RNA regulation in Lactococcus lactis
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Document Version
Publisher's PDF, also known as Version of record

Publication date:
2018

Link to publication in University of Groningen/UMCG research database

Citation for published version (APA):
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Transcriptome landscape of *Lactococcus lactis* reveals many novel RNAs including a small regulatory RNA involved in carbon uptake and metabolism

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*Published in RNA Biology, 2016* 13:353-366
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Transcriptome landscape of Lactococcus lactis reveals many novel RNAs including a small regulatory RNA involved in carbon uptake and metabolism.

ABSTRACT

RNA sequencing has revolutionized genome-wide transcriptome analyses, and the identification of non-coding regulatory RNAs in bacteria has thus increased concurrently. Here we reveal the transcriptome map of the lactic acid bacterial paradigm Lactococcus lactis MG1363 by employing differential RNA sequencing (dRNA-seq) and a combination of manual and automated transcriptome mining. This resulted in a high-resolution genome annotation of L. lactis and the identification of 60 cis-encoded antisense RNAs (asRNAs), 186 trans-encoded putative regulatory RNAs (sRNAs) and 134 novel small ORFs. Based on the putative targets of asRNAs, a novel classification is proposed. Several transcription factor DNA binding motifs were identified in the promoter sequences of (a)sRNAs, providing insight in the interplay between lactococcal regulatory RNAs and transcription factors. The presence and lengths of 14 putative sRNAs were experimentally confirmed by differential Northern hybridization, including the abundant RNA 6S that is differentially expressed depending on the available carbon source. For another sRNA, LLMGnc_147, functional analysis revealed that it is involved in carbon uptake and metabolism. L. lactis contains 13% leaderless mRNAs (lmRNAs) that, from an analysis of overrepresentation in GO classes, seem predominantly involved in nucleotide metabolism and DNA/RNA binding. Moreover, an A-rich sequence motif immediately following the start codon was uncovered, which could provide novel insight in the translation of lmRNAs. Altogether, this first experimental genome-wide assessment of the transcriptome landscape of L. lactis and subsequent sRNA studies provide an extensive basis for the investigation of regulatory RNAs in L. lactis and related lactococcal species.
INTRODUCTION

Genome-wide transcriptome analyses using RNA sequencing (RNA-seq) has allowed adding to previously annotated genomes numerous novel elements, such as non-coding regulatory RNAs (sRNAs), antisense RNAs (asRNAs), small open reading frames (sORFs) and riboswitches. In addition, RNA-seq provides excellent opportunities to correct errors in annotated ORFs, to determine operon structures and to identify alternative internal transcription start sites (TSS) within coding genes.

An important next step in these studies is the validation of these novel RNAs and to unravel their functions in the cell. Functional studies of regulatory RNAs from a variety of bacterial genomes now reveal an ever-increasing number of new regulatory mechanisms. A lot of this research has been devoted to the sRNAs, which have been shown to post-transcriptionally control numerous cellular processes. They act mostly by base pairing with their target mRNAs, thereby influencing transcription termination, mRNA stability and/or mRNA translation (1-3). In addition, some sRNAs, such as the 6S sRNA and members from the CsrB family, have been reported to bind to and thereby influence the functionality of RNA polymerase and CsrA, respectively (4). Regulatory RNAs can function as signaling regulators responding to a changing environment and preparing the cell for altered conditions, as seen e.g. in pathogenic bacteria (5, 6). Most regulatory RNAs are not translated into proteins and are therefore called non-coding regulatory RNAs although there are a number of exceptions of so-called dual-function RNA regulators; well-studied sRNAs such as RNAIII (7, 8) and SgrS (9) have been reported to act as RNA regulators but also code for (small) proteins. The dual-function sRNAs can provide valuable insights into the evolutionary development of these RNAs by studying the physiological roles of the encoded peptide and the non-coding regulatory part (10).

Different types of non-coding regulatory RNAs can be distinguished. For example, regulatory RNAs that derive from intergenic regions (IGRs) are generally named sRNAs. They are trans-encoded and affect one or more mRNA targets via imperfect base pairing. The RNA chaperone protein Hfq is often required to enable the interaction between the sRNA and its target mRNA (11-13). Hfq seems to be mainly present in GC-rich bacteria (14), and an Hfq homologue is absent in lactococcal genomes. Hfq has been used to purify and identify novel bacterial regulatory RNAs through Hfq-RNA immunoprecipitation and subsequent RNA-seq (15, 16). By studying Hfq-bound transcripts, it has recently been shown that the 3’-untranslated regions (3’-UTRs) of mRNAs can harbor functional regulatory RNAs that act in trans. Such an sRNA can be derived by cleavage of the 3’-UTR of the original mRNA molecule. Alternatively, a separate promoter in the 3’-end of the gene can lead to an sRNA that overlaps the 3’-UTR of the mRNA (17).

The second class of regulatory RNAs comprises the cis-encoded RNAs or antisense RNAs (asRNAs). An asRNA derives from the non-coding strand of a gene and its sequence is, thus,
complementary to (part of) the gene’s mRNA. Although most regulatory RNAs are small and range in size from 50 to 350 nucleotides, antisense transcription can cover whole operons (18). The functions of many asRNAs still remain to be elucidated. The fraction of asRNAs in the total RNA pool of a bacterium is significant albeit variable between bacterial species (19). Even within species the total amount of asRNAs can greatly vary, which was very recently illustrated in an *E. coli* study that devoted special attention to asRNAs (20). Base pairing between an sRNA or asRNA and its partner mRNA usually involves the repression or activation of translation of the mRNA. The binding via 16S rRNA of the small 30S ribosomal subunit can be negatively affected by blocking of the ribosomal binding site (RBS) by the regulatory RNA (21). Activation can occur through the unfolding of a secondary structure in the mRNA via interaction with the regulatory RNA and the consequent liberation of the RBS (22). Moreover, base pairing of the two RNAs can lead to degradation of both by the endoribonuclease RNase E (23). Cleavage can occur near the RBS in the 5’ leader, in the coding region (24), or it can even take place downstream of the region where sRNA and mRNA interact (25). Translation-independent stabilization of mRNAs has also been reported in which the sRNA-mRNA hybrid interferes with RNase E-mediated degradation (26, 27). Another ribonuclease that is important in mRNA regulation by sRNAs and asRNAs is RNase III, an enzyme that cleaves double stranded structures such as (a)sRNA-mRNA hybrids. Another class of *cis*-encoded regulatory RNAs are sequences at the 5’ end of mRNAs that are able to change their conformation in response to an environmental cue. So-called thermometers react to changes in temperature (28), whereas a variety of riboswitches operate as intracellular sensors by binding to small metabolites or ions. Binding of the effector molecule influences the secondary structure of the riboswitch part of the mRNA, which affects the fate of transcription and/or determines whether the coding part of the mRNA is actually translated. Riboswitches can also influence mRNA stability (29, 30). Two SAM riboswitches involved in the regulation of methionine and cysteine biosynthesis in *L. monocytogenes* were reported to act in-trans (31). Another surprising form of RNA regulation was reported in *S. aureus*, in which the 5’-UTR of the *icaR* mRNA interacts with the 3’-UTR of the same mRNA. This may either occur in cis within one mRNA molecule or in trans, involving two copies of the *icaR* transcript (32).

RNA-seq and, to a lesser extent, tiling arrays have recently greatly increased the number of sRNAs in various microorganisms such as *E. coli* (33), *B. subtilis* (34), *H. pylori* (35) and *P. aeruginosa* (36). The techniques also allowed, by exact determination of transcription start sites, the description of novel sORFs (37), operon structures and have in certain cases led to re-annotation of known ORFs. *Lactococcus lactis* is an AT-rich, Gram-positive, mesophilic lactic acid bacterium with a relatively small genome size of 2.53 Mbp (38). It is widely applied in the dairy industry where its main function is to convert lactose into lactic acid and to provide texture, flavors and aromas. Previous studies using DNA microarray and proteomics technologies have
identified genes and proteins involved in various (environmental) stress responses in *L. lactis* (39, 40). The functioning in *L. lactis* of global regulators such as CcpA (41) and CodY (42) in carbon and nitrogen metabolism, as well as quite a number of other protein regulators has been described in considerable detail (43). Notwithstanding this, the presence and roles of regulatory RNAs *L. lactis* has not yet been reported, while it is becoming increasingly clear that these molecules play pivotal roles in gene regulation in many microorganisms, especially also in coping with stressful conditions. A better understanding of whether and how regulatory RNAs are involved in the regulation of stress responses and metabolic processes in *L. lactis* could lead to an improvement of the gene regulatory model of this organism (44) and may have practical (industrial) implications. Using differential RNA sequencing (dRNA-seq), we uncovered 375 novel RNAs including sRNAs, asRNAs, long 5’-UTRs, putative regulatory 3’-UTRs, novel (small) ORFs, internal promoters, transcription start sites and operon structures.

RESULTS AND DISCUSSION

**Determination of the primary transcriptome of *L. lactis***. In order to obtain deep insight in novel RNA elements in *Lactococcus lactis*, the organism was grown in GM17 and the cultures were harvested at six time-points during growth, three each in the exponential- and stationary phases, and mixed in equal OD equivalents prior to total RNA isolation and subsequent cDNA library preparation. Selective enrichment of primary transcripts was achieved by a Terminator 5’-phosphate-dependent exonuclease (TEX) treatment that specifically degrades processed 5’-monophosphate (5’P) RNA molecules (35). In addition to primary transcript enrichment, TEX treatment also results in enriched 5’-ends of mRNAs and ncRNAs in the RNA pool. In total, 10.5 million reads were generated, of which 7.2 million reads with a PHRED score > 28 were mapped onto the genome of *L. lactis* MG1363 (38).

Both *in silico* methods and visual inspection of the data were used to classify the *L. lactis* transcripts.

**Identification of *L. lactis* sRNAs from intergenic regions and 3’-UTRs**. The TEX-treated RNA was mapped on the genome of *L. lactis* MG1363 together with the *in silico* regulatory RNA prediction output from SIPHT (45) to aid in the mining for potential regulatory RNAs. The genome-wide map was then visually inspected for sRNAs, asRNAs, long 5’-UTRs and to review and correct open reading frame (ORF) boundaries. See Figure 1A for an overview of the results of the transcription start site (TSS) typing. The RNA-seq data and mining results have been integrated in a webpage using the JBrowse viewer, and can be assessed by http://jbrowse.molgenrug.nl/.

RNAs in IGRs were annotated as sRNAs (denoted: LLMGnc_001-186). Ten of these putatively trans-acting sRNAs overlap the 3’-UTR of mRNAs. They were identified on the basis of a
transcriptome landscape of Lactococcus lactis reveals many novel RNAs including a small regulatory RNA involved in carbon uptake and metabolism. A high number of reads within the 3’-UTR, suggesting that a promoter exists for these sRNAs, although the possibility that they derive from processing of the overlapping longer mRNA cannot be excluded. Three sRNAs (LLMGnc_012/013/014) are located within a region of only six genes and show exceptionally high sequence similarity, suggesting they might have a common function. As a means to verify that the 186 sRNAs are genuine and to assess their conservation, a blast search was performed on 10 related L. lactis genomes. These genomes cover five strains each of the L. lactis subspecies lactis and cremoris. Most of the identified sRNA sequences are conserved in the subsp. cremoris strains, while this is to a lesser extent so for the 5 strains of the subspecies lactis.

To examine the consensus of sRNA promoters, we evaluated the region from -100 to -1 upstream of the sRNA TSSs using MEME (46). We found no significant difference between the sRNA promoter consensus and the canonical L. lactis promoter. Subsequently, we screened the sRNA-promoter regions for the presence of known L. lactis transcription factor binding sequences (TFBSs). TFBSs for CcpA (carbon catabolite repression) (41), CodY (nitrogen metabolism) (42), ArgR (arginine metabolism) (47) or FlpAB (metal ion homeostasis and oxidative stress) (48) were identified upstream of 16 sRNA genes. These putative regulation sites provide a link between transcription factors and sRNAs (49) and underpin the versatility of gene regulation in this bacterium. To assess the relation between the predicted TFBSs in sRNA promoters and the function of the sRNA itself, two sRNA candidates (LLMGnc_147 and S6), which are predicted to be controlled by CcpA, were studied more in detail (see below).

To evaluate the presence in L. lactis of RNAs homologous to regulatory RNAs from other prokaryotes, we used the Bacterial Small Regulatory RNA Database (BSRD) (50). A total of 37 of such homologous regulatory RNAs could thus be identified in the genome of L. lactis MG1363. Although most of these correspond to regulatory elements located in 5’-UTRs, such as riboswitches, five RNAs from the BSRD matched to sRNAs identified in this study. These include the high-abundant housekeeping RNA 6S (51) (LLMGnc_004), the non-coding catalytic subunit of RNase P (52) (LLMGnc_059) and the tmRNA or SsrA RNA (53) (LLMGnc_074), which had not been annotated previously in L. lactis.
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**Figure 1. TSS mining, 5'-UTR distribution and promoter analysis.** (A) Different types of transcription start sites identified in the *L. lactis* MG1363 genome from mapped reads of the TEX-treated RNA-seq dataset (grey arrows: annotated ORFs, blue: Reads from the + strand, green: Reads from the - minus strand, red blocks: Positions of putative regulatory RNAs predicted by SIPHT. (B) Length distribution of 5'-UTRs. 5'-UTRs up to a length of 100 nt are plotted in stepwise increments of 10 nt in grey, those larger than 100 nt are shown with increments of 25 nt (separated by the dotted line). Color code is given in the inset. The RBS consensus sequence and the consensus sequence in the first 7 nt of the 111 leaderless mRNAs were determined by MEME. (C) Top: Analysis using MEME of motifs in the 50 nt upstream of all 1819 TSSs predicted by TSSer. Curved dotted line: periodic AT stretches. Bottom: Reconstruction of the *L. lactis* promoter consensus using MAST. In both: -35 and -10 sequences are indicated.

**L. lactis** antisense RNAs and functional classification. RNAs that overlap in an antisense fashion with transcripts (including their 3'- or 5'-UTRs) were annotated as antisense RNAs (asRNAs). In total, 60 of such asRNAs were identified in the RNA-seq dataset derived from the TEX-treated RNA sample. The asRNAs were classified as being located at 5’, internal or 3’- positions relative to the gene on the opposite strand. Although many single or low-abundant antisense reads were specified throughout the genome, we only took those reads into consideration when a TSS was present immediately downstream of the conserved promoter motifs -10 (TATAAT) and/or -35 (TTGACA), allowing two mismatches. In comparison with the reads from sRNAs located in IGRs, antisense transcripts were generally less abundant. This may be explained by assuming that the perfect match between asRNAs and their target mRNAs is more stable and therefore makes them better substrates for degradation by RNases. In an *E. coli* study, the abundance of functional asRNAs that form a duplex with other RNAs appears to be very low due to cleavage by RNase III (54) Further...
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studies on the stability of the *L. lactis* asRNAs are needed to draw reliable conclusions on this matter. That asRNAs appear to be (relatively) more abundant in other organisms (19), might be species-specific and/or may have a technical origin in the different automated or manual annotation approaches used in the various studies. On the basis of what will follow, we propose to distinguish three functional classes of asRNAs; in addition to the “regulatory” asRNAs, which have a role comparable to that of *trans*-encoded sRNAs, asRNAs can have a “protective” or “meta-regulatory” function.

We identified a relatively large number of novel asRNAs and sRNAs in regions in the genome of *L. lactis* that carry (remnants of) pro-phages. More than a third of all asRNAs are specified in these areas, while 5.5% of the genome of *L. lactis* MG1363 are bacteriophage-derived sequences (38). These asRNAs target the 5’- or 3’-UTR or the coding parts of phage transcripts. The asRNAs may be native to the phage genomes but could also have evolved after integration of the phage in *L. lactis* MG1363, via mutations in AT-rich regions leading to novel promoters driving asRNA synthesis. Gene silencing by antisense transcription might suppress any harmful phage induction by targeting essential transcripts necessary for the phage to enter the lytic phase. Interestingly, all six (defective) pro-phages contain asRNAs against their respective integrase genes. *L. lactis* MG1363 is known to lack active pro-phages, although two of the six phage genomes appear to be complete (55). In addition to the 19 asRNAs, 28 sRNAs were detected in the IGRs in the genomes of these (defective) pro-phages. Although the function of these sRNAs is still unknown, they could operate such that they create a lysogen that serves as an optimal host for silent phage propagation throughout all cells in the culture. These asRNAs could serve a protective role.

Ten asRNAs were detected in loci coding for transcriptional regulators. In these cases, antisense transcription could function as a rapid off-switch under those conditions where the regulators are no longer required, and this would represent asRNAs with a meta-regulatory function.

Finally, two pairs of mRNAs (*llmg_0538* and *llmg_0539* (fabI), both involved in fatty acid biosynthesis, and *llmg_0529* and *llmg_0530* (gapA)) overlap in an antisense fashion at their 5’-UTRs, with respectively 31 and 37 complementary nucleotides. Transcription of these mRNA pairs might be influenced due to variable promoter strengths (56). Also, the consequence for translation of the mRNA-pairs is unclear. Even though the overlapping sequences do not include the RBS regions, secondary structures might affect RBS-accessibility upon interaction of the two transcripts of a pair. In addition, transcript stabilization by the base pairing may occur, as well as degradation after processing by e.g. RNaseIII. With only these two examples, overlapping 5’-UTRs are a relatively rare phenomenon in the transcriptome of *L. lactis* MG1363. Other types of overlapping transcripts have been identified in *L. monocytogenes* (57), where both 3’-UTR and 5’-UTR overlapping transcripts occur. Overlapping transcripts of operons have also been observed in *S. aureus* (18).
Long 5’-UTRs and conserved riboswitches. The RNA-seq dataset from the TEX-treated sample was also used to evaluate the 5’-untranslated regions in a genome-wide manner in order to identify putative cis-encoded intracellular sensors such as riboswitches, which can affect the expression of downstream gene(s). To detect 5’-UTRs carrying potential regulatory elements, those containing ≥100 nucleotides were examined, resulting in the identification of 129 leader sequences (LLMG_R001-129). As mentioned above, most of the 36 regulatory RNA homologs in the BSRD database specified by the genome of L. lactis MG1363 are present among this selection of 129 leader sequences. For example, several T-box sequences were identified (LLMG_R001/041/071/113/124) that use tRNA molecules to regulate the expression of aminoacyl-tRNA synthetase genes and genes involved in amino acid uptake and biosynthesis (58). Leaders were observed with putative riboswitches for flavin mononucleotide (FMN), fluoride, lysine, purine, thiamine pyrophosphate (TPP) and pre-queuosine 1 (preQ1). Four mRNAs with leaders from the pyrimidine biosynthesis pathway (LLMG_R046/054/055/081 or pyrR, carB, pyrK and pyrE) contain binding domains for the pyrimidine biosynthesis regulator protein PyrR. These leaders can form structures that result in anti-terminators in the absence of UMP-bound PyrR, after which transcription can proceed (59). When pyrimidines and UMPs are abundant in the cell, PyrR can form a stabilizing anti-antiterminator structure, preventing the RNA polymerase from further transcribing the downstream genes involved in pyrimidine biosynthesis. Only a limited number of riboswitches have been reported to date and this figure seems an underestimate considering the enormous potential contained within riboswitch RNA-ligand interactions. Clearly, current bioinformatics and genetic approaches need to be adapted to uncover novel RNA sensing structures (30). The challenge is to be able to predict binding ligand-RNA interactions, which when successful would allow devising synthetic riboswitches for novel applications in e.g. medicine or biotechnology (29). Reporting experimentally validated leader sequences could serve as guideline for further experimental research in L. lactis.

RNA-seq reveals 134 new (s)ORFs in L. lactis MG1363. Small proteins of 50 amino acid residues or less are normally not automatically annotated in bacterial genomes while biochemical detection is challenging (60). The rapid increase in the amount of high-resolution transcriptome data and the use of novel techniques such as ribosomal profiling (61) have led to the identification in bacterial genomes of ever more potentially novel genes for small proteins (≤50 amino acid residues). We scanned the leader sequences, internal promoters, sRNAs and asRNAs recognized here for the presence of small open reading frames (sORFs). To this end, the regions comprising the 250 nucleotides downstream of the TSS were analyzed using the following criteria: occurrence of a minimal RBS sequence (NNGGN_{5-14} (A/T/G)TG), the presence of an ORF of ≥20 codons using any of the three START codons AUG, UUG, GUG and a STOP codon within a 250-nt distance from the TSS. Leaderless
transcripts were also mined for ORFs ≥20 codons. A total of 134 novel ORFs were identified in this way, ranging in size from 21 to 61 codons. The putative gene products were then examined for the presence of conserved protein domains using InterProScan 5 (62). None of the 134 deduced proteins, however, contained a known protein domain. This is most likely due to the small size of the proteins as well as to the fact that only a limited number of such small proteins have been characterized in other organisms. Notwithstanding this, some of the ORFs identified here might not be actually protein encoding.

A total of 23 in-sense promoters, internal to known ORFs, were recognized on the basis of transcript abundance relative to surrounding reads and the presence of a promoter upstream of the TSS. Fifteen of these lead to transcripts that are predicted to encode a shorter version of the full-length protein. Interestingly, in four cases the shorter transcript carries an ORF in another reading frame, which would lead to a new protein. The remaining transcripts contain an ORF but lack a minimal RBS.

The transcriptome map was also inspected for differences with the currently available genome annotation of *L. lactis* MG1363 (38), genbank accession number NC_009004. The translation start site of 17 published ORFs is corrected here, since their start codons were either up- or downstream of the TSS determined in this study, leading to a shorter or longer ORF. For three genes with unknown functions (*llmg_0305, llmg_2022* and *llmg_2202*), the transcriptome map suggests that they are non-existing for the following reasons. Firstly, no sequence reads were observed for these genes. Secondly, their predicted translation products do not contain any conserved protein domain, and thirdly the TSS of the gene up- or downstream starts within the ORF itself (Figure 1A). Therefore, we propose to remove these locus tags from the genome of *L. lactis* MG1363. A corrected annotation file was created containing the information on the non-coding RNAs and antisense RNAs discussed above and used to update the genbank file with accession number NC_009004. The corrected genome has also been integrated in a JBrowse webpage (http://jbrowse.molgenrug.nl/).

**Leaderless mRNAs are abundant in *L. lactis* and carry a distinct 5’-motif.** To enable fast genome-wide TSS prediction, a dataset of a non-TEX-treated, strand-specific RNA-seq experiment was combined with that of the TEX-treated RNA-seq sample. This method, also referred to as differential RNA-seq (dRNA-seq), was applied in combination with the TSS prediction program TSSer (63). TSSer predicted 1819 TSSs, of which 884 are primary, 744 are orphan, 66 are internal and 125 are antisense. A significantly higher number of orphan TSSs and antisense TSSs were found with TSSer than by manual mining. Most likely, the latter is more restrictive because a TSS together with the probable RNA species was only annotated when a (variation of a) promoter sequence was also present at a correct distance from 5’-enriched sequence reads. Also, the dataset used for manual mining consisted only of reads derived from primary transcripts, excluding processed RNAs.

TSSs resulting in coding transcripts were used to assess the 5’-UTR length distribution in
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*L. lactis* (Figure 1B). Leader sequences larger than 10 nt contain a Shine-Delgarno sequence with the consensus aaGGAg. The most remarkable observation from the length distribution of 5'-UTRs is the relatively high number of mRNAs in *L. lactis* that do not contain a leader sequence (13% of the transcripts derived from primary TSSs), although even much higher percentages of leaderless mRNAs (lmRNAs) have been reported in other microbes such as the desert bacterium *Deinococcus deserti* (64). The gene products of lmRNAs of *Campylobacter jejuni* have been implicated in stress-responses such as DNA repair (65). Genome2D [http://genome2D.molgenrug.nl] predicts the 111 *L. lactis* lmRNAs to be predominantly involved in the Gene Ontology (GO) classes “Nucleotide Metabolism” and “DNA/RNA binding”. A motif search on these lmRNAs using MEME(46) uncovered an AUGaaa motif overlapping the AUG start codon (Figure 1B). This A-rich sequence might play a role in binding of ribosomes to the lmRNA, in addition to the conserved AUG reported earlier to be necessary for ribosomal binding (66).

*L. lactis* contains only one sigma factor, RpoD (σ70, Llmg_0521). A motif search in the genome regions 50 nt upstream of all 1819 TSSs predicted by TSSer using MEME revealed the known extended tgnTAtAAT consensus for the -10 or Pribnow box while a MAST search (67) with this promoter consensus pinpointed the standard *L. lactis* constitutive promoter sequences -35 (TTGACA) and -10 (TATAAT) (Figure 1C). As is clear from the sequence in Figure 1C, a number of AT-rich stretches can be observed in the promoter region. In contrast to the promoter consensus in the Gram-negative bacteria *C. jejuni* and *H. pylori*, a clear -35 box is present overlapping one of these AT-stretches (35, 68).

Recently, more insight in the complexity of bacterial operon structures is rapidly gained from genome-wide transcription studies employing DNA microarray technology and, lately, RNA-seq. Transcripts of operons can vary in length as a consequence of transcriptional read-through or the presence of secondary promoters, while internal promoters can lead to additional “suboperons”. We used Rockhopper (69) on both RNA-seq datasets and identified 1288 monocistrons and 432 polycistronic operons containing two or more genes in *L. lactis*. The average gene number for polycistronic operons was 2.7, identical to that established in *Neisseria gonorrhoeae* (70). The actual number of operons is expected to be significantly higher as a result of the variations mentioned above.

**Experimental validation of 12 novel RNA candidates.** Northern hybridization was performed on a selection of 12 of the newly identified RNAs to verify the RNA-seq results and to determine the lengths of these RNAs. The expression of 7 sRNAs from IGRs, 2 sRNAs from 3'-UTRs and 3 asRNAs was examined in different phases of growth of *L. lactis* in GM17 medium and under various stress conditions. All 12 RNAs are present under at least one of the conditions employed (Figure 2).

Of the 7 sRNAs from IGRs, LLMGnc_064 (~160 nt) and LLMGnc_138 are constitutively and most highly expressed. According to the Northern blot LLMGnc_138 comprises ~75 nt,
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Figure 2. Experimental validation of novel RNAs. Detection of 9 sRNAs and 3 asRNAs by Northern hybridization in total RNA isolated from L. lactis MG1363 grown in GM17 under various conditions (Ex: exponential phase, St: stationary phase, pH: 10 min acid (pH 4.5) stress, Sa: 10 min salt (2.5% w/v extra NaCl) stress, St*: 10 min starvation in PBS). The positions of the sRNAs and asRNAs are indicated with asterisks (*). As a control, all blots were probed with an oligonucleotide targeting 5S RNA. Visualization of the relevant chromosomal locus and the expression levels of the genes (as derived from the TEX-treated RNA-seq dataset) are given below each blot, // signifies that not the entire gene is shown.
albeit that the LLMGnc_138 sequence reads and the presence of a terminator structure suggest that it has a size of ~100 nt. Possibly this 100-nt RNA molecule is processed into its mature form of ~75 nt. LLMGnc_064 contains an ORF starting 61 nt from its 5'-end that theoretically encodes a small protein of 24 amino acid residues. LLMGnc_010 (~110 nt) is mainly expressed during exponential phase and at low pH. LLMGnc_072 (~290 nt) is most abundant under a high salt condition, suggesting that it is involved in regulating processes related to osmolarity. Two sRNA candidates located in 3'-UTRs, LLMGnc_172 within argR and LLMGnc_177 within zitRS, are each represented with two major RNA species on the Northern blots. The upper band identified with the LLMGnc_172 probe represents the entire argR-LLMGnc_172 transcript of ~575 nt as it also hybridized to a probe for the argR gene (data not shown), while the smallest represents LLMGnc_172 (~65 nt). In the blot probed for LLMGnc_177 (~68 nt), the upper band is likely zitRS, which is ~1400 nt in length. The middle band of a transcript of around 130 nt may be derived from zitRS transcript processing while the lowest band corresponds in size to LLMGnc_177. LLMGnc_087 (~200 nt) is expressed solely during the high-salt or low-pH conditions employed here and could, thus, play a role in *L. lactis* MG1363 coping with sudden changes in osmolarity and pH. The possibility that this RNA encodes a small protein of 31 AA residues is currently under investigation. The protein does not contain a conserved domain and, thus, no function could be predicted.

Two distinct bands are also visible on the Northern blot when probing for the LLMGnc_094 transcript. The upper one corresponds to a size of ~180 nt. Manual inspection of the RNA-seq reads of LLMGnc_094 shows that transcription termination and/or processing occurs after ~72 nt for a small fraction of the RNA. This 72-nt-long molecule is more abundant during the stationary phase and might therefore function as a small regulatory RNA during this phase of growth. Probing for LLMGnc_184 revealed several bands, the upper one of which (~200 nt) is constitutively expressed and seems to correspond with the sequence reads and a predicted terminator structure. The origin of the RNA in the double bands at around 65 nt is unknown but both seem to be more abundant during the stationary phase of growth and in starvation.

The putative as-ps118 is specified by the incomplete *L. lactis* MG1363 prophage MG-1 (55). It starts 80 nt upstream of the stop codon of ps118, directing towards the 3’-end of ps118, a gene encoding a predicted transcriptional regulator. Based on the Northern analysis, as-ps118 is ~70 nt. The RNA-seq data shows that the genes ps119 and ps118 surrounding as-ps118 are silent under the conditions examined here. It is therefore not possible to predict whether or not as-ps118 overlaps with the 3’-UTR of ps118, or even with the 5’-UTR of ps119, although this might be a very likely scenario.

The gene for the antisense RNA as-ps108 is also located on the defective prophage MG-1. The transcript overlaps the 5’-end of ps108. Since the ps108 gene is part of a putative large operon starting with ps115, as-ps108 might affect the stability of the entire transcript.
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The expression of *L. lactis* 6S RNA is carbon source dependent. The widely conserved global regulator 6S (LLMGNc_004) is located in the *L. lactis* chromosome downstream of the *mtl* operon and upstream of the genome of prophage MG-1. A catabolite-responsive element (cre) sequence was predicted immediately upstream of the putative -35 box of the promoter of 6S ([Figure 3A](#)). This suggests that expression of the 6S RNA is under the control of the carbon catabolite repression protein CcpA. Transcriptome analysis using RNA-seq indeed showed that 6S is upregulated ~3-fold after deletion of the *ccpA* gene (unpublished data). Earlier work in *E. coli* has shown that four transcriptional regulators (FIS, H-NS, LRP and StpA) can affect the expression of 6S (71). Northern hybridization using RNA from cells collected at six different points in time during growth on three alternative carbon sources revealed that *L. lactis* S6 is not only abundant during the stationary phase, as is the case in many other organisms, but also highly expressed in the exponential phase when galactose or cellobiose (but not fructose) is provided as the sole carbon source ([Figure 3C](#)). This further confirms involvement of CcpA, since CcpA repression in *L. lactis* is relieved by galactose and cellobiose, but not by fructose (72). As observed in for example *S. pneumoniae* (73), two bands of 6S could represent processed forms of the RNA since the predicted length of 6S is 202 nt ([Figure 3B](#)), which is slightly longer than calculated from the Northern analysis ([Figure 3C](#)). This might also explain why the secondary structure prediction of 6S does not contain a typical central region (74). The longest fragment changes under some of the conditions employed while the short form does not except when the cells are growing with galactose. Overexpression of 6S for 10 min affected only a limited number a transcripts, as determined by RNA-seq ([Figure 3D](#)). Besides three tRNA species and an sRNA with unknown function, two genes were significantly affected ≥2 fold. The gene for peptide deformylase (*llmg_0532*) was downregulated and that of a hypothetical protein (*llmg_1640*) transcribed as part of an operon together with an ABC transporter gene (*llmg_1639*) was upregulated. The relatively short pulse of overexpression of 6S could explain these subtle changes in the *L. lactis* transcriptome. Also, S6 could play a role in fine-tuning of the CcpA regulon, acting when CcpA repression is relieved during stationary phase and/or growth on alternative carbon sources.
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Transcriptome landscape of Lactococcus lactis reveals many novel RNAs including a small regulatory RNA involved in carbon uptake and metabolism.

Figure 3. Analysis of the L. lactis non-coding 6S RNA. (A) Genomic region of the 6S (LLMGnc_004) gene of L. lactis MG1363. Open reading frames are depicted as grow arrows, 6S is shown in blue. Solid black arrows: promoters. The nucleotide sequence of the 6S promoter (P$_{6S}$) is given in capitals, including a predicted cre-site (small letter type) upstream of the -35 box. (B) Structure of 6S RNA predicted by Mfold (83). (C) Detection of 6S RNA by Northern hybridization in samples of L. lactis grown in GM17 until the indicated ODs at 600 nm, or until OD$_{600} = 0.6$ in M17 with 1% of the indicated sugars. 5S RNA served as an RNA concentration control. In addition, one lane of the 8% polyacrylamide gel was after stained with ethidium bromide (EtBr) to visualize the relative amounts of 5S and 6S RNAs. O.N.: overnight culture, OD 2h 2.0: cells taken from a culture maintained at OD$_{600} = 2.0$ for 2 hours. (D) Volcano plot showing the differentially (p-value <0.01 and ≥2-fold) expressed genes upon overexpression of 6S RNA in comparison with the control, using a short (10 min) pulse of nisin addition to a culture of L. lactis SVDM2001 in GM17 and at an OD$_{600} = 0.45$. Indicated in yellow: differentially expressed genes, grey circles: measure of expression level.

The sRNA LLMGnc_147 is involved in carbon uptake and metabolism. As a further demonstration of the validity of the reported sRNAs and to initiate the functional analysis of these putative regulator molecules in L. lactis, we characterized one of them in more detail. LLMGnc_147 is a 102-nt-long sRNA of which the gene is located between llmg_1960.
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Transcriptome landscape of Lactococcus lactis reveals many novel RNAs including a small regulatory RNA involved in carbon uptake and metabolism and the transcriptional activator gene tenA (Figure 4B). Its promoter carries a possible cre site overlapping the -35 box, suggesting that LLMGnc_147 is under control of CcpA (Figure 4A) and related to carbon utilization. Northern analysis shows that LLMGnc_147 is highly expressed in cells growing on cellobiose (Figure 4C). A transcriptional fusion of the promoter of LLMGnc_147 to gfp confirmed that it was most active in the presence of cellobiose and to a lesser extent with galactose (Figure 4D).

To gain insight in potential mRNA targets, LLMGnc_147 was pulse-expressed for 10 min after which total RNA was isolated from the cells and subjected to RNA-seq. Clearly, one operon was highly upregulated (23 to 60-fold) as a consequence of the pulse of LLMGnc_147 RNA (Figure 5A). The six genes of this operon specify the following predicted functions: a PTS transporter (llmg_0963), two beta-glucosidases (llmg_0959/0960), a ribulose-phosphate 3-epimerase (llmg_0957), a ribose-5-phosphate isomerase B (llmg_0958) and an AraC transcriptional regulator (llmg_0962). Table 1 provides a complete list of differentially expressed genes upon pulse-expression of LLMGnc_147. Since a specific substrate has not yet been identified for this putative carbon utilization operon, we determined the effect of LLMGnc_147 overexpression on the ability of L. lactis to switch from glucose to another carbon source. Of the various sugars tested (Figure 5C) only galactose seemed to have a beneficial effect on growth, without the lag-phase seen for the control strain, upon overexpression of LLMGnc_147 (Figure 5B). As the operon llmg_0957-llmg_0963 specifies a putative ribulose epimerase and a ribose isomerase, a pentose sugar rather than galactose, a C-4 epimer of glucose, was expected to be its substrate. Possibly, galactose is imported via the PTS IIC component specified by llmg_0963. On the other hand, overexpression of LLMGnc_147 led to a slower growth phenotype on glucose, mannose, fructose and the disaccharide cellobiose. Only the latter sugar induced the LLMGnc_147 (Figure 4C/D). We assume that slower growth on these sugars after LLMGnc_147 overexpression is caused by a titration effect of the PTS IIC component on cytoplasmic components of specific phosphotransferase systems, as has been reported previously for CelB/PtcAB (75, 76).

Altogether, we show that the expression of LLMGnc_147 is controlled by galactose and cellobiose, the latter potentially via a cellobiose-specific transcriptional activator such as the AraC transcriptional regulator (llmg_0962) from the operon controlled by LLMGnc_147. This might be a remnant of hemicellulose utilization, as L. lactis has a plant origin (38). CcpA might be negatively involved in the regulation of LLMGnc_147 and llmg_0957-llmg_0963, since a cre site is located in their promoter regions. We hypothesize that LLMGnc_147 is necessary for the llmg_0957-llmg_0963 operon in order to stabilize its long transcript.
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Transcriptome landscape of *Lactococcus lactis* reveals many novel RNAs including a small regulatory RNA involved in carbon uptake and metabolism

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Transcriptome landscape of *Lactococcus lactis* reveals many novel RNAs including a small regulatory RNA involved in carbon uptake and metabolism.

Figure 4. The sRNA LLMGnc_147 is involved in carbon metabolism. (A) Genomic region of LLMGnc_147. Open reading frames are depicted as grey arrows, LLMGnc_147 in shown in blue. Solid black arrows: promoters. The nucleotide sequence of the LLMGnc_147 promoter (PLLMGnc_147) is given, including a predicted cre-site that overlaps the -35 box. (B) Structure of LLMGnc_147 using Mfold [83]. (C) Detection of LLMGnc_147 by Northern hybridization in samples of *L. lactis* grown in GM17 until the indicated ODs at 600 nm, or until OD_{600} = 0.6 in M17 with 1% of the indicated sugars. 5S RNA serves as a loading control. (D) P_{LLMGnc_147::gfp} activity in *L. lactis* MG1363 (wt) and SVDM2003 cells grown in M17 containing 1% (w/v) of the indicated carbon source. Fluorescence and optical density were measured five hours after re-inoculation from an overnight culture growing in GM17. The experiment was repeated three times and error bars are indicated.
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Transcriptome landscape of *Lactococcus lactis* reveals many novel RNAs including a small regulatory RNA involved in carbon uptake and metabolism.

Figure 5. LLMGnc_147 is involved in the utilization of galactose. (A) Volcano plot of genes that are differentially expressed (P-value <0.001 and ≥2-fold change) after pulse-expression of LLMGnc_147 via a 10-min addition of nisin to a culture at an OD<sub>600</sub> = 0.45. For clarification of symbols, see the legend to Figure 3. (B) Nisin-induced overexpression for 20 min of LLMGnc_147 (red triangles) in *L. lactis* SVDM2002 in comparison with the empty vector control (blue squares), after which the strain was re-inoculated 1:20 in fresh M17 medium containing 1% galactose. (C) Identical experimental setup as described in (B) for growth after re-inoculation in M17 with 1% (w/v) of the indicated carbon sources. The experiments were repeated twice and the lines represent averages of four microtiter plate measurements.

Conclusions

Differential RNA sequencing uncovered hundreds of novel RNAs in the genome of the lactic acid bacterium *L. lactis*, for which no sRNAs have been described so far. These may shed light on the interpretation of specific (point) mutations obtained e.g. after evolutionary
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Transcriptome landscape of *Lactococcus lactis* reveals many novel RNAs including a small regulatory RNA involved in carbon uptake and metabolism. The output of this study provides an excellent basis for further investigations on the molecular biology of *L. lactis* and a starting point for the characterization of the putative regulatory RNAs in this organism.

**MATERIALS AND METHODS**

**Bacterial strains, growth conditions, cloning and nisin induction.** For an overview of strains and plasmids used in this study, see Table 2. *L. lactis* was routinely grown as standing cultures at 30°C in M17 broth (Difco, Becton Dickinson, Le Pont de Claix, France) containing 0.5% (w/v) glucose (GM17). The restriction and ligation independent USER fusion cloning strategy (77) was employed for vector constructions. In short, vector backbone and insert fragments were amplified with PfuX7 polymerase. PCR fragments were then purified using a PCR clean-up kit (Macherey-Nagel GmbH, Germany) and inspected for correct size by agarose gel electrophoresis. PCR backbone and sRNA gene or promoter inserts were mixed at a 1:3 molar ratio, treated with the USER enzyme mix (New England Biolabs, Hitchin, UK) and directly introduced into competent cells of *L. lactis* NZ9000 by electroporation (at 2.5 kV, 25 uF, 200 Ohm) or by heat-shock (45 sec at 42°C) into chemically competent *E. coli* Dh5α cells. Colony PCR and subsequent sequencing (Macrogen, Amsterdam, The Netherlands) was used to verify construct correctness.

The promoter of LLMGnc_147, P<sub>LLMGnc_147</sub>, was PCR-amplified and cloned upstream of the GFP gene in plasmid pSEUDO_10, which was subsequently integrated in the chromosome of *L. lactis* MG1363 via single cross-over recombination (78). To stably maintain the integrated plasmid, cells were selected and continuously grown in the presence of 3 µg/ml erythromycin.

*L. lactis* NZ9000 was used for overexpression of sRNAs from the high-copy number plasmid pNZ8048. A single colony from a GM17 agar plate containing 5 µg/ml chloramphenicol was used to inoculate 10 ml of fresh GM17 medium. After overnight growth, the culture was diluted 1:100 and incubated until an OD<sub>600</sub> between 0.4-0.5 was reached. Subsequently, the cells were induced for 10 min with 7.5 ng/ml nisin (Sigma-Aldrich, Munich, Germany), and harvested as described above.

Optical density and fluorescence were measured in a Tecan F200 (Tecan Group, Männedorf, Switzerland).
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### Table 2

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Cm\(^{-}\), chloramphenicol resistance marker

Em\(^{-}\), erythromycin resistance marker

RNA isolation. For dRNA-seq, single colonies of *Lactococcus lactis* MG1363 or *L. lactis* NCDO712, grown on GM17 (1.5%) agar plates, were used to inoculate 10 ml fresh GM17 media for overnight growth at 30°C. The overnight cultures were each diluted 1:100 in 500 ml GM17. *L. lactis* MG1363 was sampled at three points in time in the exponential phase (OD\(_{600}\) of 0.9, 1.3 and 1.7) and at three time points in the stationary phase (at 30, 60 and 90 min after an OD\(_{600}\) of >2.5 was reached). To compensate for cell density, equivalent OD units were harvested by centrifugation at 10,000 rpm for 1 min; the cell pellets were immediately frozen in liquid nitrogen. Cells from *L. lactis* NCDO712 were harvested at the mid-exponential growth phase (OD\(_{600}\) of 1.0).

For RNA isolation, cell pellets were re-suspended in 400 µl TE-buffer (10 mM Tris-HCl, 1 mM EDTA, pH 7.4), after which 50 µl 10% Sodium dodecyl sulfate (SDS), 500 µl phenol/chloroform and 0.5 g glass beads (75-150 µm in diameter) were added. The mixture was cooled on ice and the cells were subsequently disrupted by two consecutive rounds of shaking for 45 sec in a Mini-BeadBeater (Biospec Products, Bartlesville, OK, USA) at 4°C, with intervening cooling on ice. After centrifugation (14,000 rpm for 10 min), the supernatants were treated with 500 µl chloroform, centrifuged as above, and the water phase was collected. Total RNA from the water phase was incubated for 30 min at 37°C with RNase-free DNase I supplemented with RiboLock RNase inhibitor (Fermentas/Thermo Scientific, Vilnius, Lithuania). The RNA was subsequently purified using standard phenol/chloroform extraction followed by sodium acetate/ethanol precipitation. RNA pellets were dissolved in TE-buffer. All solutions were treated with DEPC and autoclaved.
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RNA treatment, library preparation and RNA deep sequencing. RNA concentrations were measured using a Nanodrop ND-1000 (Thermo Fischer Scientific, Rockford, IL, USA), after which the integrity of the 16S/23S rRNA and DNA contamination were assessed using an Agilent 2100 Bioanalyser (Agilent Technologies, Waldbronn, Germany). The RNA sample from *L. lactis* MG1363 was TEX-treated at Eurofins MWG GmbH (Ebersberg, Germany). rRNA depletion was done with the Terminator System kit (Epicentre, Madison, WI, USA), preparation of a 5’-fragment cDNA library was performed as previously described (35). After PCR amplification and library purification, the library was sequenced on an Illumina HiSeq2000 v3 platform (Illumina, San Diego, CA, USA), with a paired-end protocol and read length of 101 nt, resulting in a total output of 10.5 million (M) reads. The RNA sample for the non-TEX treated library (RNA sample from *L. lactis* NCDO712) was sequenced at Otogenetics Corporation (Norcross, GA, USA) on an Illumina HiSeq2000, with a ScriptSeq™ Complete Kit (Bacteria) (Epicentre, Madison, WI, USA) including Ribo-Zero™ rRNA removal and ScriptSeq v2 library preparation for directional RNA-Seq, resulting in a total of 15.7 M reads. RNA samples from the pulse-expression of LLMGnc_147, after Ribo-Zero™ rRNA removal and library preparation using the AmpliSeq™ kit (ThermoFischer Scientific), were sequenced at the PrimBio Research Institute (Exton, PA, USA) on a Ion Proton sequencer. This resulted in 13-23 M reads per sample.

TSS calling and data analysis. RNA-seq data of TEX-treated and untreated samples was used for automated TSS calling by TSSer (63), using default parameters. Predicted TSSs were used to perform a MEME search to identify promoter motifs and Shine-Delgarno sequences in the regions -50 to -1 upstream of all TSSs using a zero or one occurrence model. From the 111 leaderless mRNAs predicted by TSSer, a MEME motif search was performed in the region +1 to +15 downstream of the TSS, based on a one occurrence per sequence model, using an E-value threshold of 0.001. Operons in *L. lactis* were predicted by Rockhopper (69), using default settings.

For manual qualitative mining, raw data reads of 101 nt of the TEX-treated sample were quality trimmed with a PHRED score >28 and subsequently aligned to the genome of *L. lactis* MG1363 (38) using Bowtie2 (79). The resulting reads were visualized with Genome2D (80) displaying known ORFs and putative regulatory RNA elements predicted in *silico* for *L. lactis* MG1363 using SIPHT (45). This data was manually inspected for novel RNA elements and re-annotation purposes. To prevent false positive calling of transcription start sites (TSS), especially with respect to lowly expressed RNAs, TSSs were inspected manually for promoter motifs (-10 and -35 boxes), after which transcription start sites were extracted using Tablet (81). Promoters of sRNAs and asRNAs were assessed for the presence of transcription factor binding sites (TFBS) by performing TFBS searches on http://genome2d.molgenrug.nl/, using -100 to -1 upstream of the TSS.

Comparative transcriptome data was quality checked as above and trimmed with a cut-off
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>30 nt, after which analyses were performed using the Transcriptome analysis webserver for RNA-seq expression data (T-REx) (82).

**Northern hybridization.** Total RNA (10 μg) was separated on an 8% denaturing polyacrylamide-(7 M)urea gel in Tris-acetate-EDTA buffer (TAE). RNAs were transferred to positively charged Zeta-Probe nylon membranes (Bio-Rad Laboratories BV, Veenendaal, The Netherlands) using a semi-dry electroblotting apparatus (Bio-Rad Laboratories BV). RNAs were covalently cross-linked to the membranes at 1200 mJ using a UVC-508 Ultraviolet Crosslinker (Ultra-Lum Inc., Carson, CA, USA), after which the blots were hybridized overnight at 42°C in PerfectHyb Plus Hybridization buffer (Sigma-Aldrich Chemie Gmbh, Munich, Germany), using appropriate 32P-labeled DNA oligonucleotides. DNA probes were labeled with 32P-γATP using Polynucleotide kinase (Fermentas/Thermo Scientific), according to the manufacturer. Nylon membranes were washed twice in 2x saline sodium citrate (SSC) buffer with 0.1% SDS, exposed to a Phosphor Screen and imaged using a Cyclone Plus Phosphor Imager and OptiQuant software (PerkinElmer, Groningen, NL).

**ACKNOWLEDGEMENTS**

We kindly thank Akos T. Kovacs and Herwig Bachmann for helpful discussions and Mikkel Jørgensen for his help with the Northern analyses. We also would like to thank Ana Solopova for her insights in *L. lactis* carbon metabolism.
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