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- 1 Title: A plasmid-encoded putative glycosyltransferase is involved in hop tolerance and
- 2 beer spoilage in Lactobacillus brevis.
- 3 Running title: A glycosyltransferase involved in beer spoilage in Lb. brevis
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- 14 **Keywords:** Lactic acid bacteria, plasmid, resistance, HorA, cell wall polysaccharide, phage.

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15 Abstract

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Lactobacillus brevis beer-spoiling strains harbor plasmids that contain genes such as horA, horC and hitA, which are known to confer hop tolerance. The Lb. brevis beer-spoiling strain UCCLBBS124, which possesses four plasmids, was treated with novobiocin resulting in the isolation of UCCLBBS124 derivatives exhibiting hop-sensitivity and an inability to grow in beer. One selected derivative was shown to have lost a single plasmid, designated here as UCCLLBS124_D, which harbors the UCCLBBS124_pD0015 gene, predicted to encode a glycosyltransferase. Hop tolerance and growth in beer was restored UCCLBBS124 pD0015 was introduced in one of these hop-sensitive derivatives on a plasmid. We hypothesize that this gene modifies the surface composition of the polysaccharide cell wall conferring protection against hop compounds. Furthermore, introduction of this gene in trans in Lb. brevis UCCLB521, a strain that cannot grow in and spoil beer, was shown to furnish the resulting strain with the ability to grow in beer while its expression also conferred phage-resistance. This study underscores how the acquisition of certain mobile genetic elements plays a role in hop tolerance and beer spoilage for strains of this bacterial species.

Importance

Lactobacillus brevis is a member of the lactic acid bacteria and is often reported as the causative agent of food or beverage spoilage, in particular that of beer. Bacterial spoilage of beer may result in product withdrawal or recall with concomitant economic losses for the brewing industry. A very limited number of genes involved in beer spoilage have been identified and primarily include those involved in hop resistance such as horA, hitA, and horC. However, since none of these genes are universal, it is clear that there are likely (many) other molecular players involved in beer spoilage. Here, we report on the importance of a plasmid-encoded glycosyltransferase associated with beer spoilage by Lb. brevis that is

- 40 involved in hop tolerance. The study highlights the complexity of the genetic requirements to
- 41 facilitate beer spoilage and the role of multiple key players in this process.

Introduction

Lactobacillus brevis is a major threat for commercial and amateur brewers as strains
of this species are the predominant bacterial contaminants associated with beer spoilage (1).
Such Lb. brevis strains can grow in beer despite the presence of ethanol, low pH and the
depletion of oxygen and nutrients (2). Moreover, hop compounds added to beer for bitter
flavor development during the fermentation process also exert antibacterial activity through
the presence of iso-α-acids (1, 2). Lb. brevis beer-spoiling (BS) strains appear to have
acquired chromosomally- or plasmid-derived genetic content to survive and grow in beer (2).
Lb. brevis resistance to ethanol (up to 10 %) and pH lower than the optimal growth
conditions (pH 4-6) seems to be associated with chromosomal genes, possibly due to the
general stressors they represent (3, 4). However, Lb. brevis BS strains are also known to
harbor plasmids that are associated with their beer-spoilage phenotype and more specifically
with hop tolerance (5-8). Plasmid-derived genes that underpin hop-resistance in Lb. brevis
include horA, horC, hitA and orf5 _{ABBC45} (1, 2). The genes horA and horC encode multidrug
transporter proteins driven by ATP and proton motive force (PMF), respectively, and were
identified as being involved in iso - α -acid extrusion from the bacterial cell (5, 7). The generation
hitA encodes a transmembrane protein involved in the transport of divalent cations such as
Mn ²⁺ in exchange of protons released from hop bitter acids (8). The <i>orf5</i> _{ABBC45} gene was
identified in Lb. brevis BS strain ABBC45 which was unable to grow in beer after it had lost
a plasmid carrying this gene. The orf5 _{ABBC45} gene encodes a predicted transmembrane protein
resembling a PMF-dependent multidrug transporter, which is presumed to be responsible for
iso-α-acid export (9).
However, these genes are not always indicative of BS ability as the presence of such

genes can be found among Lb. brevis strains that are unable to grow and consequently spoil beer (designated here as NBS strains) (10). Indeed, horA is present in the Lb. brevis NBS

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strain UCCLB556 (10). Moreover, genes identified as conferring hop-resistance are not always simultaneously present in BS strains, e.g. the BS strain UCCLBBS124 carries plasmids harboring horA and horC, however it does not possess hitA (10). Analysis of BS strain Lb. brevis BSO 464 has highlighted the importance of plasmids and genes on mobile genetic elements for bacterial growth in beer and beer spoilage ability (6). Recently, a gene predicted to encode a glycosyltransferase was identified among BS strains responsible for excess β-glucan formation (11). This gene is also present on the genome of Lb. brevis BS strain UCCLBBS124, while it is absent in that of BS strain UCCLBBS449 (10). This indicates that beer spoilage is not uniquely governed by the presence of a few genes, but rather a combination of genes acting in concert to confer beer resistance to the strain. It also suggests that other plasmid-encoded genes involved in beer spoilage are yet to be discovered. In the present study we generated plasmid-cured derivatives of Lb. brevis BS strain

UCCLBBS124 using novobiocin. This approach has been successfully employed previously to cure plasmids from lactic acid bacteria (LAB) isolates (6, 12). Plasmid-cured derivatives were assessed for their ability to grow in the presence of hop and in beer. A derivative that showed inability to grow in beer was selected and analyzed to ascertain which plasmids were responsible for this phenotype. Bioinformatic analysis of the genetic content of such plasmids revealed candidate genes required for growth in beer. These genes were used in transformation experiments to revert the NBS phenotype.

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Results and Discussion

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Derivatives with impaired growth in beer reveal loss of plasmid UCCLBBS124_D

The beer-spoiling Lb. brevis strain UCCLBBS124 (abbreviated here as UCC124) possesses four plasmids carrying genes of interest for bacterial beer spoilage (Table 3). Following exposure to novobiocin, surviving Lb. brevis UCC124 cells were plated and fifty isolated colonies (10) were randomly selected for further analysis. Thirty four of these fifty colonies displayed impaired growth in beer. PCR-based identification of the hop-resistance gene horA revealed the loss of this gene, located on plasmid UCCLBBS124 D (abbreviated here as UCC124 D) in 33 out of the 34 isolates. One derivative, designated here as MB569, was selected for genome sequencing, after which its sequence was compared to that of the WT, confirming that plasmid UCC124 D had been lost from strain MB569.

Tolerance of MB569 to iso-α-acids, ethanol and low pH

The inability of strain MB569 to grow in beer highlights the apparent importance of plasmid UCC124_D in conferring a beer spoilage phenotype on strain UCC124 (Figure 1). Beer is a harsh environment incorporating a number of stresses such as low pH, lack of nutrients, and the presence of ethanol and hop compounds. In order to understand which of these stresses imposed a negative impact on growth of MB569, the WT strain and MB569 were grown in MRS broth and mimicking conditions encountered in beer, e.g. pH4, 5.4 % ethanol, and 30 ppm iso-α-acids. Strain MB569 was shown to be capable of growth in MRS broth at neutral pH and at pH 4, while it can also grow in the presence of ethanol comparable to the WT strain (Figure 1). However, MB569 is incapable of growth in the presence of iso-α-acids unlike the WT strain UCC124 (Figure 1). This indicates that plasmid-cured derivative MB569 has lost the ability to spoil beer due to its sensitivity to the antimicrobial compounds present in hops. Therefore, based on this phenotype and the finding that MB569 lacks

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plasmid UCC124_D (when compared to its parental strain), it indicates that this plasmid is linked to hop tolerance and thus contributing to the ability of strain UCC124 to cause beer spoilage. Identification and functional annotation of genes present on plasmid UCC124_D Plasmid UCC124_D is 21 kb in size and is predicted to encompass 16 genes. Interestingly, a 7 kb region of this plasmid, contains six genes that are uniquely present among the plasmids of Lb. brevis BS strains (Table 4) (10). In order to assess the possible role of these genes in beer spoilage, the BS plasmid-specific genes UCCLBBS124_pD0014 (abbreviated here as UCC124_D14), encoding a predicted cytosine deaminase, UCCLBBS124_pD0015 (renamed gtf_{DI5}), encoding a predicted glycosyltransferase, UCCLBBS124 pD0016 (designated here as UCC124_horA), which encodes HorA (Table 4), were individually cloned into plasmid pNZ44 prior their transformation into NZ9000. The resulting plasmids were then introduced into strain MB569 to determine the ability of the obtained recombinant strains to grow in beer (where MB569 itself is unable to do so). Genes with locus tags UCCLBBS124_pD0017 (abbreviated as UCC124_D17), UCCLBBS124_pD0018 (abbreviated as UCC124_D18) and UCCLBBS124_pD0019 (abbreviated as UCC124_D19) and encoding acyl-sn-glycerol-3phosphate acyltransferases and a glycosyltransferase (Table 4) were cloned together as a cluster (as present in plasmid UCC124_D) in pNZ44 prior their introduction into NZ9000 and, subsequently, MB569. Introduction of the genes UCC124 D14, UCC124 horA, UCC124 D17, UCC124 D18 and UCC124_D19 in MB569 did not enable any obvious improvement of growth in the presence

of iso-α-acid (30 ppm) or beer (when compared to strain MB569) (data not shown).

Interestingly, expression of gtf_{D15} in MB569 was shown to confer a positive effect on its

ability to grow in MRS broth containing 30 ppm iso- α -acids, with a significant (P value <

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0.05) growth increase after 72 h compared to the non-complemented strain or MB569 carrying the control plasmid pNZ44 (Figure 2A). When Lb. brevis MB569 pNZ44:gtf_{D15} was cultivated in beer, it also exhibited an ability to grow in beer that was significantly better than that of MB569 itself (P value < 0.05) (Figure 2B). Provision of gtf_{D15} in trans in MB569 did not restore its growth in beer to the same level as the WT strain (i.e. the strain from which MB569 was derived), but nonetheless allowed survival and growth in beer for this recombinant strain across 96 h. MB569 and MB569 pNZ44 are still able to survive in the presence of iso-α-acids or beer after culture for 72 h (Figure 2A and 2B) which might be due to the presence of plasmid UCCLBBS124_C carrying the gene hor C (Table 3). The gtf_{D15} gene is predicted to encode a glycosyltransferase based on BLAST analysis and a HHPred analysis (13) predicted the protein to belong to the glycosyl transferase family 8 associated with cell wall glycosylation (99.9 % probability and E-value < 10⁻²⁸). Further sequence scrutiny suggests that the Gtf_{D15} protein is a membrane-associated protein (TMHMM Server 2.0 (14)) with a predicted signal peptide in its N-terminus that may act as a membrane anchor for the protein (http://phobius.sbc.su.se/ (15)). These predictions suggest that GtfD15 is a cell envelope-associated protein that confers protection against certain environmental stressors such as hop compounds.

Introduction of gtf_{D15} in NBS Lb. brevis strains allows growth in beer

The introduction of gtf_{D15} in MB569 was shown to significantly improve growth of the strain in MRS broth containing hop compounds (30 ppm iso- α -acids) and in beer indicating the importance of this gene for beer spoilage by Lb. brevis strain UCC124. In order to assess the potential growth-promoting effect of this gene for an NBS strain when inoculated in beer, gtf_{D15} when cloned into pNZ44 (pNZ44: gtf_{D15}) was introduced into the NBS Lb. brevis strain UCCLB521 (renamed here as UCC521) (Table 1). Remarkably, the presence of pNZ44:gtf_{DI5} in the NBS strain Lb. brevis UCC521 permitted the strain to grow significantly better (P

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value < 0.05) in MRS broth containing 30 ppm iso- α -acids and in beer compared to the strain carrying an empty plasmid which is incapable of survival or growth in these environments (Figure 2C and D). These observations reinforce our results above and highlight the significance of the gtf_{D15} gene in hop tolerance and beer spoilage. An alternative, though in our opinion less likely explanation is that strains for which we obtained no or reduced CFUs had entered a viable, but non-culturable (so-called VBNC) state as has been previously observed for beer-passaged Lb. brevis strains (16). Introduction of pNZ44:gtf_{D15} into ATCC 367, another NBS strain, did not allow improved survival or growth in the presence of hop compounds or in beer (data not shown). This suggests that a strain-specific mechanism and involvement of other genes that are absent in ATCC 367 are responsible for increased hop tolerance. Among beer spoilage-related genes, UCC521 possesses the orf5_{ABBC45} gene previously identified as involved in hop tolerance (9), unlike ATCC 367 which does not harbor known genes involved in beer spoilage. Moreover, UCC521, although a non-beer spoiler strain, was isolated from the brewery environment, unlike ATCC 367 which was isolated from silage (10). UCC521 may have acquired genes (such as orf5_{ABBC45}) or plasmids (UCC521 harbors five plasmids) throughout its presence in the brewery environment, which confer hop tolerance when combined with gtf_{D15} . This scenario has previously been observed, indicating that Lb. brevis strains can only survive and grow in beer when multiple beer spoilage-related genes are present in a particular combination (17). Selection pressures of the beer environment determine the genetic content of beer-spoiling strains. The identification of diagnostic marker genes (DMGs) are important in distinguishing BS from NBS strains, as well as predicting the ability of a given strain to grow in beer (17, 18). In the study from Bergsveinson and Ziola, proposed DMGs were not related to hop tolerance and no genes encoding glycosyltransferases were identified as DMGs. However, a glycosyltransferase-encoding gene located on the plasmid of the BS Lb.

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unique when compared to a NBS Lb. brevis strain KB290 and a BS Pediococcus damnosus strain Pc344^T (17). From these observations and knowing that gtf_{DL5} is highly prevalent in BS Lb. brevis strains (Table 4), we propose to include this gene as a DMG to assess the beer spoiling potential of Lb. brevis strains. Effect on phage sensitivity As demonstrated above gtf_{D15} was observed to play a role in hop and beer tolerance and is

brevis strain BSO 464 which showed more than 99 % nt similarity to gtf_{DI5} was described as

predicted to encode a glycosyltransferase. Since the protein is predicted to be involved in biosynthesis or modification of a cell surface-associated saccharidic polymer, the possible role of this protein in bacteriophage infection was investigated. Lb. brevis strain UCC521 is sensitive to Lb. brevis phages 3-521 and 521B (19). Plague assays employing these phages and Lb. brevis UCC521 harboring the empty vector pNZ44, or strain UCC521 containing pNZ44:gtf_{DI5} displayed similar EOP (Efficiency Of Plaquing) values with no significant difference to the WT (Table 5). However, notable differences in plaque morphology were observed, where plaques were faint and hard to distinguish on the bacterial lawn of UCC521 pNZ44:gtf_{DI5}. Moreover, overnight incubation of the different strains with the two phages led to complete lysis-in-broth of UCC521 and UCC521 containing pNZ44 with an approximately 1000-fold increase of phage titre after overnight propagation (Table 5). In contrast, UCC521 pNZ44:gtf_{D15} did not show visible lysis and was able to grow after overnight incubation with just a ten-fold increase in phage numbers after overnight propagation (Table 5). These results reinforce the role of the protein Gtf_{D15} in bacterial protection against diverse environmental hazards such as hop compounds or bacteriophages.

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Conclusions

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In this study, we identified a novel genetic component required for beer spoilage and more specifically for hop tolerance. This gene is located on plasmid UCC124_D of Lb. brevis BS strain UCC124, validating the importance of plasmids to confer a beer spoilage phenotype. Moreover, this gene had been highlighted previously as common among BS strains (10). Genes required for hop tolerance have all been identified on plasmids (5, 7, 8), reinforcing the importance of such mobile genetic elements in adaptation to the specific hurdles imposed by the beer environment. A derivative of UCC124, MB569 showed impaired growth in beer after the loss of plasmid UCC124 D and despite the presence of plasmids UCC124 B and UCC124_C which carry several genes of interest in beer spoilage. Introduction of gtf_{D15} in strain MB569 restored the hop tolerance phenotype of the strain which ultimately allowed it to grow in beer. Similar results were observed when the gene was introduced into a NBS strain confirming the notion that gtf_{DI5} is required for the development of hop tolerance and beer spoilage. Furthermore, this gene impacts on phage sensitivity of its host. This gene seems a unique trait shared among BS strains of Lb. brevis and we propose gtf_{D15} as a DMG for the detection of potential bacterial contamination of beer. The gene is predicted to encode a glycosyltransferase and analysis of its topology suggests that it is a membrane-anchored protein involved in the biosynthesis or modification of a cell surface-associated saccharidic polymer. BS strains of Lb. brevis have been shown to increase higher molecular weight lipoteichoic acids (LTA) in their cell wall, in the presence of hop bitter acids, thus believed to confer resistance to the bacteria by enhancing the barrier functions of the cell wall and preventing intrusion of hop compounds (20, 21). Moreover, lipoteichoic acids have been described as phage receptors among lactobacilli phages as seen for Lactobacillus delbrueckii phages LL-H and JCL1032 (22) but also for Lactobacillus plantarum ATCC8014-B2 (23). Therefore, we speculate that this glycosyltransferase is involved in replacing alanine residues

with sugar residues on teichoic acids thereby changing their charge and preventing iso-αacids to penetrate the membrane as well as affecting phage adsorption and/or DNA injection. This predicted glycosyltransferase shows only limited similarity (36 % amino acid similarity in 20 % query cover) with the glycosyltransferase identified in a previous study as responsible for β -glucan formation (11), and is thus believed to play a different role in beer spoilage. Future studies will focus on defining the mechanism that underpins hop tolerance and on determining how the genes identified to date (5, 7, 8) are linked to each other. Moreover, located on the same plasmid as gtf_{DI5} are genes predicted to encode a glycosyltransferase and acyltransferases (Table 2) suggesting a common action on teichoic acids with the acyltransferases involved in the acylation of alanine residues or the lipid moiety of the lipoteichoic acids (24). Follow-up work may therefore focus on determining the precise function of the glycosyltransferase (and other associated genes) in the modification of the cell wall and/or cell surface. Another question to be addressed is if and how hop tolerance is enhanced when these genes are present in a certain combination, and how such tolerance is influenced by their expression level.

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Materials and Methods

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253 Bacterial strains and cultivation media

254 Bacterial strains used in this study are listed in Table 1. Lb. brevis strains were grown in

255 MRS broth (Oxoid Ltd., UK) at 30 °C while Lactococcus lactis NZ9000 was grown in M17

256 broth (Oxoid Ltd., UK) supplemented with 0.5 % glucose. 5 µg/mL chloramphenicol (Cm5)

257 was added to the medium when indicated.

258 Plasmid curing and plasmid content analysis

> The overall experimental approach is presented in Figure S1. Plasmid curing of the BS strain UCC124 was achieved using novobiocin treatment (25). A 1 % inoculum of a WT strain overnight culture was used to inoculate 10 mL MRS broth containing 0.25 µg/mL novobiocin. Cultures were incubated at 26 °C for 72 h. After incubation, cells were diluted and plated on MRS agar. After 3 days of incubation at 26 °C, isolated colonies were randomly selected and derivatives with impaired growth in beer (no growth observed after 72 h) were checked for the presence or loss of hop-resistance genes horA, horC and $ort5_{ABBC45}$ (Table 2). A derivative showing loss of hop-resistance gene was selected and sequenced using Illumina sequencing technology. Paired-end sequence reads were generated using an Illumina HiSeq2500 system (read length 2 x 250 bp). FASTQ sequence files were generated using the Illumina Casava pipeline version 1.8.3. After Illumina sequencing the obtained sequences were mapped back against the WT reference sequence to detect mutations by single nucleotide polymorphism (SNP) or plasmid content loss. SNP analysis was performed by aligning Illumina raw reads against a reference sequence using Bowtie2 V 2.3.5 (26). The reads were then sorted using Samtools (27) and VarScan v2.3.9 was applied for the detection of variants (28). A minimum allelic variation frequency cut-off of 0.25 was applied.

Construction of plasmid vectors

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Genes of interest were amplified by PCR (Table 2) and cloned into the expression vector pNZ44 (29). PCR products and pNZ44 plasmid DNA were digested with the appropriate enzymes (Roche, USA) at 37 °C for at least 4 h, following the manufacturer's instructions (Table 2). A ratio of (3:1) was applied for the ligation of the PCR product with pNZ44 using T4 DNA ligase (Promega, USA). The mixture was incubated at room temperature for at least 4 hours prior to electrotransformation into *L. lactis* NZ9000 competent cells.

Preparation of competent cells and electrotransformation

Competent cells of L. lactis NZ9000 were prepared as previously described (30). Competent cells of Lb. brevis UCC124 were prepared using an adapted version of a previously described protocol (31): An overnight culture was transferred (1 % inoculum) to 10 mL MRS broth containing 1 % glycine and incubated overnight at 30 °C. 5 mL of the overnight culture was transferred to fresh MRS broth containing 1 % glycine (50 mL final volume) and cells were grown to an OD_{600nm} of 0.6. Cells were harvested by centrifugation at 4,000 \times g for 15 min at 4 °C and washed in ice-cold wash buffer (0.5 M sucrose, 10 % glycerol). The wash step was repeated twice and the cells were finally resuspended in 200 µL wash buffer prior to storage at -80 °C and/or electroporation (see below). All constructs were generated using L. lactis NZ9000 as the cloning host, verified by sequencing after PCR amplification using the primers pnz44F and pnz44R (Table 2) prior to their transfer into Lb. brevis strains. Electrotransformation was performed using freshly prepared competent cells as described above, where 45 µL of cells and 5 µL of plasmid construct were mixed into a pre-chilled 2 mm electroporation cuvette (Cell Projects, Kent, England) and subjected to electroporation at 1.5 kV (Lb. brevis) or 2.0 kV (L. lactis), 200 Ω, 25 μF. Following electroporation, 950 μL recovery broth was added (MRS broth supplemented with 0.5 M sucrose and 0.1 M MgCl₂ (Lb. brevis) or GM17 broth supplemented with 20 mM MgCl₂ and 2 mM CaCl₂ (L. lactis)). Cells were recovered at 30 °C for 3 h (Lb. brevis) or 2 h (L. lactis) prior to spread plating on

MRS (Lb. brevis) or GM17 (L. lactis) agar supplemented with Cm5. Presumed transformants were purified on MRS agar + Cm5 and colonies were checked by sequencing after PCR amplification using the primers pnz44F and pnz44R (Table 2) and applied to growth assays as described below.

Growth assays

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Growth profiles of the wild-type strain and its derivative were obtained by transferring an overnight culture (1 % inoculum) to MRS broth, MRS broth supplemented with 30 ppm isoα-acids or beer (fresh Heineken lager 5 % ethanol, pH 4, 23 ppm iso-α-acids). Cultures were incubated at 30 °C for 72 hours. One mL of culture was retrieved after 24, 48, 72 and 96 hours, diluted in Ringer's solution and plated on MRS agar plates. Plates were incubated at 30 °C anaerobically for 48 hours prior to colony counting. The number of viable bacteria of each strain was assessed after CFU/mL calculation. Non-inoculated controls were used in all the experiments as blank measurements. These measurements were then subtracted from each experimental condition to produce the values represented on growth curves. Statistical differences were calculated using unpaired t test method (32).

Phage activity against Lb. brevis strains and transformants

To assess phage sensitivity of Lb. brevis strains, transformants carrying genes of interest were compared to the wild-type (WT) strain using plaque assays, as previously described (33). A 10 μL volume of the appropriate phage dilution and 200 μL of Lb. brevis culture were added to 4 mL of soft agar supplemented with 10 mM CaCl₂, mixed and poured onto an MRS agar plate supplemented with 10 mM CaCl₂ and 0.5% glycine. Plates were incubated at 30 °C overnight and the resulting plaques were enumerated. Phage titre was determined as plaqueforming units per mL (PFU/mL). The ability of phages to propagate and multiply within the host cell was also tested. Lb. brevis strains were grown to early exponential phase (OD_{600nm} ~

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325 0.25), at which point phages were added to the culture (T0) at a MOI (multiplicity of 326 infection) of 1, along with 10 mM CaCl₂. The mix was further incubated at 30 °C overnight 327 (T1). The number of phages present in the medium (i.e. following removal of bacterial cells 328 by centrifugation) at T1 was then determined by plaque assay. Phage propagation efficiency 329 on a given host was then determined by dividing the phage titre (PFU/mL) at T1 by the phage 330 titre (PFU/mL) at T0. 331 GenBank accession numbers Lb. brevis UCCLBBS124: CP031169, Lb. brevis UCCLBBS124_A: CP031170, Lb. brevis 332 333 UCCLBBS124 B: CP031171, Lb. brevis UCCLBBS124 C: CP031172 and Lb. brevis 334 UCCLBBS124_D: CP031173. 335 Author Contributions: MF performed experiments and genomic analysis. DS, JM, TS and 336 VB provided materials and strains. MF, JM, TS and DS were involved in project design and 337 wrote the manuscript. All authors read and approved the final manuscript. 338 **Conflicts of Interest:** The authors declare that VB and TS are employees of Heineken. 339 Funding: Marine Feyereisen is the recipient of an Irish Research Council Enterprise 340 Partnership Scheme postgraduate scholarship (Ref. No. EPSPG/2015/7). Douwe van Sinderen is supported by a Principal Investigator award (Ref. No. 450 13/IA/1953) through 342 Science Foundation Ireland (SFI). Jennifer Mahony is in receipt of a Starting Investigator 343 Research Grant (SIRG) (Ref. No. 15/SIRG/3430) funded by Science Foundation Ireland 344 (SFI).

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Table 1. Bacterial strains and plasmids used in this study.

Strain / Plasmid	Description	References
Lb. brevis strains		
UCCLBBS124 (UCC124)	Beer-spoiling strain isolated from spoiled beer keg (Singapore)	(10)
UCCLBBS449	Beer-spoiling strain isolated from unpasteurized spoiled beer (The Netherlands)	(10)
MB569	Non-beer spoiling strain derivative of UCCLBBS124	This study
UCCLB521 (UCC521)	Non-beer spoiling strain isolated from brewery environment (The Netherlands)	(10)
MB569 pNZ44	MB569 carrying pNZ44	This study
MB569 pNZ44:gtf _{D15}	MB569 carrying pNZ44 with gtf_{DI5}	This study
UCCLB521 pNZ44:gtf _{D15}	UCCLB521 carrying pNZ44 with gtf _{D15}	This study
L. lactis strains		
NZ9000	Transformation host	(34)
Plasmids		
pNZ44	Transformation vector, chloramphenicol resistance gene	
pNZ44: <i>gtf</i> _{D15}	pNZ44 harboring gtf_{DI5}	This study

Table 2. PCR primers used in this study. Incorporated restriction sites are indicated in capital letters.

Primer name	Sequence (5' - 3')	Target	GenBank accession no.
horAF	cgcaactgaggctaacttct	horA gene in UCCLBBS124	CP031173
horAR	ggcttgctatgctaggata	horA gene in UCCLBBS124	CP031173
horCF	gtatgcctaagtgacgt	horC gene in UCCLBBS124	CP031172
horCR	cattetetgeetetatae	horC gene in UCCLBBS124	CP031172
orf5F	ctggattgaggtgaggg	orf5 gene in UCCLBBS124	CP031172
orf5R	gctgtaaagggtagtgattg	orf5 gene in UCCLBBS124	CP031172
pNZ44F	aacaattgtaacccatac	pNZ44 promoter	
pNZ44R	gaacgtttcaagccttgg	pNZ44 MCS	
pD14F	aaaaaaCTGCAGgtccgaacagcgttcggatt	Gene UCCLBBS124_pD0014 in UCCLBBS124_D	CP031173
pD14R	aaaaaaTCTAGAttaatcttcgaaatagtt	Gene UCCLBBS124_pD0014 in UCCLBBS124_D	CP031173
pD15F	aaaaaaCCATGGgcggtttggatattttatact	Gene UCCLBBS124_pD0015 in UCCLBBS124_D	CP031173
pD15R	aaaaaaTCTAGAtcactcagttttcaattccc	Gene UCCLBBS124_pD0015 in UCCLBBS124_D	CP031173
pD16F	aaaaaaCTGCAGaggcttgctatgctagg	Gene UCCLBBS124_pD0016 in UCCLBBS124_D	CP031173
pD16R	aaaaaaTCTAGAtcacccgttgctcgt	Gene UCCLBBS124_pD0016 in UCCLBBS124_D	CP031173
pD17-19F	aaaaaaCCATGGggggtagaatggttctgtt	Gene UCCLBBS124_pD0017-19 in UCCLBBS124_D	CP031173
pD17-19R	aaaaaaTCTAGAttattgataatgaccagcaa	Gene UCCLBBS124_pD0017-19 in UCCLBBS124_D	CP031173

Table 3. Lb. brevis UCC124 plasmids and genes of interest for beer spoilage.

UCC124 plasmids	Size (bp)	ORFs no.	Accession no.	Gene(s) of interest	References
UCCLBBS124_A (UCC124_A)	49,560	42	CP031170		
UCCLBBS124_B (UCC124_B)	23,078	20	CP031171	gtf family 2	(11)
UCCLBBS124_C (UCC124_C)	22,370	27	CP031172	horB, horC, orf5	(7, 9)
UCCLBBS124_D (UCC124_D)	20,971	16	CP031173	horA	(5)
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Table 4. Presence and absence of genes of UCCLBBS124_D among *Lb. brevis* BS strains.

Gene	Gene Predicted function				Lb. brevis BS strains			
		UCCLBBS124	UCCLBBS449	UCCLB95	TMW1.2108	TMW1.2111	TMW1.2112	TMW1.2113
UCCLBBS124_pD0014 = UCC124_D14	Cytosine deaminase	+	+	-	+	+	-	+
$UCCLBBS124_pD0015 = gtf_{D15}$	Glycosyltransferase family 8	+	+	-	+	+	+	+
UCCLBBS124_pD0016 = UCC124_horA	HorA	+	+	-	+	+	-	+
UCCLBBS124_pD0017 = UCC124_D17	Acyl-sn-glycerol-3-phosphate acyltransferase	+	+	-	+	+	+	+
$UCCLBBS124_pD0018 = UCC124_D18$	Glycosyltransferase family 8	+	+	-	+	+	+	+
$UCCLBBS124_pD0019 = UCC124_D19$	Acyl-sn-glycerol-3-phosphate acyltransferase	+	+	-	+	+	+	+
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476 Table 5. Effect of phages 3-521 and 521B on Lb. brevis strain UCC521 and derivatives.

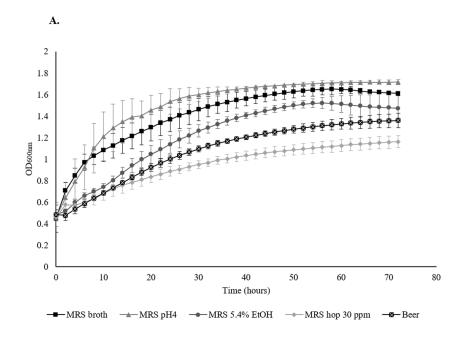
			Lb. brevis strains	
		UCC521	UCC521 pNZ44	UCC521 pNZ44:gtf _{D15}
	EOP (Efficiency Of Plaquing)	1.00	0.58 ± 0.29	0.64 ± 0.21
Phage 521B	Plaque morphology	Small clear plaques	Small clear plaques	Faint plaques
	Phage titre after O/N propagation* (PFU/mL)	2.90E+09	2.30E+09	3.00E+07
	EOP	1.00	1.52 ± 0.20	1.19 ± 0.19
Phage 3-521	Plaque morphology	Small clear plaques	Small clear plaques	Faint plaques
	Phage titre after O/N propagation* (PFU/mL)	4.30E+09	1.80E+09	4.80E+07

⁴⁷⁷ *Overnight propagation (O/N) was realized with a starting phage titre of 10⁶ PFU/mL (results

⁴⁷⁸ are average of triplicate assays).

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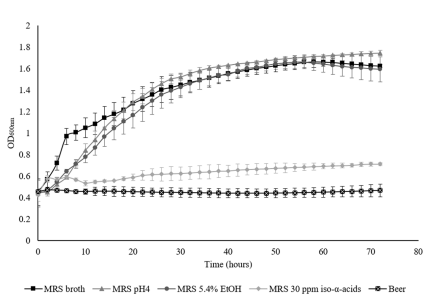


Figure 1. Growth of the WT beer-spoiling strain Lb. brevis UCC124 (A) and its plasmidcured derivative MB569 (B) in beer, MRS broth, MRS broth at pH4 and MRS broth supplemented with 5.4 % ethanol or 30 ppm *iso*-α-acids.

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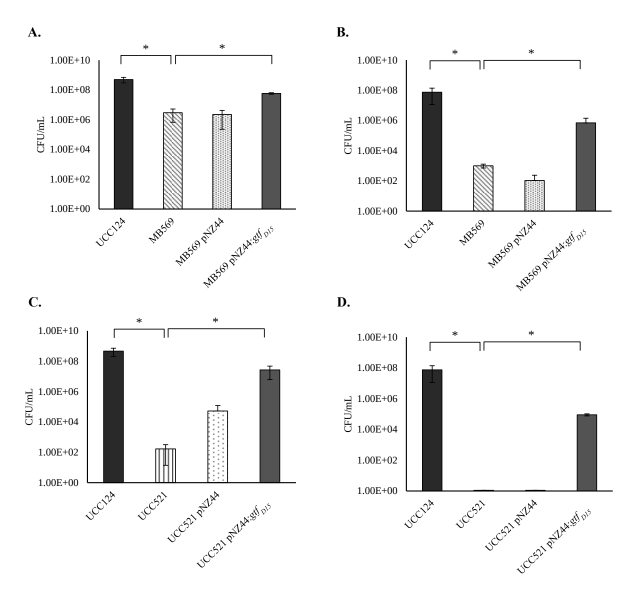


Figure 2. Number of viable bacteria (CFU/mL) of the WT BS strain Lb. brevis UCC124, the derivative MB569 +/- the empty plasmid pNZ44 and MB569 carrying the gene gtf_{DI5} after growth in (A) MRS broth containing 30 ppm iso-α-acids for 72 h and (B) beer for 96 h (P<0.05). CFU/mL of the WT BS strain Lb. brevis UCC124, the NBS UCC521 +/- the empty plasmid pNZ44 and the NBS UCC521 carrying the gene gtf_{D15} after growth in (C) MRS broth containing 30 ppm iso- α -acids and (**B**) beer for 96 h (P<0.05).