltaproteobacteria   3300022204   2675335   75   52.74   60141   89.52   0.36     Deltaproteobacteria bacterium   MAG_22309_dsfv_0022 <sup>98</sup>   Deltaproteobacteria   3300022309   2902378   66   55.15   78905   91.60   1.79     Gammaproteobacteria bacterium   MAG_02309_dsfv_022 <sup>98</sup>   Gammaproteobacteria   330000150   2847655   486   49.07   8986   98.17   3.96     Gammaproteobacteria bacterium   MAG_00160_gam_000 <sup>99</sup>   Gammaproteobacteria   330000160   2903803   318   49.10   15339   99.39   4.88     Gammaproteobacteria bacterium   MAG_00172_gam_018 <sup>96</sup>   Gammaproteobacteria   330000172   2866084   274   48.97   18904   96.95   3.05     Gammaproteobacteria bacterium   MAG_00172_gam_018 <sup>96</sup>   Gammaproteobacteria   330000188   2672010   567   48.83   6818   95.12   4.19     Gammaproteobacteria bacterium   MAG_00212_gam_1 <sup>98</sup>   Gammaproteobacteria   330000212   2103212   955   48.40   2901   78.43   5.08     Gammaproteobacteria bacterium   MAG_00215_gam_020 <sup>99</sup>   Gammaproteobacteria   330000215   2931288   507   49.02   8845   95.73   5.34     Magnetococcales bacterium   MAG_00215_gam_020 <sup>99</sup>   Gammaproteobacteria   330002105   3585593   930   52.41   5203   84.82   3.65   Nitrospinae bacterium   Nag_10050000000000000000000000000000000000
MAG_22204_dsfv_001 <sup>93</sup> Deltaproteobacteria         3500022204         26/3535         /3         32./4         60141         89.52         0.38           Deltaproteobacteria bacterium MAG_22309_dsfv_022 <sup>18</sup> Deltaproteobacteria         3300022309         2902378         66         55.15         78905         91.60         1.79           Gammaproteobacteria bacterium MAG_00150_gam_010 <sup>94</sup> Gammaproteobacteria         3300000150         2847655         486         49.07         8986         98.17         3.96           Gammaproteobacteria bacterium MAG_00160_gam_009 <sup>95</sup> Gammaproteobacteria         3300000160         2903803         318         49.10         15339         99.39         4.88           Gammaproteobacteria bacterium MAG_00172_gam_018 <sup>96</sup> Gammaproteobacteria         3300000172         2866084         274         48.97         18904         96.95         3.05           Gammaproteobacteria bacterium MAG_00188_gam_006 <sup>97</sup> Gammaproteobacteria         3300000188         2672010         567         48.83         6818         95.12         4.19           Gammaproteobacteria bacterium MAG_00212_gam_19 <sup>8</sup> Gammaproteobacteria         3300000212         2103212         955         48.40         2901         78.43         5.08           Gammaproteobacterium MAG_002
MAG_22309_dsfv_02248         Dettaproteobacteria         3300022309         29023/8         66         55.15         78905         91.60         1.79           Gammaproteobacteria bacterium MAG_00150_gam_010 <sup>94</sup> Gammaproteobacteria         3300000150         2847655         486         49.07         8986         98.17         3.96           Gammaproteobacteria bacterium MAG_00160_gam_000 <sup>95</sup> Gammaproteobacteria         3300000160         2903803         318         49.10         15339         99.39         4.88           Gammaproteobacteria bacterium MAG_00172_gam_018 <sup>96</sup> Gammaproteobacteria         3300000172         2866084         274         48.97         18904         96.95         3.05           Gammaproteobacteria bacterium MAG_0018_gam_006 <sup>97</sup> Gammaproteobacteria         3300000188         2672010         567         48.83         6818         95.12         4.19           Gammaproteobacteria bacterium MAG_00212_gam_1 <sup>98</sup> Gammaproteobacteria         3300000212         2103212         955         48.40         2901         78.43         5.08           Gammaproteobacteria bacterium MAG_0215_gam_020 <sup>99</sup> Gammaproteobacteria         3300000215         2931288         507         49.02         8845         95.73         5.34           Magnetococcales bacterium MAG_210
MAG_00150_gam_010³4         Gammaproteobacteria         3300000150         2847635         486         49.07         886         98.17         3.96           Gammaproteobacteria bacterium MAG_010160_gam_009°5         Gammaproteobacteria         3300000160         2903803         318         49.10         15339         99.39         4.88           Gammaproteobacteria bacterium MAG_00172_gam_018°6         Gammaproteobacteria         3300000172         2866084         274         48.97         18904         96.95         3.05           Gammaproteobacteria bacterium MAG_0018_gam_006°7         Gammaproteobacteria         3300000188         2672010         567         48.83         6818         95.12         4.19           Gammaproteobacteria bacterium MAG_00212_gam_1°8         Gammaproteobacteria         3300000212         2103212         955         48.40         2901         78.43         5.08           Gammaproteobacteria bacterium MAG_00215_gam_020°9         Gammaproteobacteria         3300000215         2931288         507         49.02         8845         95.73         5.34           Magnetococcales bacterium MAG_21055_mgc_1¹¹00         Ca. Etaproteobacteria         3300021055         3585593         930         52.41         5203         84.82         3.65           Nitrospinae bacterium         Nitrospi
MAG_00160_gam_00995         Gammaproteobacteria         3300000160         2903803         318         49.10         15339         99.39         4.88           Gammaproteobacteria bacterium MAG_00172_gam_01896         Gammaproteobacteria         3300000172         2866084         274         48.97         18904         96.95         3.05           Gammaproteobacteria bacterium MAG_00188_gam_00697         Gammaproteobacteria         3300000188         2672010         567         48.83         6818         95.12         4.19           Gammaproteobacteria bacterium MAG_00212_gam_198         Gammaproteobacteria bacterium MAG_00215_gam_02099         Gammaproteobacteria         3300000212         2103212         955         48.40         2901         78.43         5.08           Gammaproteobacteria bacterium MAG_00215_gam_02099         Gammaproteobacteria         3300000215         2931288         507         49.02         8845         95.73         5.34           Magnetococcales bacterium MAG_21055_mgc_1100         Ca. Etaproteobacteria         3300021055         3585593         930         52.41         5203         84.82         3.65           Nitrospinae bacterium
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MAG_00188_gam_00697         Gammaproteobacteria         3300000188         26/2010         56/         48.83         6818         95.12         4.19           Gammaproteobacteria bacterium MAG_00212_gam_198         Gammaproteobacteria         3300000212         2103212         955         48.40         2901         78.43         5.08           Gammaproteobacteria bacterium MAG_00215_gam_02099         Gammaproteobacteria         3300000215         2931288         507         49.02         8845         95.73         5.34           Magnetococcales bacterium MAG_21055_mgc_1100         Ca. Etaproteobacteria         3300021055         3585593         930         52.41         5203         84.82         3.65           Nitrospinae bacterium         Nitrospinae bacterium         Nitrospinae bacterium         Nitrospinae bacterium         2000000000000000000000000000000000000
MAG_00212_gam_198         Gammaproteobacteria         3500000212         2103212         955         48.40         2901         78.43         3.08           Gammaproteobacteria bacterium MAG_02215_gam_02099         Gammaproteobacteria         3300000215         2931288         507         49.02         8845         95.73         5.34           Magnetococcales bacterium MAG_21055_mgc_1100         Ca. Etaproteobacteria         3300021055         3585593         930         52.41         5203         84.82         3.65           Nitrospinae bacterium         Nitrospinae bacterium         Nitrospinae bacterium         2000000000000000000000000000000000000
MAG_00215_gam_020 <sup>99</sup> Gammaproteobacteria         3500000215         2951268         307         49.02         8645         95.75         3.34           Magnetococcales bacterium MAG_21055_mgc_1 <sup>100</sup> Ca. Etaproteobacteria         3300021055         3585593         930         52.41         5203         84.82         3.65           Nitrospinae bacterium
MÄG_21055_mgc_1 <sup>100</sup>
Nitrospinae bacterium
MAG_09705_ntspn_70 <sup>101</sup>
Nitrospirae bacterium MAG_10313_ntr_31 <sup>102</sup> Nitrospirae         3300010313         1933163         344         35.33         7568         90.20         3.64
Pelobacteraceae bacterium MAG_21601_9_030 <sup>103</sup> Deltaproteobacteria         3300021601         2536371         232         54.11         20074         78.15         8.39
Pelobacteraceae bacterium MAG_13126_9_058 <sup>104</sup> Deltaproteobacteria         3300013126         3576562         72         52.01         83631         91.61         1.29
Pelobacteraceae bacterium MAG_21600_9_004 <sup>105</sup> Deltaproteobacteria         3300021600         3430740         60         51.50         87025         90.32         0.65
Planctomycetes bacterium MAG_11118_pl_115 <sup>106</sup> Planctomycetes         3300011118         3767441         157         48.98         33372         89.44         1.24
Planctomycetes bacterium MAG_17991_pl_60 <sup>107</sup> Planctomycetes         3300017991         1289005         144         49.53         10179         64.20         0.00
Planctomycetes bacterium MAG_18080_pl_157 <sup>108</sup> Planctomycetes         3300018080         3144921         139         48.44         34208         90.91         3.41
Rhodospirillaceae bacterium MAG_01419_mvb_30         Alphaproteobacteria         3300001419         2811682         477         55.72         7268         94.58         4.10
Rhodospirillaceae bacterium   Alphaproteobacteria   3300004806   2085124   309   57.51   8435   87.64   2.12
Rhodospirillaceae bacterium   Alphaproteobacteria   3300005422   2281835   255   61.09   11800   85.45   0.50
Rhodospirillaceae bacterium MAG_05596_2-02_51***         Alphaproteobacteria         3300005596         1831947         329         61.19         6777         76.91         0.25
Rhodospirillaceae bacterium   MAG_06104_tlms_034 <sup>112</sup>   Alphaproteobacteria   3300006104   3186839   353   64.25   13005   89.59   2.53
Rhodospirillaceae bacterium   MAG_22225_2-02_112 <sup>113</sup>   Alphaproteobacteria   3300022225   2547095   147   61.01   26510   91.17   5.22
Ca. Omnitrophica bacterium SCGC AG-290-C17 (SAG) <sup>116</sup> Ca. Omnitrophica         3300015153         1712617         171         48.60         13921         62.84         0.00
Uncultured microorganism SbSrfc.SA12.01.D19 (SAG) <sup>117</sup> Deltaproteobacteria 3300022116 2501480 175 52.60 25257 49.13 0.00

Table 2. Characteristics of reconstructed MAGs with MGCs obtained from the IMG metagenomic data.

In *Alphaproteobacteria*, three MAGs and one genome were related to a 2-02-FULL-58-16 family in the *Rhodospirillales* order. Metagenomic samples of the studied genomes were isolated from marine ecosystems. The other non-MTB genomes of this family were also detected only in marine ecosystems<sup>51</sup>. For the first time, two MAGs containing MGCs were also detected in *Telmatospirillum* genus. Their metagenomic samples were collected from a freshwater bog. *Telmatospirillum siberiense*, the only known representative of this genus, was also isolated from freshwater peat soil<sup>52</sup>. Thus, this group possibly tends to inhabit freshwater ecosystems. Reconstruction of the MGCs revealed *mam* and *mms* genes in the studied MAGs. One MAG was referred to the *Ca*. Etaproteobacteria class. Genomes from this class previously were found in both saline and freshwater habitats<sup>15,31,53</sup>. The obtained MAG clustered with genomes isolated from freshwater environments. The MGC of the recovered MAG revealed a standard gene set inherent to MTB from this class. A further six MAGs were affiliated with the *Gammaproteobacteria*. All of these were sampled from one source and had 100% identity between their genes. Only the *mam* genes were detected in their MGCs.

The *Nitrospirae* phylum was affiliated with one MAG. A metagenomic sample of this phylum was obtained from a hot spring. Previously, other MTB and non-MTB from this phylum were also detected in hot springs<sup>54,55</sup>. Three of the recovered MAGs belonged to the SG8-4 order in the *Phycisphaerae* class of *Planctomycetes*. Apart from the reconstructed MAGs, one SAG was also obtained from the UBA1845 order in *Phycisphaerae* class. The completeness of this SAG was very low (39%), but it was also taken into analyses due to the large number of *mam* genes detected in the MGC. Another detected SAG was affiliated with *Ca*. Omnitrophica and was referred to the GWA2-52-8 family in the *Omnitrophales* order. The MGC of this genome had a set of genes that were specific to all magnetotactic representatives from this phylum.

Reconstruction of the evolutionary pathways for MGCs. The identification of putative genes involved in magnetosome biomineralization allowed investigation of MGC evolutionary pathways. These were analyzed by constructing a phylogenetic tree of concatenated MamABKMPQ sequences ("Mam tree", Fig. 3b) and comparing this tree with one based on 120 single-copy marker gene proteins ("core genome tree", Fig. 3a). Comparative analysis of the MTB position on the trees revealed some incongruences. For instance, the *Deltaproteobacteria* group from "core genome tree" was divided into three subgroups on the "Mam tree." The first subgroup comprised representatives capable of putative greigite magnetosome synthesis, while the other two subgroups included representatives with MGCs for magnetite magnetosome biomineralization. One of the magnetite subgroups included representatives of the *Pelobacteraceae*, *Syntrophia*, and *Desulfurivibrionaceae* families, which clustered with the *Nitrospirae*. According to the "Mam tree" topology, the *man* genes could be assumed to have originated in the *Deltaproteobacteria* and were inherited by the *Nitrospirae* through horizontal gene transfer. The compared trees also indicated vertical inheritance in the *Alpha*- and *Ca*. Etaproteobacteria groups, although the occurrence of horizontal transfer events was previously established in these groups<sup>27,31</sup>. These types of transfers have been confirmed to have occurred recently, which is why they cannot be detected through the tree topology analysis.

A further investigation examined whether MGC originated once or more than once. This was done by adding the Mam protein sequences recovered in this study to previously known Mam protein sequences and their non-MTB homologs and then constructing phylogenetic trees (Supplementary Fig. S3). Analysis of the constructed trees confirmed the previous results showing that all Mam protein sequences, except for MamK, formed monophyletic clades and that these clades did not contain any homolog sequences. This indicates that the MGCs for magnetite and greigite synthesis are likely to have a common origin.

The magnetosome chemical composition in genomes of every phylum where MTB were found for the first time were predicted by counting the phylogenetic distances of the concatenated sequences of six essential Mam proteins (MamA, -B, -K, -M, -P, and -Q) and conducting a principal component analysis (Fig. 4). All values clustered to three groups. First was the group that comprised *Planctomycetes*, and *Latescibacteria*, which are known to have genes for putative greigite magnetosome synthesis<sup>12,14,56</sup>. The NaphS2 family of *Deltaproteobateria*, *Ca*. Hydrogenedentes, *Ca*. Omnitrophica, and *Elusimicrobia* also clustered with this group. The other two groups comprised representatives with magnetite magnetosome synthesis genes. The first magnetite group included *Nitrospinae* and all classes of *Proteobacteria* where MTB were known. The exception was the remaining studied classes of *Deltaproteobateria*, which clustered with the second magnetite group, together with *Nitrospirae*.

#### **Discussion**

This study represents the first large-scale search of magnetosome biomineralization genes in open databases. Bioinformatic analysis of the gathered data almost doubled the number of MTB genomes from the 60 previously known; 4 genomes, 2 SAGs, and 32 MAGs were obtained as a result of this research. Besides, analysis of the database of collected MGC protein sequences revealed MamK as the most appropriate protein for MGC searching in open databases. This finding will allow the use of these putative protein sequences as markers for MTB detection in environmental samples.

This study also provides the first description of magnetosome biomineralization genes in the genomes of *Elusimicrobia*, *Nitrospinae*, and *Ca*. Hydrogenedentes. Non-MTB representatives of *Elusimicrobia* phylum were previously found as free-living<sup>57</sup> and ecto- and endosymbionts<sup>58,59</sup> of multicellular eukaryotes. MTB living symbiotically with eukaryotes have been detected previously<sup>60,61</sup>. Further investigations are needed to solve the enigma of whether MTB from *Elusimicrobia* free-living or symbiotic organisms are.

To date, little is known about Ca. Hydrogenedentes, except for its genome presence  $^{62-64}$ . More is known about Nitrospinae, where one axenic culture was previously described  $^{65}$ . However, these reports do not give an extensive understanding of the capabilities of this phylum's representatives. Thus, the detection of MGCs in genomes that belong to these phyla significantly supplements the knowledge of MTB diversity and evolution, while also providing new information about these phyla.

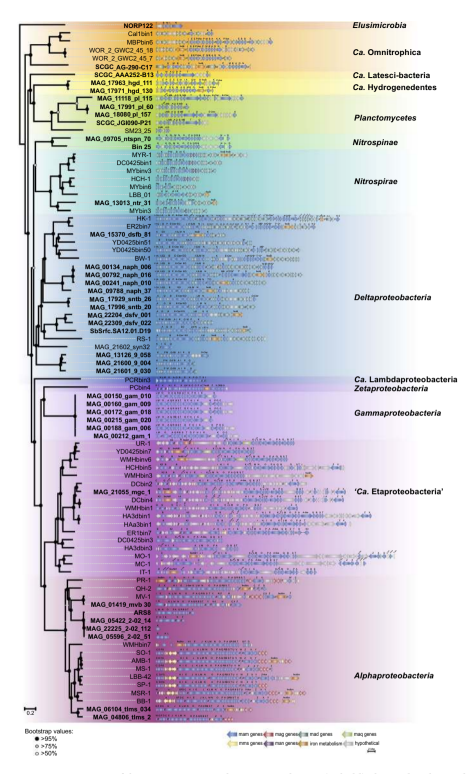


Fig. 2 Comparison of the MGC regions in the MAGs and SAGs (in bold) obtained in this study versus previously known MTB genomes. Full names for MTB strains can be found in Supplementary Table S1.

This work also gives much new information about groups where MTB were previously recognized. For instance, the relatively few genomes were affiliated with *Alpha*- and *Ca*. Etaproteobacteria, while the current belief is that representatives of these classes dominate among MTB in all natural environments<sup>12</sup>. In addition, within the *Alphaproteobacteria* class, the presence of MGCs was discovered for the first time in genomes belonging to the *Telmatospirillum* genus. This may indicate a common origin for magnetosome biomineralization genes among the *Magnetospirillum*, *Magnetospira*, and *Magnetovibrio* genera.

Furthermore, for the first time the presence of *man* genes was revealed in MGCs of the *Deltaproteobacteria*. Previously, these genes were found only in *Nitrospinae*. Whether horizontal gene transfer events occurred

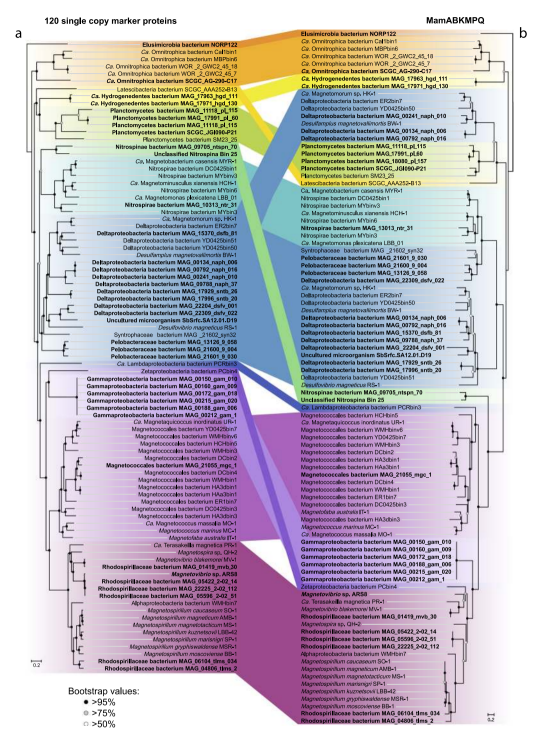


Fig. 3 Maximum-likelihood phylogenomic trees of MTB genomes. Trees were inferred from a comparison of 120 concatenated single-copy marker proteins of MTB genomes (a) and concatenated magnetosome associated protein sequences (MamABKMPQ) (b). Both trees were reconstructed with evolutionary model LG+F+I+G4. Branch supports were obtained with 1000 ultrafast bootstraps. The scale bar represents amino acid substitutions per site.

between representatives of these phylogenetic groups or their MGCs shared a common origin is not known. Further studies are required to determine which possibility is correct.

The genomes with magnetosome biomineralization genes obtained in this study allowed the investigation of the origin and evolution of the MGCs. A comparison of the "core genome" and "Mam" trees revealed clustering of the *Deltaproteobacteria* greigite subgroup sequences with the *Planctomycetes*, *Latescibacteria*, *Ca*. Hydrogenedentes, *Ca*. Omnitrophica, and *Elusimicrobia* phyla. Of these, *Latescibacteria*<sup>14</sup> and *Planctomycetes*<sup>12</sup> were already known to have MGCs for putative greigite synthesis. Note that *Ca*. Omnitrophica was also associated

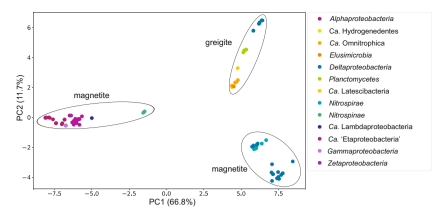


Fig. 4 The prediction of magnetosome chemical composition for phyla in which MTB genomes were found for the first time. Predictions were made using principal component analysis for a maximum-likelihood distance matrix of concatenated Mam protein sequences.

with the greigite subgroup, although it is believed that they biomineralize magnetite magnetosomes  $^{43}$ . Such assumptions are based on Ca. Omnitrophus magneticus SKK-01 however, this genome is highly contaminated (Supplementary Table S1). Thus, further investigations are needed to study Ca. Omnitrophica magnetosome chemical composition.

In addition to all mentioned findings, the latest version of the bacterial tree of life<sup>66</sup>, based on GTDB R04-RS89 reference data (Supplementary Fig. S4) helped to reveal the most ancient phylum in which MTB representatives were known. It was indicated that the *Elusimicrobia* phylum is the most closely related to the last universal common ancestor (LUCA). If the MTB of this phylum are assumed capable of greigite magnetosome synthesis, then greigite MGCs could have appeared much earlier than commonly believed, and the first MTB could have greigite, not magnetite, MGCs. The other phyla with MTB representatives in the vicinity of LUCA are *Ca.* Omnitrophica and *Proteobacteria*, although *Nitrospirae* MTB was previously thought to be the most ancient<sup>40</sup>.

Considering the existing data regarding the presence of horizontal transfer events among MTB and analyzing the discrepancies in "core genome" and "Mam" trees, the proposal could be made that horizontal gene transfers occur much more often than previously thought and are of great importance in MGC evolution.

The genomes obtained in this work require further confirmation by morphological identification. Once confirmed, these data will allow a more thorough study of the contribution of vertical and horizontal gene transfer events with respect to MGC inheritance. The data obtained in the present work will allow the study of the environmental and metabolic preferences of newly discovered MTB genomes, which may become the key to isolating them in axenic cultures. Moreover, a detailed MGC analysis could help to find as yet unidentified genes that are involved in magnetosome synthesis and to reveal much about the biomineralization process.

Generally, in this work, it was shown that MamK is the most appropriate protein for MGCs detecting in open databases. The search results allowed to receive 38 new genomes containing MGCs, that were affiliated to both taxonomic groups where MTB were found before and three new phyla. Thus, received MTB genomes permitted to unravel the MTB diversity and can be used in further MTB studies or in receiving new information about these phyla. Also, a comparison of MTB position on "mam tree" and "core genome tree" helped to reveal signs of putative horizontal gene transfers. This led to assumptions that such MGC transfers could occur with higher frequency and probably play a much more important role in MGC evolution than it was previously thought. Moreover, a proposal was made that the origin of MGC probably is more ancient than it was suggested earlier and possibly was capable of greigite magnetosomes biomineralization rather than magnetite.

Thus, all received data allowed the expansion of knowledge about MTB diversity, ecology, and evolution and has opened up new opportunities for further searches for and investigations of magnetotactic bacteria.

#### Materials and methods

The search for magnetosome biomineralization genes in open databases. The search for magnetosome biomineralization genes was conducted by collecting a database of MGC protein sequences based on currently known MTB genomes (Supplementary Table S1). The search was provided using BLASTp analysis, with identity >30% and e-value >1e<sup>-05</sup>. Searches of the IMG and NCBI genomic databases used sequences of nine essential Mam proteins from different taxonomic groups as targets. The IMG metagenomic database was searched by BLASTp using MamK sequences. The sequences obtained from BLAST analysis were further checked to separate MGC proteins from their homologs. For this, each Mam protein sequence was checked for joint clustering on the phylogenetic trees. The presence of other Mam proteins in the same scaffold provided additional support for choosing those scaffolds for further analysis. The search was conducted in April 2018.

**Genome reconstruction and analyses.** Metagenome assembled genome (MAG) reconstruction was conducted using the Busybee web<sup>67</sup>, Maxbin2<sup>68</sup>, and MyCC<sup>69</sup> with standard parameters. The DAS Tool<sup>70</sup> was used for choosing consensus assemblies for the obtained MAGs. Completeness and contamination values of genomes were obtained using lineage-specific marker genes and default parameters in CheckM v.  $1.0.12^{71}$ . RefineM v.  $0.0.24^{50}$  was used to remove contamination based on taxonomic assignments. This process, called 'decontamination',

involves the classification of obtained genes and scaffolds in each MAG relative to the gene base with a known taxonomic classification. After that, scaffolds with incongruent taxonomic classifications are removed from the MAGs. The quality metrics were assessed using the QUAST<sup>72</sup> tool. The average nucleotide identity (ANI) was calculated using fastANI<sup>73</sup>. The MGCs were determined using local BLAST and comparison with reference sequences of magnetotactic bacteria.

Phylogenetic analyses. Taxonomic assignments for the studied genomes 16S rRNA genes were obtained using the GTDB 16S r89 dataset in IDTAXA<sup>74</sup>. The GTDB-Tk v.0.1.3<sup>75</sup> 'classify\_wf' command was used to find 120 single-copy bacterial marker protein sequences, to construct their multiple alignments and to get the taxonomic assignment using the GTDB r86 database<sup>76</sup>. Amino acid sequence sets of the MamA, -B, -M, -K, -P, and -Q proteins were independently aligned using MAFFT<sup>77</sup>, curated with Gblocks v. 0.91b<sup>78</sup> with an option that allows gap positions within the final blocks, and then concatenated. These Mam protein sequences were also used to build trees with their homologs. Maximum-likelihood trees were inferred with IQ-TREE<sup>79</sup> using evolutionary models selected by ModelFinder<sup>80</sup>. Branch supports were obtained with 1000 ultrafast bootstraps<sup>81</sup>. Trees were visualized with iTOL v4<sup>82</sup>. The genomes of *Ca*. Omnitrophus magneticus SKK-01, *Ca*. Magnetoglobus multicellularis str. Araruama, *Ca*. Magnetobacterium bavaricum TM-1, and *Ca*. Magnetoovum chiemensis CS-04 were not subjected to phylogenetic analyses because they had failed the quality check (Supplementary Table S1). Taxonomic classification of the obtained genomes on phylum rank was performed using NCBI taxonomy; other ranks were named using GTDB.

#### **Data availability**

The genomes and metagenomes used during the current study are publicly available in NCBI (https://www.ncbi.nlm.nih.gov/)<sup>44,48,83-113</sup> and IMG (https://img.jgi.doe.gov/cgi-bin/m/main.cgi)<sup>114-117</sup> databases. Scaffolds of obtained MAGs could be found in Supplementary Table S6, hosted at figshare<sup>118</sup>. All data generated and analyzed in this study are also available in figshare<sup>118</sup> and in supplementary information accompany this paper. Assembly of Rhodospirillaceae bacterium MAG\_01419\_mvb\_30 could be found in RAST (https://rast.nmpdr.org/) using 'guest' as login and as password.

#### Code availability

The following tools were used for the presented analysis and described in the main text:

Busybee web, Maxbin2, MyCC, and DAS Tool with standard parameters were used for the reconstruction of metagenome-assembled genomes (MAGs).

- 1. Busybee web https://ccb-microbe.cs.uni-saarland.de/busybee
- 2. Maxbin2 https://sourceforge.net/projects/maxbin2/
- 3. MyCC https://sourceforge.net/projects/sb2nhri/files/MyCC/
- 4. DAS Tool https://github.com/cmks/DAS\_Tool
- CheckM was used to estimate obtained genomes completeness and contamination <a href="https://github.com/Ecogenomics/CheckM">https://github.com/Ecogenomics/CheckM</a>
- 6. RefineM was used to remove contamination https://github.com/dparks1134/RefineM
- 7. QUAST helped to access quality metrics http://cab.spbu.ru/software/quast/
- 8. fastANI was used to calculate ANI https://github.com/ParBLiSS/FastANI
- 9. IDTAXA helped to obtain taxonomic assignments for the studied genomes 16S rRNA genes http://www2.decipher.codes/Classification.html
- 10. GTDB-Tk was used to find 120 single-copy bacterial marker protein sequences, to construct their multiple alignments and to get the taxonomic assignment using the GTDB r86 database https://github.com/Ecogenomics/GTDBTk
- 11. MAFFT was used for aligning amino acid sequence sets of the MamA, -B, -M, -K, -P, and -Q proteins https://mafft.cbrc.jp/alignment/server/
- 12. Gblocks helped to curate sequences aligned in MAFFT http://molevol.cmima.csic.es/castresana/Gblocks\_server.html
- 13. Phylogenetic trees were inferred with IQ-TREE http://www.iqtree.org/
- 14. Obtained trees were visualized with iTOL https://itol.embl.de/

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### **Author contributions**

M.U. and L.A. created MGC protein sequences database. M.U. conducted MGCs search, analyzed obtained data and wrote the manuscript. L.A. reconstructed MGCs of obtained genomes. M.K. conducted metagenomes binning, D.G. had the initial idea for the analysis, V.K. and D.G. discussed and interpreted the results and revised the manuscript. All authors read and approved the final manuscript.

### **Competing interests**

The authors declare no competing interests.

#### **Additional information**

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