

Genome Instability in *Lactobacillus rhamnosus* GG

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We describe here a comparative genome analysis of three dairy product isolates of *Lactobacillus rhamnosus* GG (LGG) and the ATCC 53103 reference strain to the published genome sequence of *L. rhamnosus* GG. The analysis showed that in two of three isolates, major DNA segments were missing from the genomic islands LGGISL1,2. The deleted DNA segments consist of 34 genes in one isolate and 84 genes in the other and are flanked by identical insertion elements. Among the missing genes are the *spaCBA* genes, which encode pilin subunits involved in adhesion to mucus and persistence of the strains in the human intestinal tract. Subsequent quantitative PCR analyses of six commercial probiotic products confirmed that two more products contain a heterogeneous population of *L. rhamnosus* GG variants, including genotypes with or without *spaC*. These results underline the relevance for quality assurance and control measures targeting genome stability in probiotic strains and justify research assessing the effect of genetic rearrangements in probiotics on the outcome of *in vitro* and *in vivo* efficacy studies.

Recently, the concept of “generic probiotics” was introduced, as a practical solution to create access to probiotics for people in the developing world (1). This concept refers to the free use of probiotic bacteria introduced under a novel brand name, after intellectual property rights have expired. In analogy to generic drugs, also patent-expired probiotics are free to be used by others, and health and safety claims from the expired patents can be linked to the generic probiotic strains, provided that the genotype of the original strain is identical to that of the generic strain. This raises questions about the extent of genome stability in probiotic bacterial strains and its impact on probiotic functionality.

Today, only a few examples exist about active components of probiotics, which could be targeted to confirm the presence of the associated functionality (2). Lebeer et al. (3) report on the classification of several genes and molecules that contribute to probiotic and health-promoting actions of *Lactobacillus*: (i) adaptation factors, including determinants of stress resistance, metabolism in the host, and adherence to the gut mucosa, and (ii) probiotic factors directly mediating health effects including antipathogenic, epithelium barrier-preserving, and immunomodulatory molecules. In another recent study, a collection of lactic acid bacteria isolated from fermented foods has been screened for more than 30 probiotic functionality related genes. The authors defined genes as “probiotic” that are involved in survival in the gastrointestinal tract (resistance to low pH and bile salt), starch metabolism, and folate and riboflavin synthesis (4).

Many of the probiotic strains marketed today originate from intestinal or plant isolates, and the shift from their complex and highly variable niche to the relatively constant and nutrient-rich industrial production environment is likely to be linked with selection for metabolic simplification (5). This simplification can coincide with gene loss that occurs at relatively high frequencies in genomes with a large number of insertion sequences (IS). These IS elements consist of transposase genes flanked by two inverted repeats, which are mobile through transposition. Accordingly, they play an important role in the occurrence of mutations, the disruption of genes, the overexpression of genes, and chromosomal rearrangements such as deletions, duplications, or inversions (6–8).

In the present study, we carried out a comparative genome analysis to investigate if, and to what extent, the genome of the probiotic *Lactobacillus rhamnosus* GG (LGG), which contains a relatively high number of 69 IS elements (9, 10), is stable in commercial products. Our findings indicate that major genetic rearrangements occurred in some dairy product isolates, which include the deletion of genes that are hypothesized to play a role in probiotic functionality.

The results of this work emphasize the need for quality control and assurance strategies targeting the presence and maintenance of genes involved in host-microbe interactions underlying probiotic functionality, in particular in instable genome regions or genomic islands with a relatively high occurrence of mobile genetic elements. In addition, these results justify further study of the effect of genetic rearrangements in probiotics on the outcome of *in vitro* and *in vivo* efficacy studies.

MATERIALS AND METHODS

Bacterial strain and cultivation conditions. The strains used in the present study were isolated from products by incubation of dilution series on Rogosa (RO) or MRS agar plates (Tritium Microbiology, Eindhoven, The Netherlands) under aerobic or anaerobic conditions at 37°C as shown in Table 1. Species identities were confirmed by sequencing of 16S-rRNA genes (Baseclear, Leiden, The Netherlands).

DNA extraction. DNA was isolated from bacterial colonies by the use of the AGOWA mag Mini DNA isolation kit (LGC Genomics, Berlin, Germany). When DNA was directly isolated from probiotic dairy products, the isolation was preceded by the addition of 45 ml of 2% sodium

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TABLE 1 Strains used in this study, their origin, and references

Description ^a	Source ^b	Species	Strain	Culture collection no.	Novel name	SRA BioSample accession no. ^c	Source or reference
Reference*†	LMG culture collection‡	<i>L. rhamnosus</i>	GG	LMG 18243, ATCC 53103, TNO 2012.097		SAMN01831559	
Product isolate 1*	Liquid dairy formula 1‡	<i>L. rhamnosus</i>	GG	LMG 25859, TNO 2012.098	<i>L. rhamnosus</i> yoba 2010	SAMN01831560	1
Product isolate 2A*	Liquid dairy formula 2A ^c	<i>L. rhamnosus</i>	GG	TNO 2010.113		SAMN01831561	
Product isolate 2B†	Liquid dairy formula 2B‡ ^c	<i>L. rhamnosus</i>	GG				
Product isolate 3*†	Powder dairy formula 3‡	<i>L. rhamnosus</i>	GG	LMG 27229, TNO 2012.094	<i>L. rhamnosus</i> yoba 2012	SAMN01831562	This study
Product sample 4†	Liquid dairy formula 4	<i>L. rhamnosus</i> , <i>S. thermophilus</i> ^d	GG				
Product sample 5†	Liquid dairy formula 5	<i>L. rhamnosus</i>	GG				
Product sample 6†	Liquid dairy formula 6	<i>L. rhamnosus</i>	GG				
Product sample 7†	Dried supplement 7	<i>L. rhamnosus</i>	GG				
Product sample 8†	Liquid dairy formula 8	<i>L. casei</i> ^d	DN-114001				

^a *, used for comparative genome sequence; †, qPCR was conducted on the entire product.

^b ‡, qPCR was conducted on product isolates.

^c Same product, but different batches; 2A was produced 1 year before 2B.

^d Negative control samples.

^e The entire study results have been deposited at the NCBI BioSample database, Sequence Read Archive, project number SRP017797.

citrate (pH 7) to a 5-ml sample, followed by 30 min of incubation at 30°C and centrifugation for 15 min at 3,000 rpm.

qPCR. Quantitative PCR (qPCR) was adapted from a protocol reported previously (11). Primer-probe combinations for qPCR were specifically designed for the *spaC* gene (LGG_00444) and a single-copy gene, LGG_00154, by the use of Primer Express Software v2.0 (Applied Biosystems, Bleiswijk, The Netherlands), as listed in Table S1 in the supplemental material. LGG_00444 encodes a pilin subunit for binding to human intestinal mucus. LGG_00154 encodes a hypothetical membrane associated protein, designated *map*, located outside LGGISL1,2 (locus 168611 to 168105). Moreover, LGG_00154 is located on an amplicon previously selected for specific identification of *L. rhamnosus* GG (12).

The TaqMan probes contained the minor groove binder (MGB) probe in combination with a nonfluorescent quencher and a reporter. The experiment was performed using the 7500 Fast Real-Time PCR System (Applied Biosystems) with the following settings: 1 step of 2 min at 50°C and 10 min at 95°C, followed by 50 cycles of 15 s at 95°C and 1 min at 60°C. The composition of the qPCR mix included 15 µl of 2× PCR Mastermix (Diagenode, Liège, Belgium), 1.3 µl of primer forward (10 pmol µl⁻¹), 1.3 µl of primer reverse (10 pmol µl⁻¹), 1.3 µl of MGB probe (5 pmol µl⁻¹), 10 µl of ultrapure (type 1) water (Millipore, Amsterdam, The Netherlands), and 1 µl of DNA sample. Dilutions of *L. rhamnosus* genomic DNA were quantified by optical density measurement at a wavelength of 260 nm (with an extinction coefficient for double-stranded DNA of 0.020 µg ml⁻¹ cm⁻¹) and used as standards for a calibration curve (10⁸ to 1 fg µl⁻¹). The threshold cycle (*C_t*) values were derived from the qPCR and set threshold values. The corresponding amount of DNA is derived from the calibration curve.

Nextera DNA library preparation and MiSeq sequencing. Nextera DNA library was prepared for analysis on an Illumina MiSeq sequencer according to the Illumina Nextera protocol. Briefly, 50 ng of genomic DNA was tagged and fragmented in the presence of transposomes with adapters, purified, and enriched by a limited-cycle PCR. Cluster generation and sequencing was performed according to the Illumina MiSeq system *Quick Reference Guide*. Each library was sequenced paired end for 150 cycles on the MiSeq.

TruSeq DNA library preparation and HiSeq next generation sequencing. For each sample, an indexed sequencing library was prepared of 1 µg of genomic DNA according to the Illumina TruSeq DNA protocol. One microliter of each library was loaded on an Agilent Technologies

2100 Bioanalyzer using a high-sensitivity DNA assay to determine the library concentration and to check for quality. The libraries were clustered on the Illumina cBOT station and sequenced paired end for 101 cycles on the HiSeq 2000 sequencer according to the Illumina cluster and sequencing protocols.

NGS primary data analysis. The image analysis and base calling was performed on the MiSeq or HiSeq2000 system using the Illumina online basecaller. The resulting data were demultiplexed using NARWHAL software (13) and aligned to the *L. rhamnosus* GG genome (accession number NC_013198.1) using a Burrows-Wheeler aligner (14). The resulting SAM files were sorted and converted to binary SAM with Samtools (15).

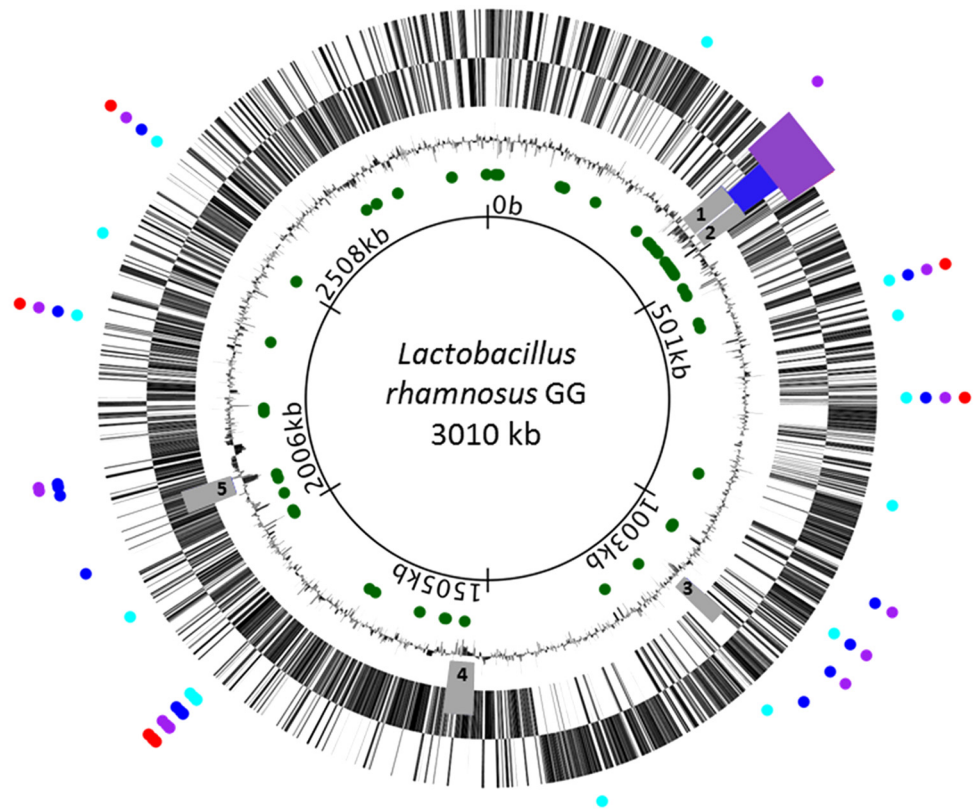
Comparative genome analysis. The genome sequence data of the three isolates and reference strain have been deposited at the NCBI BioSample database, Sequence Read Archive, project SRP017797, as indicated in Table 1. Comparative genome analysis was carried out by use of the Breseq tool, version 0.18 (16). The results of the analysis are accessible at <http://www.yoba4life.com/yoba-for-life-rd/>. The circular and linear genome maps of *L. rhamnosus* GG in Fig. 1 were created by the use of Microbial Genome Viewer, version 2.0 (17).

RESULTS

Genomic islands deleted in two propagated strains. Species identity was confirmed by sequencing of 16S rRNA genes (data not shown). The full genome sequence of three *L. rhamnosus* GG dairy product isolates and reference strain ATCC 53103, comprising approximately three million base pairs, was determined by massive parallel sequencing. Product isolate 1 was designated *L. rhamnosus* yoba 2010, and product isolate 3 was designated *L. rhamnosus* yoba 2012 (Table 1). A comparison between the genomes of the four strains and the published genome of *L. rhamnosus* GG (9, 10) showed considerable genetic variety (Fig. 1). In two of three product isolates, major parts of the genomic islands LGGISL1,2 were missing, covering stretches of 34 genes and 84 genes, flanked by IS30 and IS5 insertion elements, respectively. The missing genes in both isolates include *spaCBA*, encoding pilin subunits involved in adhesion of the strains to the intestinal mucus and persistence in the human intestinal tract (10, 18). The DNA fragment of 34 genes missing from product isolate 1 includes

A

- Mutations in *L. rhamnosus* yoba 2012 (product isolate P3)
- Mutations in *L. rhamnosus* GG (product isolate P2A)
- Mutations in *L. rhamnosus* yoba 2010 (product isolate P1)
- Mutations in *L. rhamnosus* GG ATCC53103
- Transposases
- ▬ Deletion in product isolate 2A
- ▬ Deletion in product isolate P1
- ▬ Genomic Islands LGGISL1-5



B

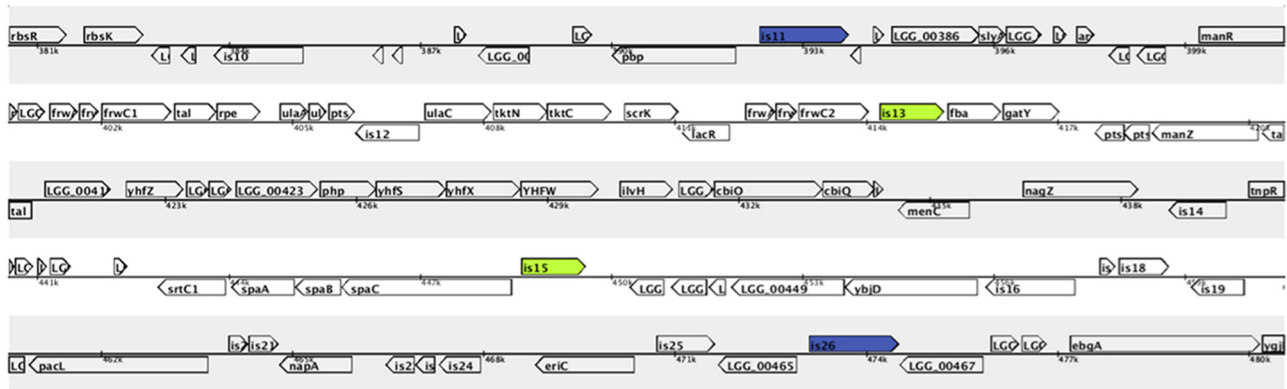


FIG 1 (A) Circular genome map of *L. rhamnosus* GG (accession number NC_013198.1). Colored circles indicate the loci of the mutations in the ATCC 53103 reference strain and three probiotic product isolates including *L. rhamnosus* yoba 2010 and 2012. The purple and blue bars indicate the sites of the missing ORFs of isolates P2A and P1, respectively. In gray are the genomic islands LGGISL1 to LGGISL5. Circles indicate from outside to inside: genes for the “+” strand; genes for the “-” strand; the GC percentage; green circles, all IS elements encoding transposases in the genome; genome positions (total length, 3,010,111 bp). (B) Linear map showing the deletions including stretches of 34 genes flanked by two transposases of the IS30 family (*is13* and *is15* in green) in product isolate 1 and 84 genes flanked by two transposases of the IS5 family (*is11* and *is26* in blue) in product isolate 2A.

two elements at both ends (*is13* and *is15*), which are 100% identical and both encode a 338-amino-acid transposase of the IS30 family (the entire genome contains four copies of this transposase gene). The DNA fragment of 84 genes missing from product iso-

late 2A also includes two elements at both ends (*is11* and *is26*), which are 100% identical and encode a 471-amino-acid transposase of the IS5 family (the entire genome contains 13 copies of this transposase gene). Strikingly, the LGGISL1,2 region contains

TABLE 2 Results of qPCR on DNA extracted from bacterial isolates from commercial products, control strain ATCC 53103, and *Lactobacillus rhamnosus* yoba 2012^a

Cultivation condition/sample no.	Product 2B				Product 4				ATCC 53103				Product 3/ <i>L. rhamnosus</i> yoba 2012			
	<i>map</i>		<i>spaC</i>		<i>map</i>		<i>spaC</i>		<i>map</i>		<i>spaC</i>		<i>map</i>		<i>spaC</i>	
	<i>C_T</i>	Quantity	<i>C_T</i>	Quantity	<i>C_T</i>	Quantity	<i>C_T</i>	Quantity	<i>C_T</i>	Quantity	<i>C_T</i>	Quantity	<i>C_T</i>	Quantity	<i>C_T</i>	Quantity
RA/1	17.5	3.0E+06	33.8	3.46E+02	18.0	2.3E+06	35.2	1.5E+02	17.5	3.11E+06	18.3	3.7E+06	17.6	2.9E+06	18.6	3.2E+06
RA/2	17.7	2.7E+06	34.0	3.00E+02	17.6	2.9E+06	18.5	3.4E+06	17.5	3.04E+06	18.4	3.6E+06	17.8	2.6E+06	18.7	2.9E+06
RA/3	17.5	3.0E+06	18.4	3.46E+06	17.5	3.0E+06	35.9	9.6E+01	17.5	2.96E+06	18.3	3.8E+06	17.7	2.7E+06	18.7	2.9E+06
RA/4	17.2	3.6E+06	32.5	7.53E+02	16.6	5.0E+06	17.5	6.0E+06	ND	ND	ND	ND	ND	ND	ND	ND
RN/1	17.4	3.2E+06	18.2	3.92E+06	27.2*	9.9E+03	29.8	3.8E+03	17.7	2.77E+06	18.5	3.4E+06	17.4	3.1E+06	18.3	3.7E+06
RN/2	17.3	3.4E+06	19.4	1.91E+06	27.2*	1.0E+04	30.5	2.4E+03	17.4	3.14E+06	18.3	3.9E+06	17.8	2.6E+06	18.6	3.1E+06
RN/3	17.7	2.7E+06	32.2	8.78E+02	17.2	3.6E+06	35.2	1.5E+02	17.6	2.81E+06	18.1	4.2E+06	17.6	2.8E+06	19.0	2.4E+06
RN/4	17.1	3.8E+06	34.0	3.02E+02	17.4	3.1E+06	OR		ND	ND	ND	ND	ND	ND	ND	ND
MA/1	17.4	3.3E+06	36.6	6.56E+01	17.7	2.6E+06	35.2	1.5E+02	17.6	2.94E+06	18.4	3.5E+06	17.5	3.0E+06	18.4	3.5E+06
MA/2	17.6	2.8E+06	36.0	9.14E+01	17.7	2.8E+06	OR		17.5	3.00E+06	18.4	3.5E+06	17.7	2.8E+06	18.6	3.2E+06
MA/3	17.6	2.9E+06	OR		17.7	2.6E+06	OR		17.2	3.68E+06	18.1	4.2E+06	17.7	2.6E+06	18.6	3.1E+06
MA/4	17.4	3.2E+06	37.5	3.65E+01	17.5	3.1E+06	34.8	1.9E+02	ND	ND	ND	ND	ND	ND	ND	ND
MN/1	17.3	3.4E+06	36.6	6.38E+01	ND	ND	ND	ND	17.4	3.27E+06	18.5	3.3E+06	17.3	3.4E+06	18.5	3.3E+06
MN/2	29.1*	3.2E+03	32.0	1.03E+03	ND	ND	ND	ND	17.8	2.46E+06	18.9	2.6E+06	17.8	2.5E+06	18.9	2.6E+06
MN/3	17.7	2.6E+06	38.6	1.90E+01	ND	ND	ND	ND	17.8	2.55E+06	18.9	2.6E+06	17.9	2.3E+06	19.0	2.4E+06
MN/4	OR*		OR		ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND

^a Abbreviations used cultivation conditions in column 1: R, Rogosa agar; M, MRS agar; A, aerobic; N, anaerobic. Other abbreviations: OR, out of range; ND, not determined. Quantities are expressed in fg/μl. *, the *C_T* of the control gene *map* is much larger than the average value of 17.5; hence, no mathematical comparison of the data is possible (see the text).

a very high density of insertion elements; for example, the deleted DNA sequence of 82 kb in isolate 2A contains 16 insertion elements (*is11* to *is26*), which is almost 10 times more than the average occurrence of these elements in the entire genome. Besides the 2 large deletions, a range of 9 to 23 deletions, duplications, substitutions, and point mutations (silent and nonsilent) were identified in the reference strain and the three product isolates (see Tables S2 to S4 in the supplemental material).

Genetic heterogeneity in commercial products. Two qPCRs were carried out targeting *spaC* and *map*, as a control gene, in order to investigate whether the large deletions, including the *spaCBA* genes, already had occurred in the *L. rhamnosus* GG strains present in the commercial products or later during the isolation of the variants by cultivation on selective nutrient agar. In a first series of experiments, DNA was directly isolated from seven commercial products. Six of these products contained *L. rhamnosus* GG according to the product label specifications, and one product contained a *Lactobacillus casei* strain (Table 1). It should be noted that two of these products also served as the source for the comparative genome analysis of product isolates 1, 2A, and 3, mentioned before (Table 1). The qPCR results confirmed the presence of the *map* and *spaC* genes in all products containing the LGG label. However, in two of these products the detected quantity of *spaC* genes appeared to be significantly lower than the *map* control gene. The ratios of *spaC* to *map* were 0.38 and 0.31 for products 2B and 4, respectively, while the average ratio in the four other products was 1.11 ± 0.15 . This indicates that these two products contain a heterogeneous population of *L. rhamnosus* GG variants consisting of genotypes with or without *spaC*.

In a follow-up experiment, a qPCR targeting *map* and *spaC* was conducted on 14 and 10 DNA samples extracted from a corresponding number of single colonies obtained from the commercial products 2B and 4, respectively. In order to evaluate whether environmental effects could select for a *spaC* minus genotype, colonies were selected after cultivation under aerobic and anaerobic conditions on MRS or RO agar. Positive controls included DNA isolates obtained from 12 colonies of cultivated ATCC 53103 strain and 12 colonies of product isolate 3/*L.*

rhamnosus yoba 2012, which, based on the genome sequencing results, had a genome very similar to the reference strain (Fig. 1 and Table 2). As negative controls, one sample without DNA and four samples with DNA extracts from a *Streptococcus thermophilus* isolate were included.

In total, 52 qPCRs were run on *L. rhamnosus* GG strains targeting *map* and *spaC*. Each qPCR was performed on DNA isolated from a unique colony. The *C_T* value of the calibration curve of *map* varied between 18.3 and 38.3, corresponding to 2.3×10^6 fg/μl and 23 fg/μl, respectively, with an *R*² value of 0.993. The *C_T* value of the calibration curve of the *spaC* varied between 19.1 and 34.1, corresponding to 2.3×10^6 fg/μl and 230 fg/μl, respectively, with an *R*² value of 0.997. The average *C_T* of 48 qPCRs targeting the *map* gene was 17.54 ± 0.24 , indicative of the highly reproducible DNA extractions and qPCR. The results of 4 of 52 qPCRs were excluded from further data analysis, because the corresponding *C_T* values of the *map* gene completely deviated from the control samples, suggesting that the amplification of the DNA had not run well, for instance due to poor DNA isolation and/or contamination of the sample. The *C_T* values of the control sample without DNA scored higher than 35, and the four samples with DNA extracts from a *Streptococcus thermophilus* isolate scored higher than 33 (data not shown).

Table 2 and Fig. 2 show that in all 24 colonies derived from the two positive control strains (i.e., ATCC 53103 and *L. rhamnosus* yoba 2012, originating from product isolate 3), the *spaC* and *map* genes were present in a ratio of ~1 (mean, 1.14; standard deviation, 0.12). This ratio confirms that *spaC* was present in all colonies after cultivation under the four described conditions. In contrast, the *C_T* value of *spaC* required at least 12 cycles more for 11 of the 14 colonies isolated from product 2B and 8 of the 10 colonies isolated from product 4 (Table 2), meaning that the *map/spaC* ratio in these colonies was >10,000. Since the source of the DNA for each qPCR was a single colony and the ratio *map* to *spaC* was ~1 in all control samples, this high ratio indicates the factual absence of *spaC* in 11 of 14 colonies and 8 of 10 colonies isolated from product 2B and 4, respectively. Figure 2A and B illustrate the ²log(*spaC:map*) ratio of the conducted qPCR for all samples derived from products 2B, 3, and 4 and the reference strain.

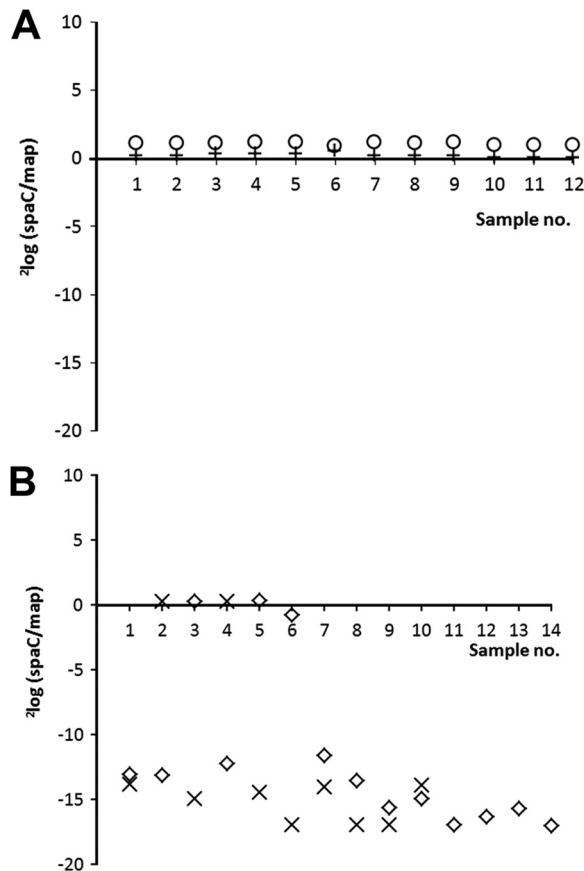


FIG 2 (A) qPCR derived $2\log(\text{spaC}/\text{map})$ ratio on DNA extracted from several isolates of control strain ATCC 53103 (+) and *L. rhamnosus* yoba 2012 (○). (B) qPCR-derived $2\log(\text{spaC}/\text{map})$ ratio on DNA extracted from several bacterial isolates of commercial product 2B (◇) and product 4 (×). The data are extracted from Table 2. Note: when the C_t value of *spaC* was below the detection limit, indicating that *spaC* was absent, then the $2\log(\text{spaC}/\text{map})$ ratio was arbitrarily set at 17.

As already indicated by the outcome of the comparative genome analysis (Fig. 1) for product 1 and 2A, the results from the qPCR analyses confirmed that the declared probiotic *L. rhamnosus* GG was not present either as a homogenous culture in products 2B and 4 (Table 1). In these products, the majority of the *L. rhamnosus* bacteria lack the pilin subunit-encoding *spaC* gene.

DISCUSSION

We conclude from our work that genetic alterations occurred in a major fraction of the population of probiotic *L. rhamnosus* GG bacteria present in some of today's commercialized products. A typical example of genetic alterations concerns the deletion of the genomic islands LGGISL1,2, including the *spaCBA* genes encoding multisubunit pilins with mucus-binding proteins (10, 18). Transmission electron microscopy using uranyl acetate staining confirmed the presence or absence of pili in strains with or without *spaC*, respectively (data not shown). Phenotypic analysis with *L. rhamnosus* yoba 2010 (product isolate 1) and product isolate 2A lacking these pili showed that these isolates had human mucus-binding abilities of only $1.6\% \pm 0.3\%$ and $1.4\% \pm 0.5\%$, respectively, compared to the *L. rhamnosus* GG reference strain (F. Douillard and W. M. de Vos, unpublished data). We reason that these

L. rhamnosus GG variants without *spaCBA* genes will have a reduced residence time in the gastrointestinal tract and even an altered probiotic functionality in case the claimed probiotic effect is related to adhesion of the strain to the intestinal mucus. Recently, the relevance of these genes for probiotic functionality was further demonstrated by showing that the SpaCBA pili could directly induce anti-inflammatory pathways or indirectly lead to anti-inflammatory signaling by promoting the release of anti-inflammatory factors such as the MSP1 and MSP2 soluble proteins (18).

The deleted regions observed in *L. rhamnosus* GG product isolates 1 and 2A are completely overlapping and comprise DNA stretches of 34 genes (LGG_00412 to LGG_00445) and 84 genes (LGG_00383 to LGG_00466), respectively. They both appear to overlap with the genomic islands LGGISL1 (LGG_00393 to LGG_00427) and LGGISL2 (LGG_00433 to LGG_00450), previously identified as DNA sequences deviating in codon usage, base composition, and dinucleotide frequency from the rest of the genome, suggesting that they originated from horizontal gene transfer. Typical functions of the genomic island LGGISL1 include the capacity to transport or metabolize sugars, while LGGISL2 island appears to encode a set of genes for multisubunit pili (SpaCBA) and a sortase, required for the assembly of pilus structures (10).

The deleted DNA segments were both flanked by two identical insertion elements, encoding IS30 transposases in one product isolate and IS5 transposases in the other. Although ISs are best known to be involved in acquisition of accessory functions, these elements are also known to be involved in chromosomal deletions (7). Since it was found that both missing stretches of DNA were flanked by two identical copies of ISs, they could be involved in the deletions, possibly through the activity of a composite transposon. In case the result of this genetic alteration leads to increased fitness under the conditions used for cultivation, there will be an enrichment for bacterial genotypes without (part of the) genomic islands, as has been observed in the present study for some of the dairy products.

Cai et al. reported that gene decay in lactobacilli is known to be associated with the transition from dynamic and nutritionally variable environments, such as the human gastrointestinal tract and plants, to the relatively constant and nutrient-rich dairy niche (5). This event has also recently been simulated in a laboratory experiment (19). Hence, propagation of strains isolated from the intestinal gut, as is the case for many probiotics, including *L. rhamnosus* GG, likely results in metabolic simplification and loss of genes, which could be linked to probiotic functionality. A practical example refers to the industrial processing of probiotics, when strains are grown for several generations in relatively large volumes (20) without selective pressure for maintenance of genes with probiotic functionality.

We reason that from a genome stability point of view, future research should evaluate whether traditionally fermented dairy products, where the lactobacilli have been cultivated for numerous generations under nutrient-rich conditions, could be a better source of selection for large scale production of genetically stable probiotic strains than the human gastrointestinal tract. Typically, this was the approach followed by Elia Metchnikoff more than a 100 years ago (21). As a drawback of this approach, one could reason that some specific probiotic functionalities may only be found in isolates from the intestinal tract, illustrating the need for innovative cultivation methods that keep selective pressure on persistence of genes, which are important for health-enhancing

microbe-host interactions. In the context of the present study, one might think of growth conditions that maintain selective pressure on the presence of carbohydrate transporting and metabolizing genes encoded by LGGISL1. In addition, it might be worthwhile to avoid exposure to stress conditions that might activate transposons (7). However, pretreatment of cells under nonmutagenic conditions (42°C) appeared to protect them against subsequent increases in the mutation frequency due to oxidative stress, showing that sequential exposure to certain stresses may also be used to protect industrial strains against elevated mutation rates and in this way potentially enhance the stability of these strains (22).

Although we describe here the occurrence of genetic instability, including mutations and deletions of several genes, it should also be noted that the overall genome similarity between the four *L. rhamnosus* GG variants was >97%, including deleted regions and >99.9% when excluding deleted regions. Even though a single point mutation might lead to a loss of functionality, when checking the mutations and deletions presented in Tables S2 to S4 in the supplemental material, one may conclude that many of the other probiotic functionalities are still present; these include exopolysaccharide production, tolerance to bile salts, stimulation of the immune system, and also all of the probiotic effects beyond the gut, as reviewed by Lebeer et al. (3). Two specific examples to mention are the *L. rhamnosus* GG secreted proteins MSP1 and MSP2, which have been reported to enhance survival and growth of intestinal epithelial cells and which are still encoded in all of the four sequenced genomes (23). Interestingly, the corresponding genes for these proteins are supposed to remain genetically stable because the secreted proteins are also involved in essential cell metabolic activities such as cell separation (24).

Our results contribute to the debate about both regulatory and scientific aspects of functional claims linked to probiotic products and required quality control measures. The European Food and Safety Authority (EFSA) has scientifically evaluated numerous article 13.1 health claims. The NDA Panel of the EFSA considered that in most of the cases the information provided was not sufficient to characterize a number of foods or constituents with respect to the claimed effects, including some (but not all) probiotic bacteria (25). The NDA Panel proposed species identification by DNA-DNA hybridization or 16S rRNA gene sequence analysis and/or sequence analysis of other relevant genetic markers. Strain identification is proposed by pulsed-field gel electrophoresis of genomic DNA, randomly amplified polymorphic DNA analysis, or other internationally accepted genetic typing molecular methods (26). We reason that even EFSA-proposed strain-specific identification methods such as DNA fingerprinting or other methods, such as enzyme-linked immunosorbent assays, the use of monoclonal antibodies, or strain-specific PCR (27), are not sufficient to guarantee that strains with a claimed functionality are effectively present in probiotic products. Furthermore, and as a general statement, we think that products containing probiotic strains that have a significantly altered genetic content or functionality should be tested in trials to confirm their efficacy. However, it should be noted that bacteria are living entities that always evolve. The suggestion that products cannot be sold because of the potential occurrence of minor genetic alterations as part of an evolutionary process would make it probably impossible to ever sell a probiotic.

Until the impact of genetic instability in general and the absence of pili in particular on probiotic functionality is well under-

stood, we propose a quality assurance approach involving validation steps in the production and release processes of probiotic products. The validation is aimed to confirm the genetic stability of the overall genome, especially around mobile genetic elements that could induce the simultaneous deletion of several genes. In addition, it remains relevant to confirm that strains, which are marketed today, have the same genetic makeup as the strains used in clinical studies on which their health claims have been based. Hereto, and as a starting point for further research, we propose to check whether the results presented here could provide a complementary explanation for the variation of probiotic properties (based on *in vitro* assays) observed in 16 *L. rhamnosus* GG product isolates and suggested to be linked to different processing conditions (28).

Finally, we note that when strains such as *L. rhamnosus* GG are genericized, as has recently been reported (1), it is important that they possess the characteristics that make them efficacious; otherwise, people relying on them will not benefit.

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ADDENDUM IN PROOF

During the review of this paper, there have been two more reports of the genetic (in)stability of probiotics. Douillard et al. (Appl. Environ. Microbiol. 79:1923–1933, 2013) found that the genomes of two *L. rhamnosus* GG strains isolated from two different products are virtually identical. Averina et al. (Russ. J. Genet. 48:1103–1111, 2012) noted the loss of the *galA* and *tet(W)* genes from the genome of *Bifidobacterium longum* subsp. *longum* B379M during cultivation and maintenance under laboratory conditions. Interestingly, *tet(W)* is located between two *IS30* elements, as is described for one of the deletions observed in the present research paper. The latter results indicate that the genetic instability of probiotics may be a rather widespread phenomenon, which further supports a recommendation that regulating authorities, the food industry, and academia focus on controlling the retention of key probiotic genes.

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