Unleashing Natural Competence in *Lactococcus lactis* by Induction of the Competence Regulator ComX

Joyce Mulder,a,b,c Michiel Wels,b Oscar P. Kuipers,a,c Michiel Kleerebezem,c,d Peter A. Bronb,c

Molecular Genetics, University of Groningen, Groningen, The Netherlands; NIZO B.V., Ede, The Netherlands; BE-Basic Foundation, Delft, The Netherlands; Host-Microbe Interactomics Group, Animal Sciences, Wageningen University, Wageningen, The Netherlands

**ABSTRACT** In biotechnological workhorses like *Streptococcus thermophilus* and *Bacillus subtilis*, natural competence can be induced, which facilitates genetic manipulation of these microbes. However, in strains of the important dairy starter *Lactococcus lactis*, natural competence has not been established to date. However, in silico analysis of the complete genome sequences of 43 *L. lactis* strains revealed complete late competence gene sets in 2 *L. lactis* subsp. *cremoris* strains (KW2 and KW10) and at least 10 *L. lactis* subsp. *lactis* strains, including the model strain IL1403 and the plant-derived strain KF147. The remainder of the strains, including all dairy isolates, displayed genomic decay in one or more of the late competence genes. Nisin-controlled expression of the competence regulator *comX* in *L. lactis* subsp. *lactis* KF147 resulted in the induction of expression of the canonical competence regulon and elicited a state of natural competence in this strain. In contrast, *comX* expression in *L. lactis* NZ9000, which was predicted to encode an incomplete competence gene set, failed to induce natural competence. Moreover, mutagenesis of the *comEA-EC* operon in strain KF147 abolished the *comX*-driven natural competence, underlining the involvement of the competence machinery. Finally, introduction of nisin-inducible *comX* expression into *nisRK*-harboring derivatives of strains IL1403 and KW2 allowed the induction of natural competence in these strains also, expanding this phenotype to other *L. lactis* strains of both subspecies.

**IMPORTANCE** Specific bacterial species are able to enter a state of natural competence in which DNA is taken up from the environment, allowing the introduction of novel traits. Strains of the species *Lactococcus lactis* are very important starter cultures for the fermentation of milk in the cheese production process, where these bacteria contribute to the flavor and texture of the end product. The activation of natural competence in this industrially relevant organism can accelerate research aiming to understand industrially relevant traits of these bacteria and can facilitate engineering strategies to harness the natural biodiversity of the species in optimized starter strains.

**KEYWORDS** *Lactococcus lactis*, natural competence, comparative genomics
a chemically defined medium (7). When Gram-positive bacteria enter a state of natural competence, exogenous DNA translocates through the DNA uptake machinery, a multiprotein complex comprising the proteins ComEA, ComEC, ComFA, and ComFC and a nuclease (EndA in S. pneumoniae) encoded by the late competence (com) genes (8, 9). Other late competence genes encode proteins that compose pilus-like structures (ComGA-GG) or protect internalized DNA against degradation (SsbA, SsbB, DprA, and RecA) (8–10). Expression of these genes is positively regulated by the competence master regulator ComX, which acts as an alternative sigma factor (11–13). In S. thermophilus, expression of comX is initiated upon formation of the quorum-sensing ComRS complex, comprising the pheromone-like peptide ComS and transcriptional regulator ComR and encoded by the comRS operon (14, 15). Addition of a synthetic peptide that resembles the active competence pheromone has proven to be a successful strategy to induce natural competence in several bacterial species. For example, addition of a synthetic ComS peptide to S. thermophilus cultures in the early logarithmic phase of growth enabled the activation of natural competence and highly efficient DNA transformation (14, 16). Analogously, other streptococci, including S. pneumoniae, utilize the comCDE regulatory module to control natural competence, involving the competence-stimulating peptide (CSP) (encoded by comC) and a two-component system (encoded by comD and comE [17, 18]), and the addition of synthetic CSP leads to development of natural competence in this species.

Strains of Lactococcus lactis are of great importance in the dairy industry, primarily in the production of cheese, butter, and buttermilk (19). So far, a comRS- or comCDE-like system has not been identified in L. lactis. Nevertheless, complete sets of late competence genes appear to be present in several L. lactis genomes (20–22; this study). In addition, increased expression of competence genes has been observed in L. lactis subsp. lactis IL1403 and KF147 under specific conditions that included carbon starvation (23, 24). Unfortunately, in neither of these strains, or any other L. lactis strain, could natural competence development be experimentally established (20, 24). As an alternative route to establish natural competence, overexpression of comX has been employed, aiming to enhance expression of the complete late competence regulon. Such an approach has been successful in S. thermophilus (25) but failed in L. lactis IL1403 (20). Nevertheless, the observations that complete sets of late competence genes are apparently present in some of the L. lactis genomes (26, 27) and that their expression can be induced under specific conditions (23, 24) deserve a more dedicated bioinformatic and experimental effort.

Here, we present a comparative genomics analysis of 43 L. lactis genomes to assess their potential to enter a state of natural competence. Moreover, by employment of controlled expression of ComX, we demonstrate enhanced expression of the late competence regulon and concomitant induction of natural competence, which was successful only in strains predicted to encode a complete late competence machinery. The discovery of natural competence in L. lactis will enable transfer of genetic information without the use of genetically modified organisms (GMO), resulting in the improvement of the industrial performance of strains of this species and the enhancement of the quality of fermented products.

RESULTS
Genomic analyses show complete sets of competence genes in several L. lactis strains. To evaluate whether L. lactis strains possess the genetic capacity to enter a state of natural competence, late-competence-associated genes were initially identified in the L. lactis KF147 genome by using the known competence genes of the naturally competent Streptococcus thermophilus LMD-9 (7, 14). This strain was selected for this primary analysis based on previous work that reported that many of the late competence genes were induced in this strain under starvation, nongrowth conditions (24). Similar proteins (similar both in length and sequence) were identified to be encoded within the KF147 genome for all selected late competence genes of S. thermophilus.
LMD-9, corroborating that a complete competence gene set is present in this *L. lactis* strain (see Table S1 and Fig. S1 in the supplemental material).

Subsequently, the identified KF147 competence protein sequences were used as a reference set for identification and comparison to the orthologous groups (OGs) of genes carried by 42 other *L. lactis* strains (21, 22, 26–35). Full-length protein sequence identity to the KF147 OGs was calculated for all 42 *L. lactis* strains (Fig. 1). This analysis revealed considerable genomic decay in several of the strains of both *L. lactis* *subsp.* *lactis* and *L. lactis* *subsp.* *cremoris*. Moreover, for the OGs that were intact, there was a clear distinction between the levels of identity observed for strains belonging to *L. lactis* *subsp.* *lactis* (which includes strain KF147) and *L. lactis* *subsp.* *cremoris* (Fig. 1), exemplifying the genetic distinction between these two subspecies (36). Among the strains belonging to *L. lactis* *subsp.* *cremoris*, only strains KW2 and KW10 appeared to encode full-length homologues of all the late competence proteins selected in KF147, albeit with identity scores ranging from 56 to 99% (Fig. 1). Notably, when the late competence gene set of the KW2 strain instead of that of strain KF147 was used to determine full-length protein sequence identity levels, it was apparent that late competence proteins displayed a high degree of conservation within *L. lactis* *subsp.* *cremoris* but were distinct from their orthologues in *L. lactis* *subsp.* *lactis* (see Fig. S2 in the supplemental material). Among the 28 strains belonging to *L. lactis* *subsp.* *lactis*, at least 10 appeared to encode a full set of late competence proteins.

**FIG 1** Genomic analysis of 43 *L. lactis* strains to assess genetic capacity to develop natural competence. A concatenated core genome single-nucleotide polymorphism (SNP) tree of 43 *L. lactis* strains was combined with full-length protein identity scores (%) for the selected subset of late-competence-associated proteins in comparison to their homologues in *L. lactis* strain KF147, which was used as a reference. Protein identity scores are depicted within each cell and reflected by gray scales based on the *L. lactis* KF147 query protein sequences, in which at least 90% full-length alignment is considered indicative of gene presence. Genetic events leading to competence gene decay (black cells in the figure) are specified as a premature stop codon within the first 90% of the gene (a), transposon insertion (b), prophage insertion (c), or absence of gene, a mutated/alternative start, or lengthened/fused protein at least more than 25% of its total length (d), followed by the position within the protein sequence where the event is detected relative to its N terminus. Source of isolation: P, plant; D, dairy; S, soil; W, water; H, human body; F, fruit (72). References for the genome sequences are given when available.
The genomic decay within these late competence genes in the *L. lactis* subsp. *cremoris* strains displayed several conserved disruptive mutations in specific genes, including IS982 insertions in *comEC* (strains SK11, A76, and UC509.9) and *comGA* (strains SK11 and A76), although with some variation with respect to the precise position of insertion (Fig. 1; see Fig. S3 in the supplemental material). Various strains of *L. lactis* subsp. *cremoris* contained conserved premature stop codons within one or more of their late competence genes, suggesting that these strains derive from a common ancestor, in which conserved and strain-specific mutations have shaped the decay pattern of the late competence genes. For example, strains SK110, AM2, SK11, A76, UC5099, B40, FG2, HP, and LMG6897 share similar mutational events in *comEA*, *comEC*, *comFA*, and *comGD*, whereas strains N41, NCDO763, and MG1363 harbor common mutations in *comEC* (see Fig. S4 in the supplemental material). In contrast, the disruptive mutations observed in the late competence genes of strains of *L. lactis* subsp. *lactis* appeared to be more scattered (Fig. 1), suggesting that degenerative mutations accumulated more recently in this subspecies. Nevertheless, several strains (KF282, KF24, N42, CV56, ML8, KLDS, UC317, and KF67) contain a (remnant of a) prophage insertion within the *comGC* gene (see Fig. S5 in the supplemental material). Remarkably, these phage sequences are always inserted at the same position within the *comGC* sequence, suggesting site-specific integration at a conserved sequence element within the *comGC* gene.

In summary, these findings indicate that in the majority of *L. lactis* strains, one or more late competence functions are compromised, suggesting that these strains are not able to develop a state of natural competence. The analysis also implies that in some strains, including *L. lactis* KF147, the genetic capacity to enter a state of naturally competent appears to be intact. Finally, it is noteworthy that within the present panel of strains, there are no dairy isolates that appear to encode a complete set of intact late competence proteins, which may reflect the high level of genome decay that has been reported for strains in the milk environment before (37–39).

**Moderate overexpression of the late competence regulon regulator ComX results in a state of natural competence in *L. lactis* KF147.** In order to test whether the identified competence machinery can be activated and is functional, we set out to overexpress the predicted competence regulator ComX. From the subset of strains predicted to harbor a complete set of competence genes, *L. lactis* KF147 harbors a chromosomal copy of nisRK but does not produce nisin (40), allowing nisin-inducible *comX* expression by cloning of this gene under the control of PnisA in pNZ8150 (41). This *comX* expression strategy led to a dose-dependent inhibition of growth (Fig. 2A), which was not observed in the control strains harboring pNZ8150 (see Fig. S6 in the supplemental material) (41) or pNZ8040, a vector enabling nisin-inducible expression of pepN (Fig. S6) (42). Hence, the observed growth retardation is not caused by the addition of nisin or the overexpression of proteins as such but is specifically caused by the presence of ComX.

To investigate the impact of elevated ComX levels on the expression levels of the late competence genes, their transcript levels were determined by reverse transcription-quantitative PCR (RT-qPCR) on RNA derived from *L. lactis* KF147 harboring pNZ8150 or pNZ6200, either uninduced or moderately or fully induced with nisin. Under uninduced conditions, *comX* expression levels were 2.5- to 6-fold increased in *L. lactis* KF147 harboring pNZ6200 compared to the pNZ8150-harboring cells, which likely reflects low-level “promoter leakage” due to the presence of PnisA on a high-copy-number plasmid (Fig. 2B). Induction of *comX* expression in *L. lactis* KF147 harboring pNZ6200 with either 0.03 or 2 ng/ml nisin for 2 h led to 15- to 20-fold and 1,500- to 4,000-fold induction of *comX* expression relative to that in the uninduced control of the same strain, respectively (Fig. 2B). Similarly, expression of the late competence genes *comEA*, *comFA*, and *comGA* was induced, illustrating the strongly enhanced expression of the late competence regulon as a consequence of the elevated levels of its regulator, ComX (Fig. 2B). These induction conditions for the activation of late competence genes were employed to test whether the corresponding phenotype could also be observed, by
adding pIL253 (43) to the culture medium at the same time point that comX induction was initiated using a range of nisin concentrations. As expected, no pIL253 transformants were obtained for *L. lactis* KF147 harboring pNZ8150 under any of the conditions tested (data not shown). In contrast, pIL253 transformants were obtained for *L. lactis* KF147 harboring pNZ6200 following induction with nisin concentrations ranging from 0.005 to 0.1 ng/ml nisin, with an approximate transformation rate of 10\(^{-7}\) to 10\(^{-6}\) (transformants/total cell number/\(\mu\)g plasmid DNA). The highest transformation rates were obtained after induction with 0.03 ng/ml nisin (Fig. 2C). Both strain identity and pIL253 presence was confirmed by PCR in all transformants tested (Fig. 2D). Notably, full nisin induction (2.0 ng/ml nisin) of comX expression in pNZ6200-harboring *L. lactis* KF147 did not result in any transformants. To check whether comX of the *L. lactis* subsp. *cremoris* strain MG1363 is still functional, transformation of *L. lactis* KF147 harboring pNZ6201 upon nisin induction was also tested. Similar results were obtained when comX of *L. lactis* subsp. *cremoris* MG1363 was expressed in *L. lactis* subsp. *lactis* KF147 (Fig. 2E and F), indicating that comX derived from an *L. lactis* subsp. *cremoris* strain is
also fully functional. Taken together, these results demonstrate that activation of moderate expression, but not high-level expression, of endogenous comX in L. lactis KF147 elicits the natural competence phenotype in this strain. The observation that this does not occur at a high level of comX expression may be a consequence of the observed growth defect under these conditions, which may interfere with completion of the competence machinery assembly and/or recovery of potential transformants after plating. Such a notion is supported by the observation that expression of a heterologous copy of comX (derived from L. lactis subsp. cremoris) induced less severe growth defects upon maximal nisin induction and still led to detectable natural competence development, albeit with reduced efficiency compared to that with moderate nisin induction levels.

ComX-induced transformation in L. lactis depends on the late competence operon comEA-EC. The experiments described above do not provide direct proof for a functional dependency of the observed transformation phenotype on the expression of the late competence genes, although this is likely, considering the fact that these genes encode the DNA uptake machinery. Therefore, we constructed a comEA-EC-negative derivative of L. lactis KF147 through the integration of a linear fragment harboring a tetracycline resistance-encoding tetR gene flanked by regions homologous to the 5′ and 3′ regions surrounding the comEA-EC operon. The procedure for moderate comX induction was applied to transform this linear mutagenesis fragment into strain KF147 harboring pNZ6200. Integrants with the anticipated genotype ΔcomEA-EC::tetR (Fig. 3A) were obtained with an efficiency similar to that observed for pIL253 transformation. Subsequent comX expression induction experiments in the ΔcomEA-EC::tetR derivative of strain KF147 (NZ6200) harboring pNZ6200 showed that full induction of comX led to a growth rate reduction in this strain (Fig. 3B) similar to that observed for the parental KF147 strain. Importantly, nisin concentration-dependent comX overexpression and corresponding upregulation of expression of the comFA and comGA, but not comEA, genes was similar to what was established in L. lactis KF147 harboring pNZ6200 (see Fig. S7 in the supplemental material). However, in contrast to the case for the parental strain KF147, transformation of NZ6200 harboring pNZ6200 with plL253 did not yield any transformants (Fig. 3C), establishing the involvement of the comEA-EC operon in comX-induced competence in L. lactis KF147.

Expansion of the natural competence phenotype to a broader set of L. lactis strains. In order to employ the comparative genomics analysis performed in this study as a predictor for competence potential, L. lactis subsp. cremoris strains KW2 (28), and NZ9000 (41, 44) and L. lactis subsp. lactis IL-9000 (45) were tested for transformability

FIG 3 Natural competence of L. lactis KF147 depends on the late competence operon comEA-EC. (A) By use of a linear mutagenesis fragment, a comEA-EC-negative derivative of KF147 harboring pNZ6200 (NZ6200) could be obtained with high efficiency. (B and C) NZ6200 displayed a similar growth defect upon full nisin induction (B), but transformation with pIL253 is not feasible (C).
upon \(\text{comX}\) induction. Strains NZ9000 and IL-9000 are derivatives of MG1363 and IL1403, respectively, which contain the \(\text{nisRK}\) genes integrated in their chromosome, thereby allowing the use of the nisin-controlled expression system. Strain KW2 does not harbor \(\text{nisRK}\) in its genome, and to facilitate the use of the nisin induction system in this strain, pNZ9531 was first introduced into this strain, thereby expressing \(\text{nisRK}\) in this background from a low-copy-number plasmid vector that is compatible with the pNZB150 backbone of the pNZ6200 and pNZ6201 vectors used for \(\text{comX}\) expression (46). The plasmid pNZB150, pNZ6200, or pNZ6201 was transformed into electrocompetent cells of IL-9000, NZ9000, and pNZ9531-harbor ing \(L.\text{lactis}\) KW2. These transformants were induced with 0, 0.03 or 2 ng/ml nisin, and the induction of the competence phenotype was evaluated in these strains by transformation with \(\text{pIL253}\) (for NZ9000 and IL-9000) or pNZ6202 (a tetracycline-selectable pIL253 derivative). As anticipated, none of the conditions employed allowed the activation of natural competence in strain NZ9000 (Table 1), which is in good agreement with the \(\text{comEC}\) and \(\text{coiA}\) mutations observed in its parental strain MG1363 (Fig. 1). In contrast, transformants were obtained for pNZ6200 and pNZ6201 harboring derivatives of IL-9000, despite its incomplete \(\text{dprA}\) gene, and when these plasmids were transformed to KW2 harboring pNZ9531 (Table 1). Notably, although the efficiency of transformation appeared to be the highest for the cells in which moderate \(\text{comX}\) expression was induced (i.e., with 0.03 ng/ml nisin), transformants also were obtained under uninduced conditions and upon high-level induction of \(\text{comX}\) expression (i.e., with 2 ng/ml nisin). The observed transformation under uninduced conditions might be caused by the previously reported higher levels of “leakage” of the \(\text{nisA}\) promoter activity in \(L.\text{lactis}\) strains harboring the \(\text{nisRK}\) expression vector pNZ9531 (46), whereas the \(\text{dprA}\) mutation in the IL-9000 parental strain (IL1403) (Fig. 1) may require lower levels of \(\text{comX}\) expression to activate competence, as the \(\text{DprA}\) function has been associated with competence shutoff (47, 48). The observation that high-level \(\text{comX}\) expression still allowed competence development in KW2 and IL-9000 despite the growth-inhibitory consequences of this level of induction, which is in contrast with the results obtained for strain KF147, remains to be determined. Finally, in pNZ9531-harbor ing KW2, induced expression of the \(\text{comX}\) derived from \(L.\text{lactis}\) subsp. \(\text{cremoris}\) (i.e., as expressed from pNZ6201) allowed competence development, confirming the bidirectional functional exchangeability of the \(\text{comX}\) genes from these two \(L.\text{lactis}\) subspecies. These results confirm the predictions made by comparative genomics (Fig. 1) with respect to the capacity to develop a natural competence phenotype in \(L.\text{lactis}\) strains, and they establish that strains of both subspecies have the capacity of natural competence which can be induced by controlled expression of the \(\text{comX}\)-encoded regulator from either of the subspecies.

## DISCUSSION

This study demonstrates that the \(L.\text{lactis}\) strains KF147, KW2, and IL1403 possess a functional DNA uptake machinery, which can be activated by the ComX regulator. This

<table>
<thead>
<tr>
<th>(L.\text{lactis}) strain</th>
<th>Plasmid content</th>
<th>Transformation rate ((\text{pIL253/pNZ6202})^a) with the following ([\text{nisin}]) (ng/ml):</th>
</tr>
</thead>
<tbody>
<tr>
<td>IL-9000</td>
<td>pNZ6200, pNZ8150</td>
<td>1.5 × 10⁻⁷ ± 8.2 × 10⁻⁸, 1.7 × 10⁻⁷ ± 1.5 × 10⁻⁷, 3.4 × 10⁻⁷ ± 1.7 × 10⁻⁷</td>
</tr>
<tr>
<td>KW2</td>
<td>pNZ9531 + pNZ6200, pNZ9531 + pNZ6201, pNZ9531 + pNZB150</td>
<td>1.5 × 10⁻⁶ ± 1.1 × 10⁻⁶, 1.1 × 10⁻⁶ ± 1.1 × 10⁻⁶, 1.0 × 10⁻⁵ ± 9.2 × 10⁻⁷, 6.2 × 10⁻⁶ ± 5.7 × 10⁻⁶, 5.2 × 10⁻⁶ ± 3.1 × 10⁻⁶</td>
</tr>
<tr>
<td>NZ9000</td>
<td>pNZ6200, pNZB150</td>
<td>ND, ND, ND</td>
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</tbody>
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Note: The transformation rate was calculated as number of pIL253 or pN6202 transformants/total number of cells/μg DNA. Values are means ± standard deviations. ND, not detected.

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TABLE 1 Assessing transformation with plasmid pIL253/pNZ6202 by controlled expression of \(\text{comX}\) in \(L.\text{lactis}\) IL-9000, NZ9000, and pNZ9531-harbor ing KW2.
implies that identification of a complete set of late competence genes through comparative genomics represents an appropriate approach to predict the capacity of a strain to enter a state of natural competence, and it seems likely that most, if not all, of the other strains identified here as carrying complete gene sets can be made naturally competent via the same strategy of comX overexpression. It should be noted that the expression of a much larger set of over 100 genes is regulated upon addition of the competence pheromone in streptococci (12, 49, 50). For instance, development of natural competence usually occurs in concert with increased expression of proteins involved in DNA recombination, thereby facilitating integration of acquired DNA (51), a feature that has been observed in this study for \( L. \) lactis KF147, as well suggesting expression of such proteins in \( L. \) lactis KF147 upon competence development. Nevertheless, we show that the dedicated assessment of only the canonical late competence genes is a valid predictor for competence potential in \( L. \) lactis strains.

It is commonly assumed that the \( L. \) lactis ancestor strain prior to subspeciation into \( L. \) lactis subsp. lactis and \( L. \) lactis subsp. cremoris originated from a plant-associated niche and that strains adapted to increase their fitness in the nutritionally rich dairy environment (40, 52, 53). Remarkably, none of the dairy isolates of \( L. \) lactis that were analyzed here appear to carry a complete set of late competence genes, suggesting that during the adaptation to the dairy niche, there was no significant environmental fitness benefit associated with the capacity to become naturally competent. This may relate to a real lack of fitness benefit of this phenotype within the dairy environment or may be due to highly consistent suppression of the phenotype during growth in milk, thereby preventing the possible fitness benefit to become apparent, which may allow the decay of encoding genes without an apparent fitness cost for the bacteria. The latter scenario appears to be in agreement with the observed activation of the expression of late competence genes in \( L. \) lactis during carbon starvation conditions (23, 24), which are not likely to occur within the dairy niche, as it is very rich in lactose. The genomic decay events associated with dairy-derived \( L. \) lactis strains include prophage disruption of the comGC locus in strains of \( L. \) lactis subsp. lactis (28, 54) and insertion of IS982 into several com genes in strains of \( L. \) lactis subsp. cremoris (40, 55–57). Notably, the phylogenetic relatedness of \( L. \) lactis subsp. cremoris strains predicted on the basis of competence gene decay events displayed a topology that was remarkably similar to that observed for the core genome relatedness of these strains (see Fig. S4 in the supplemental material). Importantly, typical dairy environment-associated lactic acid bacteria quite commonly display genomic decay as a consequence of the adaptation to this nutritionally rich environment (37–39). For example, loss-of-function events have been observed in \( S. \) thermophilus, Lactobacillus helveticus, and Lactobacillus bulgaricus upon prolonged culturing in milk, with mutations accumulating in genes encoding transport-, energy metabolism-, and virulence-associated functions, implying that these functions do not contribute to fitness in the dairy niche (37, 39, 58, 59). Analogously, experimental evolution of \( L. \) lactis KF147 to enhanced fitness and growth in milk was shown to be associated with suppression of gene repertoires associated with the import and utilization of a variety of typically plant environment-associated carbon sources, as well as mutations leading to functional reconstitution and elevated transcription of the peptide import system (opp) of this strain (52). Paradoxically, dairy strains of \( S. \) thermophilus still possess the genetic and phenotypic capacity to develop natural competence (14, 25, 39, 60), suggesting that competence development in this species contributes to fitness in this habitat. In contrast to the case for \( L. \) lactis, where carbon starvation has been associated with induction of late competence expression (23, 24), similar conditions have not been implied in competence regulation in \( S. \) thermophilus. This may suggest that \( S. \) thermophilus actively expresses the competence phenotype in the dairy environment, which may contribute to this species’ fitness in the milk environment.

In nature, natural competence in bacteria is commonly a transient phenotypic state with a small window of opportunity to take up DNA (61), the activation and shutdown of which are subject to subtle regulation (47, 48) to prevent futile activation of the
costly process and to sustain genomic stability. Analogously, optimal induction in *L. lactis* was achieved with a moderate level of ComX induction, whereas high-level induction of this regulator failed to lead to competence development (strain KF147) or led to significantly reduced levels of transformation (strains KW2, and IL-9000). Analogously, full induction of the late competence gene expression in *Lactobacillus sakei* was achieved by moderate levels of induction of its central regulator *sigH* (62). Moreover, high-level *comX* expression was consistently associated with reduced growth efficiency of the strains used in this study, which is illustrative of the tight connection between competence and growth (63). Previous *comX* expression studies using *L. lactis* IL1403, the parental strain of IL-9000, failed to elicit natural competence (20), which may have been due to inappropriate expression levels of *comX* or might have been caused by the fact that the endogenous *comX* gene of IL1403 was used, which contains an alternative start codon and appears to be truncated.

Taken together, the results of this study show that in *L. lactis* strains that carry complete late competence gene sets, a state of competence can be induced by controlled expression of *comX*; in particular, moderate expression of this regulator appears to be effective in activation of this phenotype. Naturally competent *L. lactis* strains could internalize plasmid and linear DNA from their environment with similar efficiencies. The conditions that naturally activate *comX* expression and contribute to the regulation of competence development in *L. lactis* remain to be established. Unraveling the *in situ* control mechanisms of natural competence in *L. lactis* would offer opportunities to exploit this phenotype for strain improvement purposes in this industrially important species.

**MATERIALS AND METHODS**

**Bacterial strains, plasmids, and media**. The strains used in this study are listed in Table 2. The publicly available draft genome sequences of 43 *L. lactis* strains (21, 22, 26–35) were used for comparative genomics of late competence genes by employing OrthoMCL to obtain orthologous group (OG) sequences in order to construct an orthologous gene matrix (64; M. Wels et al., unpublished data). *L. lactis* strains were routinely cultivated in M17 (Tritium, Eindhoven, The Netherlands) supplemented with 1% (wt/vol) glucose (GCDM) (Tritium, Eindhoven, The Netherlands). Upon recovery after electrottransformation or natural transformation, *L. lactis* cells were cultivated in recovery medium (M17 supplemented with 1% glucose, 200 mM MgCl₂, and 20 mM CaCl₂). *Escherichia coli* TOP10 (Invitrogen, Breda, The Netherlands) was routinely cultivated in TY (Tritium, Eindhoven, The Netherlands) at 30°C with agitation. Antibiotics were added when appropriate: 5.0 μg/ml chloramphenicol, 10 μg/ml erythromycin, and 12.5 μg/ml tetracycline.

**DNA manipulations**. Plasmid DNA from *E. coli* and *L. lactis* was isolated using a Jetstar 2.0 maxiprep kit (ITK Diagnostics bv, Uithoorn, The Netherlands). Notably, phenol-chloroform extraction was performed prior to loading on the Jetstar column for plasmid isolation from *L. lactis* cultures (67). Primers were synthesized by Sigma-Aldrich (Zwijndrecht, The Netherlands). PCR was performed using KOD polymerase according to the manufacturer’s instructions (Merck Millipore, Amsterdam, The Netherlands). PCR products and DNA fragments in agarose gel were purified using the Wizard SV gel and PCR clean-up system (Promega, Leiden, The Netherlands). PCR-grade chromosomal DNA was isolated by using InstaGene Matrix (Bio-Rad, Veenendaal, The Netherlands). Ligations were performed using T4 ligase, and, when applicable, the products were transformed into either electrocompetent *E. coli* TOP10 (Invitrogen, Breda, The Netherlands) or *L. lactis* NZ9000, IL-9000, KW2, or KF147 (68).

**Plasmid and mutant construction**. To enable controlled expression of *comX* in *L. lactis*, the *comX* gene was amplified by PCR using the primer pair C1-C2 or C3-C4 and *L. lactis* KF147 or MG1363 chromosomal DNA as a template, respectively. The resulting 502-bp *comX* amplicons were digested with KpnI (introduced in primers C2 and C4) and ligated into KpnI-ScaI-digested pNZ8150 (41), yielding pNZ2600 and pNZ2601, respectively. These *comX* overexpression vectors were transformed into electrocompetent *L. lactis* KF147, NZ9000, IL-9000, and KW2 (68). The natural competence potential in *L. lactis* strains was evaluated using pIL253 (43) when possible, but because of incompatibility of antibiotic resistance markers in strain KW2, an alternative plasmid in which the erythromycin resistance (*eryR*) gene was replaced by a tetracycline resistance (tetR) gene was constructed. To this end, a 1,644-bp tetR amplicon was generated using primers C15 and C16 with pGhost8 (69) as a template and cloned as a PstI-SacI fragment into similarly digested pIL253, yielding pNZ2602. A *comEA-EC* deletion derivative of the *L. lactis* KF147 mutant was constructed using double-crossover recombination. To construct the mutagenesis fragment, the 5’ and 3’ flanking regions of the *comEA-EC* operon were amplified using chromosomal DNA of strain KF147 as a template and primer pairs C7-C8 and C11-C12, respectively. The tetracycline resistance-encoding gene tetR was amplified from pN72103 (70) using primers C9 and C10. Splicing by overlap extension PCR (SOE PCR) (71) was employed to join...
the three amplicons using the compatible sequence overhangs introduced by the primers in the individual PCRs (Table 2) and primers C7 and C12 for amplification in this PCR. The 6-kb SOE amplicon was purified from 1% agarose gels and transformed to naturally competent *L. lactis* KF147 (see Results).

The anticipated comEA-EC deletion in the resulting derivatives of *L. lactis* KF147, yielding *L. lactis* NZ6200, was confirmed by PCR using the C13 and C14 primers.

**Induction of competence in** *L. lactis*. Cells harboring pNZ6200, pNZ6201, or pNZ8150 were grown overnight in GCDM with appropriate antibiotics, followed by subculturing (1:65) in the same medium to an optical density at 600 nm (OD600) of 0.3, at which point Ultrapure nisin A (Handary, Brussels, Belgium) was added to the medium at a final concentration of 0.005, 0.01, 0.03, 0.05, 0.07, 0.1, 0.5, or 2 ng/ml. In parallel, 1 μg of plasmid DNA was added. Samples were incubated for 2 h at 30°C, after which 5 ml recovery medium was added and incubation was continued for another 2 h. Bacteria were pelleted by centrifugation at 4,000 g for 8 min, and transformants were enumerated by plating of serial dilutions on GM17 plates. KF147 transformants were subjected to PCR analysis to assess the presence of the transformed plasmid with primers PS1 and PS2, whereas the strain-specific primers SS1 and SS2 were used to confirm strain identity.

### Table 2

<table>
<thead>
<tr>
<th>Strain, plasmid, or primer</th>
<th>Relevant features or sequence</th>
<th>Reference(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Strains</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Lactococcus lactis</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>KF147</td>
<td>Plant-derived strain belonging to <em>L. lactis</em> subsp. <em>lactis</em></td>
<td>22</td>
</tr>
<tr>
<td>NZ6200</td>
<td>ΔcomEA-EC::tetR derivative of strain KF147</td>
<td>This study</td>
</tr>
<tr>
<td>IL-9000</td>
<td>Dairy-derived strain belonging to <em>L. lactis</em> subsp. <em>lactis</em> harboring nisRK integrated into its genome</td>
<td>21, 45</td>
</tr>
<tr>
<td>NZ9200</td>
<td>Dairy-derived strain belonging to <em>L. lactis</em> subsp. <em>cremoris</em> harboring nisRK integrated into its genome</td>
<td>29, 44</td>
</tr>
<tr>
<td>KW2</td>
<td>Plant-derived strain belonging to <em>L. lactis</em> subsp. <em>cremoris</em></td>
<td>28</td>
</tr>
<tr>
<td><strong>Escherichia coli TOP10</strong></td>
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<tr>
<td>pIL253</td>
<td>Cloning host</td>
<td></td>
</tr>
<tr>
<td>pNZ8040</td>
<td>Emr; high-copy-number plasmid replicative in <em>L. lactis</em></td>
<td>43</td>
</tr>
<tr>
<td>pNZ1213</td>
<td>Cmr; pNZ123 derivative containing pepN downstream of the nisin promoter</td>
<td>42</td>
</tr>
<tr>
<td>pNZ8150</td>
<td>Cmr; pNZ123 derivative with Scal site downstream of the nisin promoter for translational fusion</td>
<td>41</td>
</tr>
<tr>
<td>pNZ8200</td>
<td>Cmr; pNZ8150 derivative containing comX from <em>L. lactis</em> KF147 downstream of the nisin promoter</td>
<td>This study</td>
</tr>
<tr>
<td>pNZ8201</td>
<td>Cmr; pNZ8150 derivative containing comX from <em>L. lactis</em> MG1363 downstream of the nisin promoter</td>
<td>This study</td>
</tr>
<tr>
<td>pNZ8202</td>
<td>Tet; pIL253 derivative with eryR replacement by tetR from pGhost8</td>
<td>This study</td>
</tr>
<tr>
<td>pGhost8</td>
<td>Tet; vector with thermosensitive replicon, tetR</td>
<td>69</td>
</tr>
</tbody>
</table>

**Plasmids**

- plpL253
- pNZ8040
- pNZ1213
- pNZ8150
- pNZ8200
- pNZ8201
- pNZ8202
- pGhost8

**Primers**

- C1
- C2
- C3
- C4
- C5
- C6
- C7
- C8
- C9
- C10
- C11
- C12
- C13
- C14
- C15
- C16
- S51
- S52
- Q1
- Q2
- Q3
- Q4
- Q5
- Q6
- Q7
- Q8
- Q9
- Q10

*Underlining indicates restriction sites used in subsequent cloning procedures. Emr, erythromycin resistance; Cmr, chloramphenicol resistance; Tetr, tetracycline resistance.*
Analysis of competence gene expression. RNA was isolated from L. lactis cultures using the High Pure RNA isolation kit (Roche Diagnostics Nederland B.V., Almere, The Netherlands), including an on-column DNase treatment. Eluted RNA was again treated with DNase (1 U; Thermo Fisher Scientific, Waltham, MA, USA) for 45 min at room temperature to remove remaining DNA, followed by DNase inactivation by the addition of EDTA to a final concentration of 25 mM and then heating at 75°C for 15 min. cDNA was prepared using 10 ng total RNA and random hexamer primers in the reverse transcription reaction (Applied Biosystems, Foster City, CA, USA) according to the manufacturer’s protocol. Control RNA samples that were not reverse transcribed were included as negative controls to ensure the absence of DNA contamination. Transcripts of competence genes were quantified using 2 RNAs samplesthat werenotreversetranscribedwereincludedasnegativecontrolstoensuretheabsenceofDNA contamination. Transcripts of competence genes were quantified using 2 μl cDNA and locus-specific primers for each competence-associated target gene (primers Q1 to Q10 [Table 2]) in SYBR green-quantified PCR (Bio-Rad, Veenendaal, The Netherlands). Transcript copy numbers were calculated using a template standard curve and normalized to the housekeeping control transcript of rpoA. These RT-qPCR analyses were performed in triplicate for each sample using the Freedom EVO 100 robot system (Tecan, Männedorf, Switzerland), and amplicon identities were verified using melting curve analysis. The nonparametric Mann-Whitney U test (one-tailed) was used to determine whether gene expression levels were significantly different between uninduced and induced conditions (P < 0.05).

SUPPLEMENTAL MATERIAL
Supplemental material for this article may be found at https://doi.org/10.1128/AEM.01320-17.

SUPPLEMENTAL FILE 1, PDF file, 1.4 MB.

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