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Authors	Lugli, Gabriele A.;Milani, Christian;Duranti, Sabrina;Mancabelli, Leonardo;Mangifesta, Marta;Turroni, Francesca;Viappiani, Alice;van Sinderen, Douwe;Ventura, Marco
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University College Cork, Ireland Coláiste na hOllscoile Corcaigh





Tracking the Taxonomy of the Genus *Bifidobacterium* Based on a Phylogenomic Approach

Gabriele Andrea Lugli,^a Christian Milani,^a Sabrina Duranti,^a Leonardo Mancabelli,^a Marta Mangifesta,^b Francesca Turroni,^{a,c} Alice Viappiani,^b Douwe van Sinderen,^{d,e} Marco Ventura^{a,c}

aLaboratory of Probiogenomics, Department of Chemistry, Life Sciences and Environmental Sustainability,

University of Parma, Parma, Italy

^bGenProbio srl, Parma, Italy

^cMicrobiome Research Hub, University of Parma, Parma, Italy

dAPC Microbiome Institute, National University of Ireland, Cork, Ireland

eSchool of Microbiology, National University of Ireland, Cork, Ireland

ABSTRACT For decades, bacterial taxonomy has been based on *in vitro* molecular biology techniques and comparison of molecular marker sequences to measure the degree of genetic similarity and deduce phylogenetic relatedness of novel bacterial species to reference microbial taxa. Due to the advent of the genomic era, access to complete bacterial genome contents has become easier, thereby presenting the opportunity to precisely investigate the overall genetic diversity of microorganisms. Here, we describe a high-accuracy phylogenomic approach to assess the taxonomy of members of the genus *Bifidobacterium* and identify apparent misclassifications in current bifidobacterial taxa belonging to the genus *Bifidobacterium* by employing their overall genetic content. The results of this study demonstrate the potential of this whole-genome approach to become the gold standard for phylogenomics-based taxonomic classification of bacteria.

IMPORTANCE Nowadays, next-generation sequencing has given access to genome sequences of the currently known bacterial taxa. The public databases constructed by means of these new technologies allowed comparison of genome sequences between microorganisms, providing information to perform genomic, phylogenomic, and evolutionary analyses. In order to avoid misclassifications in the taxonomy of novel bacterial isolates, new (bifido)bacterial taxons should be validated with a phylogenomic assessment like the approach presented here.

KEYWORDS genomics, phylogenomics, ITS, next generation sequencing, bifidobacteria, *Bifidobacterium*

S ince the 1960s, bacterial taxonomy has been determined by using the DNA-DNA hybridization approach in order to measure the degree of genetic similarity between two microbial genomes (1). Another accepted method that was and is still widely used in bacterial taxonomy is comparative analysis of 16S rRNA gene-based sequences (2). Unfortunately, the DNA-DNA hybridization method suffers from reproducibility issues and does not provide an accurate measure of actual sequence identity between genomes (3). Similar limitations affect the 16S rRNA gene approach; for example, very recently diverged species that have undergone intense evolutionary pressures may possess highly similar 16S rRNA gene sequences that may nonetheless ignore a wide phylogenetic gap between such taxa (4, 5). To overcome the limitations of these techniques, a multigenic approach that relies on multiple conserved molecular markers, such as the *clpC*, *dnaB*, *dnaG*, *dnaJ1*, *purF*, and *rpoC* genes, was shown to be more

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Address correspondence to Marco Ventura, marco.ventura@unipr.it.

G.A.L. and C.M. contributed equally to this article.

reliable for species discrimination compared to single-gene phylogeny (5, 6). Having easy access to next-generation sequencers (NGS) has in recent times allowed the development of a new bioinformatics approach for phylogeny that is based on wholegenome sequencing followed by comparative genomics. Comparative genomics has proven to be accurate in strain discrimination and has been applied extensively for phylogenetic characterization of novel bacterial species, in particular those residing in complex communities, e.g., gut microbiota members (7–9).

Members of the genus *Bifidobacterium* are among the main representatives of the mammalian gut microbiota, particularly during the first six to 12 months following birth (10, 11). This group of microorganisms is known for a claimed ability to confer a range of health benefits to the host (12, 13, 61), although the associated genetic attributes for such beneficial or probiotic activities remain largely obscure. Bifidobacteria are wide-spread inhabitants of the mammalian, avian, and insect intestinal tracts (11, 13, 14, 62), yet a large part of the currently recognized bifidobacterial taxa, i.e., 49 species and 9 subspecies, has been isolated from the human gut (15, 16). Nonetheless, various ecological attempts have recently been made to survey bifidobacterial populations in other mammals (10, 17–21), which has resulted in the identification of several scientifically accepted and putative novel bifidobacterial species.

The majority of the currently proposed novel (bifido)bacterial taxa are identified through partial sequencing of several molecular marker genes and comparison to the currently recognized type strains (19–21). In the current study, we describe a methodology that is based on whole-genome comparisons and is aimed at unambiguously redefining the taxonomy of members of the genus *Bifidobacterium*. Notably, similar approaches based on whole-genome comparisons have been employed in the reclassification of members of the genus *Bacillus* (22, 23). Applying this approach, we were also able to phylogenetically characterize and position seven novel bifidobacterial strains which were isolated from animal feces, i.e., those of goose, hamster, rabbit, and monkey, and which are related to *Bifidobacterium choerinum*, *Bifidobacterium hapali*, *Bifidobacterium saguini*, *Bifidobacterium stellenboschense*, and *Bifidobacterium tissieri*.

RESULTS AND DISCUSSION

Pangenome reconstruction among members of the genus Bifidobacterium. Recently, species- and genus-level comparative genomic analyses based on pangenome reconstruction have been shown to be crucial in providing information regarding the overall gene content, while also generating information on the resistome, metabolic capabilities, and mobilome of such taxonomic ranks (9, 24-28). In recent years, the number of sequenced bifidobacterial strains has increased from a few dozen to several hundred, and thus we felt it was opportune to explore the genomic biodiversity within different species of the genus Bifidobacterium (12, 15, 16, 29). Currently, the number of sequenced bifidobacterial strains for each species ranges from just one to 83 (see Table S1 in the supplemental material). In order to increase the amount of genetic data available for those bifidobacterial species for which only one or a few strains have been sequenced, we decided to decode the genomes of 13 additional bifidobacterial strains belonging to the species Bifidobacterium asteroides, Bifidobacterium pseudolongum, and Bifidobacterium thermophilum (Table 1). Furthermore, the public NCBI genomic database contains 55 type strain genomes corresponding to each known bifidobacterial (sub)species, as well as complete or draft genome sequences of 233 additional strains belonging to the species Bifidobacterium adolescentis, Bifidobacterium animalis, B. asteroides, Bifidobacterium bifidum, Bifidobacterium breve, Bifidobacterium dentium, Bifidobacterium longum, Bifidobacterium pseudocatenulatum, B. pseudolongum, and B. thermophilum (see Table S1).

These gathered data sets were then employed for the genomic approach to study of the phylogeny of the genus *Bifidobacterium*. Thus, a pangenome analysis of available type strains was undertaken to determine putative orthologous genes between the 55 (sub)species of the genus *Bifidobacterium* sequenced to date. The analysis resulted in the identification of 26,201 clusters of orthologous genes (COGs), representing the

TABLE 1 General features of sequenced bifidobacterial strai
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		Average		Genome		No. of				
Cuestes	Chunin	coverage	No. of	length	Average GC	predicted	No. of	No of	Dislagical aginin	GenBank
species	Strain	(1010)	contigs	(qa)	content (%)	ORFS	trinas	rkinas"	Biological origin	accession no.
B. asteroides	1460B	104.83	38	2,121,817	60.46	1,653	45	2	Honeybee hindgut	PCHJ0000000
B. pseudolongum	1370B	89.46	17	1,902,036	62.97	1,571	52	4	Feces of pig	PCHI0000000
B. pseudolongum	1520B	115.03	17	2,008,481	63.12	1,632	52	4	Feces of hamster	PCHH0000000
B. pseudolongum	1549B	118.25	48	1,990,203	63.19	1,686	55	4	Feces of Brahma chicken	PCHG0000000
B. pseudolongum	1595B	147.96	16	1,936,418	63.02	1,593	53	4	Feces of pig	PCHF00000000
B. pseudolongum	1691B	65.49	59	2,148,724	63.06	1,810	52	4	Feces of hippopotamus	PCHE00000000
B. pseudolongum	1734B	84.15	25	2,111,856	63.27	1,778	62	4	Feces of wallaby	PCHD0000000
B. pseudolongum	1619B	235.22	31	2,050,408	63.27	1,722	53	3	Feces of llama	PCHC00000000
B. pseudolongum	1744B	101.65	51	2,143,581	63.05	1,801	55	4	Feces of bear	PCHB00000000
B. pseudolongum	1747B	56.17	52	2,143,079	63.18	1,830	60	4	Feces of giraffe	PCHA0000000
B. pseudolongum	1524B	99.95	23	2,062,414	63.15	1,700	52	4	Feces of hamster	PCGZ0000000
B. thermophilum	1542B	89.45	36	2,359,132	60.28	1,820	47	3	Feces of pig	PCGY0000000
B. thermophilum	1543B	115.32	16	2,316,133	60.41	1,751	50	3	Feces of pig	PCGX0000000

^aPredicted number of rRNA loci.

pangenome of the *Bifidobacterium* genus. The collected COGs allowed the identification of genes that are shared between the genomes of the 55 bifidobacterial type strains, i.e., the core genome of the *Bifidobacterium* genus (Fig. 1). Furthermore, dispensable genes present in two or more strains and unique genes retrieved in a single type strain were unveiled (Fig. 1). The pangenome size, when plotted versus the number of included bifidobacterial genomes, shows that the power trendline has yet



FIG 1 Pangenome of the genus *Bifidobacterium*. (a) Number of core genes (gray), unique genes (blue), and dispensable genes (orange) identified in the pangenome analysis (internal pie chart). The number of unique genes of each type strain of the genus sequenced to date is highlighted in different colors (external pie chart). (b) Representation of the pangenome size based on sequential addition of the 55 type strains genomes. (c) Average of new genes upon sequential addition of the bifidobacterial type strain genomes.

to reach a plateau and data sets from the last strains added to the analysis still substantially expand the total gene pool by circa 330 genes per added genome (Fig. 1). Therefore, according to these data, the pangenome of the genus *Bifidobacterium* can be considered open (25) (Fig. 1). This open pangenome profile is typically associated with genera in which the constituent species occupy multiple environments with mixed microbial communities and have extended their total set of genes through horizontal gene transfer events, e.g., *Escherichia* (9, 30).

In order to assess intraspecies variability, we selected bifidobacterial species for which at least five sequenced genomes are available in the NCBI databases, or which were decoded as part of the current study. Specifically, a total of 256 different strains were employed for the reconstruction of 10 species-specific bifidobacterial pangenomes (see Fig. S1 and S2 in the supplemental material). Notably, and similar to those of the pangenome of the overall Bifidobacterium genus (Fig. 1), the power trendlines do not reach a plateau when species-specific pangenome sizes are plotted versus the number of included genomes (see Fig. S1 and S2). Furthermore, the species-specific pangenome analyses also revealed that the average number of new genes added by inclusion of additional genomes tend to decrease for 7 out of 10 analyzed species. In contrast, the comparative genomics analysis of B. animalis, B. breve, and B. longum revealed that the pangenome expansion seems to remain stable at a minimum of circa 43, 75, and 101 new genes added for each further iteration, respectively (see Fig. S1), suggesting that the biodiversity of these species has been extensively explored. In this regard, it is worth mentioning that a portion of these "new" genes corresponds to truncated genes predicted at the edges of contigs within partially sequenced genomes. In this context, the pangenome's expansion trend in B. breve and B. longum reflects the high number of strains sequenced for both these two species (12, 16), whereas the limited pangenome expansion of *B. animalis* is probably due to the monophyletic origin of the subspecies B. animalis subsp. lactis, as previously suggested (29).

Overall, these data indicate that availability of a very large number, perhaps as many as thousands, of strains representing populations residing in a wide range of different hosts will be pivotal to obtain a comprehensive overview of the biodiversity that characterizes these common gut commensals.

Phylogenetic and phylogenomic analyses of bifidobacterial type strains. While 16S rRNA gene-based comparative analysis has for many years been considered to represent the gold standard for phylogenetic investigations, the recent advent of NGS and associated bioinformatic tools has led to the introduction of novel genome-wide approaches, such as the core-genome-based supertree (31, 32). We therefore decided to compare the phylogeny of the type strains of the genus Bifidobacterium reconstructed through alignment of the 16S rRNA gene and concatenated alignment of the bifidobacterial core genome. The 16S rRNA genes were retrieved from the 55 type strains of each taxon belonging to the genus Bifidobacterium and were used to construct a phylogenetic tree (Fig. 2). Moreover, the predicted 26,201 clusters of orthologous genes identified by comparative genomics analysis of the bifidobacterial type strains allowed the identification of 262 shared COGs, representing the core bifidobacterial genome coding sequences (Fig. 2). After exclusion of paralogs, concatenation of the remaining 236 core protein sequences was used to build a Bifidobacterium phylogenomic tree (Fig. 2). These analyses showed that 34 and 52 nodes were supported by bootstrap values greater than 50 for the 16S rRNA gene- and core gene-based trees, respectively (Fig. 2). These data clearly support the notion that increasing sequence lengths used in the phylogenomic tree leads to improved robustness of the results. Furthermore, the phylogenomic analyses are based on a core genome, which is represented by amino acid sequences. Remarkably, these protein sequences involve a variation of 20 amino acids, while the sequence of the 16S rRNA gene is based on only four bases. The use of amino acid sequences clearly enhances the robustness of the resulting phylogenomic tree.



FIG 2 Phylogenetic and phylogenomic trees of the genus *Bifidobacterium*. (a) Proteomic tree based on the concatenation of 262 bifidobacterial core genes identified in the pangenome analysis of the 55 type strains. Phylogenetic groups are highlighted in different colors. (b) 16S rRNA gene-based tree of the 55 type strains of the genus. Bifidobacterial species are colored according to their phylogenetic membership group. Each tree was constructed by the neighbor-joining method, and the genome sequence of *Scardovia inopinata* JCM 12537 was used as outgroup. Bootstrap percentages above 50 are shown at node points, based on 1,000 replicates of each phylogenetic tree.

The bifidobacterial phylogenomic tree clearly delineates the presence of seven previously described bifidobacterial phylogenetic groups, i.e., the B. adolescentis, B. asteroides, B. bifidum, Bifidobacterium boum, B. longum, B. pseudolongum, and Bifidobacterium pullorum groups (33, 34). Notably, the number of bifidobacterial taxa included does not affect the consistency of the core gene-based tree (Fig. 2) (9, 33). Comparison of the phylogenomic and 16S rRNA gene-based trees revealed discrepancies for the B. bifidum and B. longum groups (Fig. 2), while the phylogeny of the B. adolescentis, B. asteroides, B. boum, B. pseudolongum, and B. pullorum groups is conserved in both trees. In detail, the four members of the B. bifidum group cluster together only in the core gene-based tree, while they are scattered across the 16S rRNA gene-based tree. In contrast, the core of the *B. longum* group is consistent in both trees, represented by B. longum subsp. suis LMG 21814, B. longum subsp. longum LMG 13197, B. longum subsp. infantis ATCC 15697, Bifidobacterium saguini DSM 23967, B. breve LMG 13208, Bifidobacterium eulemuris DSM 100216, and Bifidobacterium lemurum DSM 28807 (Fig. 2). Nevertheless, the B. longum group identified in the core gene-based tree includes six additional taxa compared to that in the 16S rRNA gene-based tree.

The presented comparison between the phylogenetic reconstruction based on alignment of the 16S rRNA gene and the concatenated alignment of the bifidobacterial core genome suggests that the latter permits the reconstruction of a more robust and consistent overview of bifidobacterial evolution and may for this reason be considered the preferential approach for genus- and species-wide phylogenetic investigations.

Evaluation of intraspecies variability between bifidobacterial genomes. The 256 bifidobacterial genomes collected above were also used to perform whole-genome comparisons, focusing specifically on the species B. adolescentis, B. animalis, B. asteroides, B. bifidum, B. breve, B. dentium, B. longum, B. pseudocatenulatum, B. pseudolongum, and B. thermophilum. The genomes of bifidobacterial taxa belonging to the same species were subjected to average nucleotide identity (ANI) evaluation. These analyses revealed consistent species classification in seven out of 10 phylogenetic groups, highlighting ANI values ranging from 95.31% to 99.98% (see Table S2 and Fig. S3–S6 in the supplemental material). In this regard, it should be noted that two strains displaying an ANI value of <95% are considered to belong to two distinct species (35, 36). Thus, all the strain pairs that belong to the species B. adolescentis, B. animalis, B. bifidum, B. breve, B. dentium, B. longum, and B. pseudocatenulatum possess ANI values higher than 95%, confirming taxonomic assignment to the same species (35, 36). Conversely, the genomes belonging to the species B. asteroides, B. pseudolongum, and B. thermophilum exhibit cases of ANI values lower than 93.3%, which would cast doubt on the correct taxonomic classification of several strains previously assigned to these species (see Table S2).

More specifically, within the species *B. thermophilum*, the chromosomal sequence of strain JCM 1207 generates ANI values of 90.16% and 90.28% compared to the genome sequences of *B. thermophilum* DSM 20210 and *B. thermophilum* DSM 20212, respectively (see Table S2). These values are the lowest retrieved between strains of the same species in this intraspecies analysis, highlighting that the JCM 1207 strain should not be classified as a *B. thermophilum* species (see Fig. S6 in the supplemental material). Further ANI analyses encompassing all bifidobacterial type strains showed that *B. thermophilum* JCM 1207 possesses a higher genome sequence identity with *B. boum* LMG 10736, sharing an ANI value of 94.9% (33), which is very close to the threshold for species discrimination (35, 36).

Besides, different clusters were detected within the species *B. pseudolongum* (see Fig. S6). In particular, the genome sequence of *B. pseudolongum* subsp. *pseudolongum* LMG 11571 displays high ANI values compared to strains 1370B (99.21%) and 1595B (99.21%), both sequenced in the current study. Nonetheless, the chromosomal sequences of these three strains generate corresponding ANI values lower than 93.99% compared to 11 other *B. pseudolongum* strains, which among each other display ANI values above 95.81% (see Table S2). These data suggest that *B. pseudolongum* subsp. *pseudolongum* LMG 11571, along with strains 1370B and 1595B, may represent a distinct bifidobacterial species.

The most intriguing data, due to the high genomic diversity observed for the six analyzed strains, were retrieved for members of the species *B. asteroides*. In fact, the genomes belonging to 1460B, Bin2, Hma3, and Bin7 strains produced ANI values that are below the species threshold level compared to the genome of the type strain *B. asteroides* DSM 20089 (see Table S2) and resulted in four clusters, i.e., four putative distinct bifidobacterial species (as reported in Fig. S3).

Further inspection of genomic sequence identity at subspecies level revealed that the chromosomes of the 30 *B. animalis* strains analyzed constitute two clear clusters, encompassing seven *B. animalis* subsp. *animalis* and 23 *B. animalis* subsp. *lactis*, with strains A6, 08, 11, and RH without a (stated) subspecies classification falling in the latter group (see Fig. S3). Moreover, the low genome variability observed in the *B. animalis* subsp. *lactis* strains confirms the previously reported observation that this subspecies is a strict monophyletic taxon (29, 37, 38).

Interestingly, the intraspecies analysis performed on *B. longum* highlights several inconsistencies in the subspecies classifications (see Fig. S5). In this context, a clear cluster composed of 19 *B. longum* subsp. *infantis* strains was identified, as well as a smaller cluster representative of *B. longum* subsp. *suis*. Notably, the latter cluster includes the genomes of AGR2137, Su859, CMCC P0001, BXY01, and JDM301 strains previously classified as *B. longum* subsp. *longum*. Remarkably, strains named CMCC

P0001, BXY01, and JDM301 displayed ANI values above 99.89% compared to each other, confirming their previously described association (12). Nonetheless, genome comparisons between the seven identified *B. longum* subsp. *suis* generated ANI values above 98%, and for this reason no distinct phylogenetic groups could be identified based on genomic identity approaches (see Table S2). The largest *B. longum* cluster contains 57 *B. longum* subsp. *longum* strains, of which three strains, named 157F, CCUG 52486, and CECT 7210, had erroneously been classified as *B. longum* subsp. *infantis* strains.

Overall, ANI evaluation of all 258 available bifidobacterial genomes indicated that 16 strains should be taxonomically reclassified, while it also revealed the putative existence of six novel bifidobacterial species.

Genome sequencing as the most current standard for taxonomic classification of bifidobacteria. The pipeline for accurate bifidobacterial taxonomic classification described here was also exploited to precisely taxonomically classify putative novel bifidobacterial taxa isolated as part of a recently published study aimed at the exploration of the biodiversity of bifidobacterial communities of 291 animals, including goose, hamster, rabbit, and monkeys (10) (Table 2). Subsequent selection on mupirocin medium (39) allowed the isolation of strains belonging to members of the genus Bifidobacterium, which were further characterized by amplification and sequencing of the 16S rRNA gene and internal transcribed spacer (ITS). While 16S rRNA gene sequence comparisons have been used in (bifido)bacterial taxonomy for decades, the ability to distinguish closely related bifidobacterial taxa using ITS sequences has recently been described (40). 16S rRNA gene sequence analysis revealed that seven isolated strains, named Rab10A, Ham19E, Goo31D, Tam1G, Tam10B, Uis4E, and Uis1B, showed 16S rRNA sequence-based identity values that ranged from 96.2% to 98.6% with respect to known bifidobacterial type strains listed in Table 2, while the hypervariable ITS seguence displayed values ranging from 65.7% to 89% (Table 2). Notably, these data highlight a high degree of sequence diversity between these seven isolates and the known bifidobacterial species, especially for the hypervariable ITS sequence, thus suggesting that they present novel bifidobacterial species.

In order to get insights into genome-wide genetic relatedness between the putative novel species and currently known taxa of the genus Bifidobacterium, the genomes of these seven isolates were sequenced. The reconstructed genome length ranged from 2,155,882 to 3,111,005 bp, with an average fold coverage ranging from 55.52 to 139.27 (Table 2). Using the ANI system based on whole genome sequence comparisons, the seven sequenced strains were compared with the currently recognized bifidobacterial type strains (9, 41). Interestingly, the seven isolates exhibited ANI values below the threshold for species recognition compared to all 55 available type strains, with the highest ANI value obtained against B. saquini DSM 23967 (94.55%) (Fig. 3) (36, 42). Furthermore, an ANI analysis involving only these seven strains revealed that the highest ANI value (91.76%) was obtained between the genomes of Rab10A and Ham19E (Fig. 3). Genome-to-Genome Distance Calculator (GGDC) analysis, which is based on in silico DNA-DNA hybridization (DDH) of genome-to-genome comparison (43), was employed to validate the ANI results based on the genomic relatedness between bifidobacterial taxa. The seven sequenced strains' genomes compared with the closest related type strains' genomes exhibited estimated DDH values below 70%, ranging from 28.5% to 55.2% between pairs Uis1B/B. hapali DSM 100202 and Tam1G/B. saquini DSM 23967, respectively (Table 2).

Taken together, ANI and GGDC analyses clearly indicate that Rab10A, Ham19E, Goo31D, Tam1G, Tam10B, Uis4E, and Uis1B belong to novel bifidobacterial species (Fig. 3). Recently, based on this genomic approach, Tam10B was formally accepted by the International Committee on Systematic Bacteriology (ICSB) as a novel species of the genus *Bifidobacterium*, and was accordingly named *Bifidobacterium vansinderenii* (44).

Phylogenomic approach for the evaluation of novel bifidobacterial taxa. The availability of genome sequences of the seven putative novel bifidobacterial species

strains	
bifidobacterial	
of novel	
features	
2 General	
TABLE 2	

	Average coverage	No. of	Genome length	Average GC	No. of predicted							
Strain	(fold)	contigs	(dq)	content (%)	ORFs	tRNA	rRNAa	16S identity (%, species)	ITS identity (%, species)	ANI value (%, species)	GGDC ^b value (%, species)	Biological origin
Rab10A	104.3	72	2,276,351	65.45	1,825	52	4	98.36, B. choerinum	87.2, B. choerinum	91.8, B. choerinum	41.30, B. choerinum	European rabbit
Ham19E	108.6	44	2,155,882	62.53	1,733	53	4	97.32, B. choerinum	68.6, B. animalis	87.81, B. choerinum	28.50, B. choerinum	European hamster
G0031D	115.4	6	2,166,761	64.3	1,681	52	4	97.65, B. choerinum	84.7, B. choerinum	91.47, B. choerinum	39.30, B. choerinum	Domestic goose
Tam 1G	55.52	62	2,639,899	56.13	2,215	59	4	99, B. saguini	89, B. saguini	94.55, B. saguini	55.20, B. saguini	Emperor tamarin
Tam 10B	138	68	3,111,005	62.46	2,522	60	4	96.4, B. tissieri	72, B. hapali	89, B. tissieri	31.60, B. tissieri	Emperor tamarin
Uis1B	66.93	80	2,789,387	61.91	2,281	60	č	96.2, B. tissieri	65.7, B. biavatii	88.04, <i>B. hapali</i>	28.50, B. hapali	Pygmy marmoset
Uis4E	139.27	44	2,820,211	65.81	2,247	62	9	96.8, B. stellenboschense	81.3, B. stellenboschense	93.45, B. stellenboschense	50.20, B. stellenboschense	Pygmy marmoset
^a Predicte	d number of	f rRNA loci.										

^bGGDC, Genome-to-Genome Distance Calculator.



FIG 3 Graphical representation in three-dimensional columns of the average nucleotide identity (ANI) between novel bifidobacterial taxa. The *x* axis displays the 62 bifidobacterial taxa used in the comparison (55 type strains and seven novel taxa), the *y* axis reflects the ANI percentages between strains, and the *z* axis displays the seven novel taxa. Thus, each column represents the ANI value observed between a given bifidobacterial species pair. The higher ANI values, involving Rab10A, Ham19E, Goo31D, Tam10B, Uis4E, and Uis1B, are highlighted with different colors, showing ANI values below 95%.

also allowed updating of the phylogeny of the genus *Bifidobacterium*. A comparative genomics analysis was undertaken to determine putative orthologous genes between the 55 sequenced type strains of the genus *Bifidobacterium* and the seven putative novel taxa, resulting in the identification of 27,868 BifCOGs (*Bifidobacterium*-specific clusters of orthologous genes). Analysis of the predicted BifCOGs identified 259 COGs that were shared among all these genomes, representing the core bifidobacterial genome coding sequences (core BifCOGs). This core BifCOG collection represents an updated version of a previously published core genome of the genus *Bifidobacterium* (9, 33). The concatenation of 232 core BifCOG protein sequences (note that 27 core BifCOGs were excluded, as they constitute paralogs within the bifidobacterial pangenome) was used to build a *Bifidobacterium* phylogenomic tree (Fig. 4).

The updated bifidobacterial phylogenomic tree confirmed the seven bifidobacterial phylogenetic groups previously described (33, 34). Moreover, the seven putative novel species appear to be distributed across the whole tree. As expected through the ANI analysis, Rab10A, Ham19E, and Goo31D cluster in the proximity of *B. choerinum* LMG 10510 and fall within the *B. pseudolongum* group. Interestingly, Rab10A, Ham19E, and Goo31D were isolated from three different animals, i.e., rabbit, hamster, and goose, respectively, while *B. choerinum* LMG 10510 is an isolate from piglet feces. Thus, these data indicate that the *B. pseudolongum* group is currently the most variable phylogenetic bifidobacterial group in terms of ecological niches represented by animal species, in which group strains were isolated from chickens, geese, hamsters, oxen, pigs, rabbits and rats. Nonetheless, one has to keep in mind that related bifidobacterial species



FIG 4 Phylogenomic tree of the genus *Bifidobacterium* based on the concatenation of the deduced amino acid sequences of 232 core genes of Goo31D, Ham19E, Rab10A, Tam1G, Uis4E, and Uis1B and members of the genus *Bifidobacterium*. The amino-acid-deduced core gene-based tree shows the division intoseven phylogenetic groups of the *Bifidobacterium* genus as represented by different colors. The phylogenetic tree was constructed by the neighbor-joining method, with genome sequence of *Scardovia inopinata* JCM 12537 as outgroup. Bootstrap percentages above 50 are shown at node points, based on 1,000 replicates of the phylogenetic tree.

appear to be widespread among mammals (10). Moreover, strains Tam1G and Uis4E are included in the *B. longum* group and show high phylogenetic relatedness with *B. saguini* DSM 23967 and *B. stellenboschense* DSM 23968, respectively (Fig. 4). Furthermore, Tam10B and Uis1B cluster together with *B. tissieri* DSM 100201.

Phylogenomic analysis indicates that 13 bifidobacterial taxa do not belong to any previously described phylogenetic group (Fig. 4). Nevertheless, based on the relatedness among these bifidobacterial type strains, two new phylogenetic clusters are proposed, namely, the *Bifidobacterium psychraerophilum* group and the *Bifidobacterium bombi* group (Fig. 4). The results presented highlight once again the potential of the phylogenomic approach to establish detailed phylogenetic reconstruction of the entire *Bifidobacterium* genus and detailed taxonomic characterization of novel bacterial species.

Conclusions. Next-generation sequencing has significantly influenced microbial taxonomy by giving access to genome sequences of essentially all known bacterial taxa. In fact, the deciphered genome sequences of bifidobacterial type strains now provide genetic information for genomic, phylogenomic, and evolutionary analyses, and facilitate the determination of the gene contribution of each isolated microorganism. Here,

301 sequenced strains belonging to the genus Bifidobacterium were compared to perform inter- and intraspecies analyses aimed at redefining bifidobacterial taxonomy and unveiling discrepancies in species assignment. Overall, ANI analyses identified inconsistencies in classification of a total of 16 strains within the B. asteroides, B. pseudolongum, and B. thermophilum species, unveiling the existence of six additional clusters of strains that may represent novel putative bifidobacterial species. Furthermore, we validated the potential of the phylogenomics approach in the identification of novel species through the sequencing of new bifidobacterial isolates. Seven of these strains may be characterized as novel bifidobacterial species, showing genomic compositions related to those of B. choerinum LMG 10510, B. hapali DSM 100202, B. saguini DSM 23967, B. stellenboschense DSM 23968, and B. tissieri DSM 100201, while maintaining ANI values below the threshold for species assignment. The phylogenomic analysis of these seven putative novel species also revealed their localization into the updated phylogeny of the genus Bifidobacterium, with three strains belonging to the B. pseudolongum group and two novel taxa falling into the B. longum group. In light of these results, we propose to implement the current taxonomic scheme for the classification of novel bifidobacterial taxa through a (phylo)genomic assessment of a proposed new taxon.

MATERIALS AND METHODS

Bifdobacterial genome sequences. We retrieved complete and partial genome sequences of 55 *Bifdobacterium* type strains from the National Center for Biotechnology Information (NCBI) public database, in additional to genomes of a further 233 taxa that belong to this genus (see Table S1). Furthermore, 13 bifdobacterial genomes were sequenced to perform intraspecies analyses, creating a pool of 301 bifdobacterial genomes. We also analyzed the genome sequences of seven novel bifdobacterial taxa deposited at DDBJ/ENA/GenBank, available under accession numbers MVOG0000000, MVOH00000000, NEWD0000000, NMWU00000000, NMWV00000000, and NMYC00000000 (Table 2).

Isolation of bifidobacterial species. Fecal samples were collected from several zoological parks as described previously (10). In order to acquire the fecal material as fresh as possible and to be sure of its origin, it was collected immediately following defecation. Fecal samples consisted of 6 to 10 g of fresh material, which was cooled to 4°C immediately after collection and transferred to the Laboratory of Probiogenomics of Parma University (Parma, Italy). Bifidobacterial isolation from stool samples was performed starting from 1 g of fecal sample mixed with 9 ml of phosphate-buffered saline (PBS) solution. Serial dilutions and subsequent plating were performed using the de Man-Rogosa-Sharpe (MRS) agar, supplemented with 50 μ g/ml mupirocin (Delchimica, Italy) and 0.05% (wt/vol) L-cysteine hydrochloride. Bifidobacterial cultures were incubated for 48 h at 37°C in a chamber (Concept 400; Ruskin) with anaerobic atmosphere (composed of 2.99% H₂, 17.01% CO₂, and 80% N₂). Colonies were randomly picked and restreaked to isolate purified bacterial strains. All colonies were subjected to DNA isolation and characterized as previously described by Turroni et al. (45) and Ventura et al. (46).

DNA extraction and amplification of 16S rRNA and ITS sequences. Fecal samples maintained at -80° C were subjected to DNA extraction using the QIAamp DNA Stool minikit following the manufacturer's instructions (Qiagen). Partial 16S rRNA gene sequences were amplified from extracted DNA using the primer pair Probio_Uni/Probio_Rev, which targets the V3 region of the 16S rRNA gene sequence (47), while partial ITS sequences were amplified using the Probio-bif_Uni/Probiobif_Rev primer pair, which targets the hypervariable region of the bifidobacterial ITS sequences (40). Results were then subjected to a BLAST search against the GenBank database.

Genome sequencing and assemblies. DNA extracted from the bifidobacterial isolates was subjected to whole-genome sequencing using a MiSeq system (Illumina, UK) at GenProbio srl (Parma, Italy) following the supplier's protocol (Illumina, UK). Fastq files of the paired-end reads obtained from targeted genome sequencing of the isolated strains were used as input for the genome assemblies through the MEGAnnotator pipeline (48). The MIRA program (version 4.0.2) was used for *de novo* assembly of each bifidobacterial genome sequence (49).

Genome annotation. Protein-encoding open reading frames (ORFs) were predicted using Prodigal (50). tRNA genes were identified using tRNAscan-SE v1.4 (51), while rRNA genes were detected using RNAmmer v1.2 (52). Gene annotation was defined by means of RAPSearch2 (Reduced Alphabet based Protein similarity Search 2) (53) in a nonredundant protein database provided by the National Center for Biotechnology Information (NCBI) and a hidden Markov model (HMM) search (http://hmmer.org/) of the manually curated Pfam-A protein family database (54). Results were inspected by Artemis (55), which was used for genome analyses of predicted genes and for manual editing where necessary.

Pangenome and identification of shared and unique genes. Genomes of bifidobacterial type strains (see Table S1), together with bifidobacterial taxa that belong to the same species (Table 1; see also S1), as well as the seven sequenced genomes of novel strains (Table 2) were subjected to a pangenome calculation using PGAP (Pan-Genomes Analysis Pipeline) (56). In order to reduce genome content redundancy, all analyses included a single type strain of each (sub)species. The ORF content of all assessed genomes was organized into functional gene clusters using the GF (gene family) method, which

involves comparison of each protein to all other proteins using BLAST analysis, followed by clustering into protein families named as *Bifidobacterium*-specific clusters of orthologous genes (BifCOGs; cutoff E value of 1×10^{-5} and 50% identity across at least 80% of both protein sequences), using MCL (a graph-theory-based Markov cluster algorithm) (57). Pangenome profiles were built using an optimized algorithm incorporated into PGAP software, based on a presence/absence matrix that included all identified BifCOGs in the analyzed genomes. Consequently, the unique protein families for each bifidobacterial genome were classified. Protein families shared between all genomes, named core BifCOGs, were defined by selecting the families that contained at least one single protein member for each genome.

Phylogenomic comparison between strains. The concatenated core genome sequences of the genus *Bifidobacterium* were aligned using MAFFT (Multiple Alignment using Fast Fourier Transform) (58), and the corresponding phylogenomic tree was constructed using the neighbor-joining method in Clustal W version 2.1 (59). The core genome tree was built using FigTree (http://tree.bio.ed.ac.uk/software/figtree/). For each genome pair, a value for the average nucleotide identity (ANI) was calculated using the program JSpecies version 1.2.1 (35). Clusters based on ANI values between bacterial strains were constructed using Multiple Experiment Viewer (MeV) software (60). The Genome-to-Genome Distance Calculator (GGDC) version 2.1 was employed to estimate the DNA-DNA hybridization (DDH) between bifodbacterial taxa using the recommended "Formula 2" (identities/high-scoring segment pairs length) (43).

Accession number(s). The 13 bifidobacterial genome sequences have been deposited at DDBJ/ENA/ GenBank under the accession numbers reported in Table 1.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/AEM .02249-17.

SUPPLEMENTAL FILE 1, PDF file, 1.6 MB. **SUPPLEMENTAL FILE 2,** XLSX file, 0.1 MB.

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We declare that we have no competing interests.

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