Comparative genomics of *Lactobacillus reuteri* from sourdough reveals adaptation of an intestinal symbiont to food fermentations

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*Lactobacillus reuteri* is a dominant member of intestinal microbiota of vertebrates, and occurs in food fermentations. The stable presence of *L. reuteri* in sourdough provides the opportunity to study the adaptation of vertebrate symbionts to an extra-intestinal habitat. This study evaluated this adaptation by comparative genomics of 16 strains of *L. reuteri*. A core genome phylogenetic tree grouped *L. reuteri* into 5 clusters corresponding to the host-adapted lineages. The topology of a gene content tree, which includes accessory genes, differed from the core genome phylogenetic tree, suggesting that the differentiation of *L. reuteri* is shaped by gene loss or acquisition. About 10% of the core genome (124 core genes) were under positive selection. In lineage III sourdough isolates, 177 genes were under positive selection, mainly related to energy conversion and carbohydrate metabolism. The analysis of the competitiveness of *L. reuteri* in sourdough revealed that the competitiveness of sourdough isolates was equal or higher when compared to rodent isolates. This study provides new insights into the adaptation of *L. reuteri* to food and intestinal habitats, suggesting that these two habitats exert different selective pressure related to growth rate and energy (carbohydrate) metabolism.

*Lactobacillus reuteri* persist in intestinal microbiota of vertebrate animals as well as in food fermentations1–4. *L. reuteri* colonizes humans and animal hosts2,4; the phylogenetic differentiation of strains of *L. reuteri* originating from different hosts reflects co-evolution of *L. reuteri* with its vertebrate hosts4. This evolutionary adaptation differentiates the species *L. reuteri* in host-adapted phylogenetic lineages comprised of isolates from rodents (lineages I and III), humans (lineages II and VI), pigs (lineages IV and V), and poultry (lineage VI)4,5.

*L. reuteri* also occur in industrial sourdoughs6 and cereal fermentations in tropical climates1,7. Sourdoughs are typically maintained by continuous propagation, a process which rapidly selects for the most competitive microbiota. Major selection criteria for fermentation microbiota in cereal ecosystems are rapid growth in cereal substrates, and acid resistance1,8–10. Food isolates of *L. reuteri* match to host-adapted lineages11 and maintain host-specific physiological traits12–14, including the ability to colonize the lineage-specific hosts11,15.

The differentiation of *L. reuteri* into host-adapted lineages implies that an extra-intestinal habitat did not exist for a majority of the evolution of this species14. However, the occurrence of *L. reuteri* in the human-made habitat sourdough provides the opportunity to study the “reverse adaptation” of vertebrate symbionts to an extra-intestinal habitat. This study employed comparative genomics of *L. reuteri* to evaluate the genetic determinants of this adaptation or selection process. Genome sequences of intestinal strains of *L. reuteri* were retrieved from public databases and compared to four genome sequences of rodent-lineage sourdough isolates16. The sourdough isolates *L. reuteri* LTH2584, TMW1.112 and TMW1.656 originate from SER sourdough, a sourdough that is used industrially for production of a baking improver8. This sourdough has been maintained by continuous propagation since about 1970. *L. reuteri* LTH2584, TMW1.112 and TMW1.656 were isolated from this sourdough in 1988, 1994, and...

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19986,9; all of these strains produce reutericyclin, a tetramic acid derivative with antimicrobial activity against Gram-positive bacteria6,16. L. reuteri LTH5448 was isolated from a different sourdough processed at the same facility in 20008,17; this strain does not produce reutericyclin but maintains the reutericyclin genomic island and reutericyclin resistance16,17. Comparative genomics analyses included analyses of the core genome as well as gene gain and gene loss events that were studied on the basis of the pan-genome. We also performed positive selection analysis for these core genes of the whole species. Finally, the competitiveness of sourdough isolates of L. reuteri in model sourdoughs was compared to the competitiveness of closely related intestinal isolates.

Results
Phylogenetic analysis of 16 sequenced L. reuteri strains including 4 sourdough strains. The phylogenetic analysis was carried out with all available genome sequences of L. reuteri, including 4 genome sequences of sourdough isolates16. A phylogenetic tree was constructed based on the core genome of L. reuteri (Fig. 1A). Strains of L. reuteri were grouped into 5 clusters corresponding to the host-adapted lineages I (rodent), II (human), III (rodent), IV (pig) and VI (poultry and human). Sourdough strains were assigned to the rodent-adapted lineages I and III, in agreement with previous analyses11. L. reuteri LTH5448 clustered with lineage I rodent isolates; L. reuteri LTH2584, TWM1.112 and TWM1.656 were grouped into lineage III together with the rodent isolates L. reuteri 100-23 and mlc3. L. reuteri LTH2584, an SER sourdough isolate obtained in 1988, was more closely related to L. reuteri TWM1.656, which was isolated from SER sourdough in 1998, than to L. reuteri TWM1.112, which was isolated from the same sourdough in 19946.

A gene content tree was constructed to study the gain and loss of genes among these strains. Here, strains sharing more genes were clustered together (Fig. 1B). The topology of the gene content tree was different from the core genome phylogenetic tree, indicating gene loss or acquisition of genes by horizontal genetic transfer. Three clusters corresponding to lineages II, IV and VI were maintained but the gene content tree highlighted differences between strains in each cluster. For example, the four lineage II L. reuteri MM4-1A, MM2-3, DSM 20016 and JCM 1112 were not separated in the core genome phylogenetic tree but differentiated in two groups by calculating the gene content tree (Fig. 1B). L. reuteri DSM20016 and JCM1112 were derived from the same original isolate, F275, and differences between these two strains may reflect loss of genes during propagation in the laboratory18. The two lineage III strains L. reuteri 100-23 and mlc3 showed a quite different gene content. Remarkably, all four sourdough isolates were grouped together despite their divergent phylogenetic origin. L. reuteri LTH5448 was more closely related to L. reuteri LTH2584 than to L. reuteri TWM1.112 and TWM1.656.

Comparative analysis of sourdough strains. To understand how the intestinal strains adapted to sourdough, and to identify genes that are unique to sourdough isolates, the gene content similarity and dissimilarity of these strains was analysed. L. reuteri LTH2584, TWM1.112, TWM1.656 and 100-23 shared 1535 core genes (Fig. 2A); this core genome is higher than the core genome of the whole species, reflecting that all these strains are grouped in lineage I. L. reuteri LTH2584 and 100-23 had more unique genes than L. reuteri TWM1.112 and TWM1.656 (Fig. 2A), which contributed to the distinct position of the former two strains in the gene content tree. Sourdough isolates shared 1523 core genes (Fig. 2B). Genes that were shared by all sourdough isolates but absent in
other strains include the chromosomally encoded reutericyclin genomic island, a putative aspartate racemase, a LytTr-domain protein with putative regulatory function, and components of a putative ABC-transporter (Table 2). Genes that were only present in some of the sourdough isolates include a glycoyltransferases with putative function in protein glycosylation (LTH2584 and TMW1.656) and a putative hydroxyglutarate dehydrogenase (LTH2584 and TMW1.112) which catalyses the use of α-ketoglutarate as electron acceptor. Of note, distributed genes that are present in sourdough isolates of _L. reuteri_ and other strains include several putative enzymes of the shikimic acid pathway for biosynthesis of aromatic amino acids (Table 2). In summary, only genes coding for reutericyclin biosynthesis are unique to all sourdough isolates of _L. reuteri_.

**Figure 2.** Comparative analyses of genes shared between sourdough isolates of _L. reuteri_ and _L. reuteri_ 100-23, and among sourdough isolates. (A) Venn diagram of core, distributed and unique gene numbers between the lineage III strains _L. reuteri_ LTH2584, TMW1.656, TMW1.112, and 100-23. (B) Venn diagram of core, distributed and unique gene numbers among the Lineage III sourdough isolates _L. reuteri_ LTH2584, TMW1.656, TMW1.112, and the Lineage I sourdough isolate _L. reuteri_ LTH5448.

Positive selection of the core genes contributing to the adaptation of sourdough isolates. Analysis of positive selection aimed to identify the selective pressure on the core genome of _L. reuteri_, and to determine whether sourdough and intestinal strains are subjected to a differential selective pressure. Initially, positive selection was analysed in all 16 strains of _L. reuteri_. A total of 124 core genes were under positive selection (Fig. 3 and Table S2), representing 10.36% of the core genome. Among the genes that are under positive selection, 22% relate to metabolism, including transporters and enzymes for protein, amino acid, carbohydrate, and lipid conversion. Several genes under positive selection were listed as “general functional prediction only”, but most of these predicted functions were also related to metabolic functions. Other abundant genes under positive selection relate to DNA replication, recombination, and repair. When compared with the composition of the core genes, COG categories “translation, ribosomal structure and biogenesis” and “general functional prediction only” were significantly enriched among genes under positive selection in all 16 core genomes of _L. reuteri_ (P = 0.04, 0.03, one-sided binomial test). For the 20 genes in the former category, 8 are tRNA associated genes, 6 are ribosomal protein genes, 3 are 23S RNA-specific pseudouridylate synthases, 2 are translation elongation factor genes, and 1 is methylase of polypeptide chain release factor gene (Table S2). For the latter category, most predicted functions relate to metabolism. For example, 6 out 21 were hydrolases, and other were some reductases, permeases and esterases.

To identify the selective pressure acting on the sourdough isolates, the branch-site model and its null model were used to compare the function categories under positive selection in the branch comprising _L. reuteri_ LTH2584, TMW1.112, and TMW1.656 to all other strains in the species. A total of 177 core genes were under positive selection in these lineage III sourdough isolates (Fig. 3 and Table S3). Of these, 135 genes were under positive selection only in this branch and the remaining 42 were under positive selection in the sourdough isolates as well as the remainder of the species. Of the core genes under positive selection in the sourdough branch, 33% related to metabolism (Fig. 3 and Table S3). Three COG categories were significant enriched in the lineage III sourdough isolates, “Energy production and conversion” (P = 9.9 * 10^-9), “Carbohydrate transport and metabolism” (P = 0.03) and ”Defense mechanisms” (P < 2.2 * 10^-16) (Fig. 3). Examples of gene in these COG categories that are under positive selection include key metabolic enzymes such as maltose phosphorylase, lactate dehydrogenase, alcohol dehydrogenase, and several sugar transport enzymes (Table S3).

**Competitiveness of _L. reuteri_ strains in sourdough: experimental design.** To determine whether genomic adaptation of _L. reuteri_ to the sourdough environment increases the competitiveness of strains, competition experiments sourdoughs were carried out. Competition experiments in back-slopped sourdoughs are a sensitive tool to determine the competitiveness of strains because even small differences in competitiveness result in predominance of the more competitive strain after few refreshments. Experiments were performed with fermentation cycles of 1, 2, or 3 days. The selection of strains used in the competition experiments included the sourdough isolates _L. reuteri_ LTH5448, LTH2584, TMW1.112 and TMW1.656; and the rodent isolates...
This study employed comparative genomics to demonstrate that the evolution of L. reuteri is shaped by positive selection of the core genome in addition to the gain and loss of accessory genes. Moreover, the identification of core genes under positive selection and the analysis of competitiveness in sourdough demonstrate that sourdough selection of the core genome in addition to the gain and loss of accessory genes. Moreover, the identification of core genes under positive selection and the analysis of competitiveness in sourdough demonstrate that sourdough selection of the core genome in addition to the gain and loss of accessory genes. Moreover, the identification of core genes under positive selection and the analysis of competitiveness in sourdough demonstrate that sourdough selection of the core genome in addition to the gain and loss of accessory genes. 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strains *L. reuteri* LTH2584, TWM1.112, and TWM1.656 were isolated from the same sourdough in 1988, 1994, and 1998, respectively. The phylogenetic relatedness of these strains suggested that the later isolates may be isolates of the same organism after 10 years of adaptation to sourdough fermentation. However, successive contamination of the same sourdough with different strains of rodent origin is an alternative explanation for the isolation of highly related strains from the same sourdough.

**Role of reutericyclin production for competitiveness in sourdoughs.** Reutericyclin production may contribute to competitiveness of *L. reuteri* in sourdough. This study demonstrated that a reutericyclin sensitive derivative of *L. reuteri* TMW1.656 and the reutericyclin producing wild type strain exhibited comparable competitiveness in sourdough, indicating that the ecological advantage of reutericyclin is about equivalent to the cost of reutericyclin production. The reutericyclin gene cluster was acquired by horizontal gene transfer by few Lineage I and Lineage III sourdough isolates of *L. reuteri* and is thus unlikely to represent a sourdough-specific metabolic trait.

**Evolution of the intestinal symbiont *L. reuteri* by horizontal gene transfer and positive selection.** The evolution of pathogens is driven by gene loss, acquisition of genes by horizontal gene transfer, and by positive selection of the core genome. Only few genes were reported to be under positive selective pressure in the pathogenic *Listeria monocytogenes* while up to 34% and 92%, respectively, of the core genome were positively selected in specific lineages of the host-adapted genera *Streptococcus* and *Campylobacter*. The evolution of the host adapted gut symbiont *L. reuteri* was previously attributed to gene loss and acquisition of lineage-specific accessory genes. The congruent clustering of *L. reuteri* strains in the phylogenetic tree and the gene content tree confirms a major role of the gain or loss of host-specific metabolic and genetic traits in the evolution of the species. This study additionally demonstrates that positive selection of the core genome shapes the evolution of *L. reuteri*. The expression of ribosomal proteins, rRNA and other transcription factors is regulated by the bacterial growth rate and a high density of ribosomal genes relates to rapid growth of *L. sanfranciscensis* in sourdough. Positive selection in the functional category translation indicates that the intestinal ecological niches harbouring *L. reuteri* exert selective pressure for rapid growth. The proportion of the core genome that was under positive selection in *L. reuteri* matches the corresponding proportion in *S. mutans*. *S. mutans* and *L. reuteri* are phylogenetically related host-adapted organisms colonizing the upper intestinal tract but contrast with respect to their impact on the hosts. *S. mutans* is a pathogen and *L. reuteri* is considered as probiotic but both species apparently use comparable ecological strategies for colonization and persistence.

![Figure 3](image_url)

**Figure 3. Proportions of positive selection of core genes in each COG category in all *L. reuteri* strains (gray bars), and the branch of sourdough isolates comprising *L. reuteri* LTH2584, TWM1.112, and TWM1.656 (white bars).** Site model M2a and its null model M1a were compared to infer genes under positive selection in the whole species. Branch-site model and its null model were compared to study genes under positive selection across the COG categories. COG categories that were significantly enriched in all strains of the species *L. reuteri*, or in the Lineage III sourdough isolates are marked by an asterisk.
Reverse evolution or selection of *L. reuteri* in an extra-intestinal habitat? The persistence of *L. reuteri* over 10 years in sourdough fermentation provides a unique opportunity to study the adaptation or selection of a host-specific gut symbiont to an extra-intestinal environment. Several lines of evidence suggest that SER sourdough isolates are distinct from intestinal strains. First, sourdough isolates cluster separately in the gene content tree, indicating that horizontal gene transfer and the loss of genes relates to the transition to sourdough\(^{16}\). Second, the functional categories “energy production and conversion” and “carbohydrate metabolism”, which are key elements for competitiveness in sourdough\(^{28,29}\), were significantly enriched among the positively selected genes in SER isolates. Third, sourdough isolates of *L. reuteri* displayed a higher relative fitness in sourdough when compared to rodent isolates. *L. reuteri* LTH5448 also achieved a high proportion of cell counts in competition with rodent isolates *L. reuteri* mlc3 and lpuph in experiments with 2 d fermentation time (data not shown). The differences in competitiveness of rodent and sourdough isolates, however, are smaller than differences between individual strains, reflecting the relatedness of the rodent forestomach and sourdough environments\(^{30}\). Because the time between sourdough contamination with *L. reuteri* and the isolation of the specific strains is unknown, it is not possible to discriminate whether the specific differences of the sourdough isolates reflect selection for a specific subset of rodent isolates, or “reverse evolution” of a gut symbiont to food fermentations.

In conclusion, this study demonstrated that gene loss and gene gain as well as selective pressure on the core genome drive the evolution of *L. reuteri*. Remarkably, the gene content of sourdough isolates of *L. reuteri* differed from intestinal isolates, and genes under positive selection in sourdough strains included maltose phosphorylase, alcohol dehydrogenase, and lactate dehydrogenase, genes which are known to contribute to competitiveness in cereal fermentations. The study improves our understanding of the adaptation of bacteria to food fermentations as an evolutionary recent man-made habitat. It will also improve our ability to use food fermentations as model systems for more complex, intestinal ecosystems\(^{31}\).
Methods

Strains, media and growth conditions. The sourdough isolates \textit{L. reuteri} LTH2584, TMW1.112, TMW1.656 and LTH5448\textsuperscript{9,17} and the rodent isolates \textit{L. reuteri} 100-23, mlc3, and lpuph\textsuperscript{32} were grown anaerobically at 37°C in mMRS\textsuperscript{8}. Sugars were autoclaved separately. Solid media contained additional 20 g agar per liter.

Whole-genome alignment and phylogenetics. Genome sequences of the 12 \textit{L. reuteri} were retrieved from Genbank (Table S1). Genome sequences of sourdough isolates\textsuperscript{16} were re-annotated on the RAST server\textsuperscript{33}.
after gap closing by PCR amplification and Sanger sequencing. Primers binding to up- and down-stream locus of the target gap were selected after alignment of the genomes with Mauve\(^{41}\), and are shown in Table S1. Sequencing was performed by service of Macrogen Co. (Rockville, Maryland, USA).

All 16 genomes were aligned with Mugsy\(^{35}\). Homologous blocks present in each genome were concatenated with an in-house perl script. The most disordered regions were eliminated using GBLOCKS\(^{39}\). The disordered regions includes sites containing at least one gap, and sites that are too divergent as they may not be homologous or may be saturated by multiple substitutions. The core genome size of \textit{L. reuteri} was about 1.2 Mbp. A maximum-likelihood core genome tree was constructed using RaxML\(^{37}\). The tree was inferred under the general time-reversible nucleotide substitution model (GTR), with gamma-distributed rate heterogeneity of four rate categories (+I+G) (Γ4).

Bootstrap support values were calculated from 1000 replicates.

\textbf{Gene clustering and construction of a gene content tree.} Protein sequences longer than 50 amino acids from all genomes were combined and searched using BLAST with an all-against-all style with default parameters. The protein sequences with identities and coverage above 70% were clustered into families using the program orthoMCL\(^{38}\). The inflation value of 2 was used for the MCL clustering. Core genes were defined as those shared by all of the 16 strains; distributed genes as those shared by 2 to 15 strains, and unique genes as those only contained in one strain.

A matrix of the presence or absence of each gene for each genome was created. A dissimilarity distance between genomes based on gene content (binary data for presence or absence of each protein family) measured by one minus the Jaccard coefficient (Jaccard distance) was calculated from this matrix\(^{39}\). A gene content tree was constructed using the hierarchical clustering (UPGMA) method based on these distances by MEGA\(^{40}\).

\textbf{Analysis of positive selection.} For each cluster of the single-copy core genes, protein sequences were aligned with MUSCLE\(^{41}\). These alignments were reverse-translated to codon-based nucleotide alignments by PAL2NAL\(^{42}\). Positive selection analysis based on each of these alignments was performed by CODEML implemented in PAML\(^{43}\). Nonsynonymous (amino acid altering) synonymous (silent) substitution ratios ($\omega$), with $\omega = 1$, $< 1$, or $> 1$ indicate neutral, purifying, or positive selection, respectively. Positive selection was analysed on each family of core genes shared by all 16 \textit{L. reuteri} isolates using the site models M1a and M2a\(^{44,45}\). The model M1a (nearly neutral) allows all sites to be purifying selection ($\omega_0 < 1$) or neutral selection ($\omega_0 = 1$); the model M2a allows all sites to be positive selection ($\omega_0 > 1$). A likelihood ratio test (LRT) was carried out to infer the occurrence of sites subject to positive selective pressure through comparing M1a against M2a. Branch-site model and the one-ratio null model were used to analyze positive selection across the \textit{L. reuteri} LTH2584/TWM1.112/TWM1.656 branch. Branch-site model allows $\omega$ to vary both among sites in the protein and across branches on the tree and aim to detect positive selection affecting a few sites along particular lineages (called foreground branches)\(^{46}\). Two models were used, the null model does not allow positive selection for the foreground branch, and the alternative model assumes that the foreground branch may have some sites under positive selection. For the alternative model, three classes of $\omega$ were defined: $\omega < 1$, $\omega = 1$ and $\omega > 1$, while in the null model, $\omega$ was fixed to 1. A likelihood ratio test (LRT) was carried out to infer positive selective pressure across the \textit{L. reuteri} LTH2584/TWM1.112/TWM1.656 branch through comparing the results from these two models. The LRT statistic (twice the log-likelihood difference between the null and the alternative models) was compared with the chi-square distribution with 2 degrees of freedom for M2a vs. M1a, and one degree of freedom for branch-site model vs. the null model.

For Clusters of Orthologous Groups of proteins (COG) analysis, we constructed a local COG database\(^{47}\), and then ran rpsblast using the sequence sets mentioned above as queries. We focused on the top three hits of each alignment and counted each category for comparison using in-house Perl script.

\textbf{Competitiveness of \textit{L. reuteri} in sourdough: experimental design.} The persistence of strains was analyzed in back-slopped rye sourdough fermentations; experiments were carried out with fermentation times of 1, 2, and 3 days. Competition experiments were carried out with six strain combinations; sourdoughs were inoculated with \textit{L. reuteri} LTH2584 and 100-23; \textit{L. reuteri} LTH5448 and 100-23; \textit{L. reuteri} LTH5448 and mlc3; \textit{L. reuteri} LTH5448 and lpuph; \textit{L. reuteri} LTH2584 and LTH5448; or \textit{L. reuteri} LTH2584, TMW1.112, TMW1.656, and LTH5448.

\textbf{Sourdough preparation and differential enumeration of cell counts.} Competition experiments in sourdough were performed essentially as described\(^{49}\). In brief, sourdough was prepared by mixing 10 g rye flour with 10 ml of autoclaved tap water and 1 ml of bacterial inoculum. For binary and quaternary strain combinations, 0.5 and 0.25 mL, respectively, of the cell suspensions of individual strains were mixed to obtain 1 mL of bacterial cocktail as inoculum. Dough was fermented at 37°C for 1, 2, or 3 days and back-slopped over 10 fermentation cycles. At each back-slopping step, 1 g of ripe sourdough from the previous cycle was mixed with 9.5 g of fresh rye flour and 9.5 ml of autoclaved tap water. The competition experiments were performed in duplicate and analyses were carried out with two technical replicates.

At each fermentation cycle, sourdoughs were analysed with respect to the pH, differential cell counts, and qPCR with strain specific primers. Viable cell counts were enumerated by surface-plating of appropriate dilutions on mMRS agar. Individual strains were differentiated on the basis of the colony morphology. Differential enumeration was possible for the binary strain combinations \textit{L. reuteri} LTH2584 vs. 100-23, \textit{L. reuteri} LTH2584 vs. LTH5448, \textit{L. reuteri} LTH5448 vs. mlc3, and \textit{L. reuteri} LTH5448 vs. lpuph, but not for \textit{L. reuteri} LTH5448 vs. 100-23. In the quaternary strain combination, the combined total of \textit{L. reuteri} LTH2584, TMW1.112 and TMW1.656 was differentiated from \textit{L. reuteri} LTH5448.
Analysis of sourdough microbiota by qPCR. Total DNA was isolated from sourdough48 and gene copy numbers of _L. reuteri_ were quantified by strain-specific qPCR. Strain-specific primers are listed in Table 1. Standard curves to convert detection threshold cycles to gene copy numbers were established by analysis of 10-fold serial dilutions of target DNA of known concentration. The strain-specific primers to for strain specific quantification are shown in Table 1.

Calibration curves to convert gene copy numbers to cell counts in sourdough were Calibration curves to convert gene copy numbers to cell counts in sourdough were established with sourdoughs that were fermented with single strains. Samples were mixed with 2 volumes of sterile saline, and serially diluted with saline. From each of the dilutions, cell counts were determined by further dilution and surface plating and the gene copy numbers were quantified by qPCR as described above. Calibration curves were established in duplicate after 1 and 3 d of fermentation (Figure S2).

Quantitative PCR analyses were carried out in duplicate in MicroAmp Fast Optical 96-well reaction plates capped with MicroAmp Optical Adhesive Film (Applied Biosystems, Burlington, ON, Canada). The PCR reaction mixture consisted of 12.5μl Fast SYBR Green Master Mix (Applied Biosystems), 0.4μM of each primer (Table S1), 2μl of template DNA and sterile Milli-Q water to final volume of 25μl. Melting curves were obtained by a stepwise increase of the temperature from 60 to 95 °C at 0.05 °C/s) and melting-curve data were analyzed to verify amplification of the correct targeted PCR products. The detection limit was 10^2 copy numbers/g sourdough for the strain-specific primers.

### Table 2. Distributed genes specific to sourdough strains. Hypothetical proteins and phage-related proteins were excluded from the list. Protein Accession numbers refer to the genome of _L. reuteri_ LTH2584 unless otherwise specified.

| Gene or gene cluster | (Putative) function [ref] |
|----------------------|----------------------------|
| **Exclusive to all sourdough strains** | |
| _L. reuteri_ LTH2584, TMW1.112, TMW1.656, and LTH5448 | |
| Reutericyclin genomic island | Reutericyclin biosynthesis and resistance [16] |
| Components of an ABC transporter | Unknown [16] |
| Aspartate racemase (WP_003670574.1), | |
| LytTr DNA-binding domain (WP_006729038.1) | Unknown |
| **Exclusive to some sourdough strains:** | |
| _L. reuteri_ LTH2584 and TMW1.112 | |
| GntR (WP_006916030.1), | |
| Membrane transport protein (WP_006916028.1), | Unknown |
| Hydroxyglutarate dehydrogenase (WP_006916027.1) Use of ketoglutarate as electron acceptor; | |
| **L. reuteri_ LTH2584 and TMW1.656** | |
| Two GT8_A4GalT_like proteins (WP_020807754.1, WP_020807753.1) | Bread spectrum glycosyltransferases with putative function in protein glycosylation; |
| YkuD (WP_003664366.1) | peptidoglycan crosslinking |
| **Predominantly sourdough strains** | |
| _L. reuteri_ TMW1.112, TMW1.656 and mlc3 | Putative components of shikimate pathway for biosynthesis of aromatic amino acids |
| _L. reuteri_ TMW1.112, LTH5448 and 100-23 | |
| Homocysteine methyltransferase (100_23|ZP_03072304) | Oxidative stress response |
| S-methylmethionine transporter (100_23|ZP_03072305) | |
| **L. reuteri_ LTH2584, TMW1.112, TMW1.656, and 100-23** | |
| Lr100-23|ZP_03073418.1 | NADPH-dependent FMN reductase, rRNA |
| Lr100-23|ZP_03073416.1 | methyltransferase |
| Lr100-23|ZP_03073414.1 | Lipid metabolism |
| Lr100-23|ZP_03073413.1 | serine protease |
| Lr100-23|ZP_03073412.1 | Metal-dependent beta-lactamase superfamily |
| Lr100-23|ZP_03073411.1 | unknown |
| Lr100-23|ZP_03073410.1 | unknown |
| Lr100-23|ZP_03073409.1 | Histidine kinase |
| Lr100-23|ZP_03073446.1 | Cardiolipin synthase |
| Lr100-23|ZP_03073445.1 | Acetyltransferase (GNAT) family |

[^48]: DNA isolation from sourdough
Competitiveness of isogenic reutericyclin-positive and reutericyclin-negative and reutericyclin-sensitive isogenic strains of \textit{L. reuteri}. Competition experiment between reutericyclin-positive wild type strain \textit{L. reuteri} TMW1.656 and its reutericyclin-susceptible mutant \textit{L. reuteri} TMW1.656Δ\textit{ArcN}\Delta\textit{ArcT} \textsuperscript{70} were performed using white wheat flour with a dough yield of 200. Sourdough was propagated every 24h with 1% inoculum for 5 days. The ratio of wild type to mutant strains at the end of each fermentation cycle was determined by qPCR with primers listed in Table 1.

Calculation of the relative fitness of \textit{L. reuteri} in sourdough. The differential cell counts and the strain-specific gene copy numbers were used to calculate the relative fitness of the respective strains of \textit{L. reuteri}. The fitness (\(w\)) of strain \(x\) relative to that of strain \(y\) was calculated based an equation derived from \textsuperscript{49}:

\[
w(x, y) = \ln \frac{x_f / x_0}{y_f / y_0}
\]

where \(x_0\) and \(y_0\) denote the strain specific cell densities or gene copy numbers at the beginning of each fermentation cycle and \(x_f\) and \(y_f\) are cell densities at the end of each of fermentation cycle. For each competition experiment, the relative fitness was plotted as average of 20 replicates (replicate experiments with 10 fermentation cycles each).

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