Genetics of Lactococci

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Introduction

Lactococci have been used for centuries in dairy fermentations. These Gram-positive, generally non-pathogenic, non-motile and non-sporulating bacteria are members of the Streptococcaceae family, which includes food, commensal and virulent species (Fig. 1). *L. lactis* is a relatively simple bacterium, with a 2.4 Mbp genome. Many of its functions of interest are non-redundant, which facilitates functional genetic studies of non-essential genes. Lactococci are also presumed to be devoid of virulence factors (although isolated cases did report *L. lactis* as the infectious agent in human and bovine infections; (1, 2)). The goal of this updated chapter is to confront previous and present information in different areas of lactococcal genetics, keeping in mind the relevance of findings to related bacteria, especially pathogens. Work on pathogens has long been focused on surface and secreted virulence factors, while work on lactococci has gone deeper in characterizing basic metabolic properties, nutrient uptake, and survival. Genes in basic metabolic pathways (e.g., respiration, metal homeostasis, amino acid metabolism) are now known to be essential not only for fitness, but also for virulence. Numerous lactococcus researchers who shifted their focus to pathogens have contributed to this understanding. The overall non-virulence of lactococci has also been useful in determining how metabolic and virulence factors participate in bacterial ‘everyday life’ outside the animal host. The deep knowledge of *L. lactis* physiology has led to new concepts and general findings, for example by, i) establishing the bases for dialog between Firmicutes, ii) giving concrete *in vivo* data on the biomedical or probiotic potential of recombinant and wild type lactic acid bacteria (LAB) (3, 4), and iii) establishing the existence of an inverse correlation between bacterial mRNA concentration and stability (5).

This chapter, organized in five sections, highlights major work in lactococci, including metabolic capacities, physiology, stress response, interbacterial effects, and studies leading to novel uses of lactococci for protein delivery or as probiotics. Prophages have a primordial role in fermentation processes, genetic diversity, expression and cell lysis; this important area of
research, including abortive infection mechanisms is not a focus of this review (see (6-12)). Comparative *L. lactis* genome organization analyses are considered in the following publications and references therein (13, 14).

I. **Basic features of lactococci:** A primary description of *L. lactis*, and relationship to other Streptococccaccaee are presented.

II. **Metabolic options for lactococci:** As lactococci are “industrial” bacteria, their studies focused on optimizing growth and flavor production during fermentation. Lactococci shift to a respiration metabolism when provided with an exogenous heme source. Metabolic flexibility provides a valuable prototype for lifestyle of other lactic acid bacteria and certain streptococcal pathogens, which respire when supplied with heme and also menaquinones. Nitrogen metabolism is extensively studied in *L. lactis*, as it determines the capacity to grow in dairy medium for food fermentation. Remarkably, nucleotide metabolism coordinate intracellular pools with the envelope state.

III. **Lactococcal stress responses:** Lactococci, though largely consumed through fermented foods, have been considered as non-colonizers of the animal host. Lactococci express numerous and diverse factors that allow bacteria to respond and survive in a crowded or hostile environment, and were identified by simple selective systems. Unique structures discovered on the lactococcal envelope (pellicles and pili) may extend survival in harsh environments, notably the gut, and narrow the divide between food bacteria and pathogens.

IV. **L. lactis and inter-bacterial dialogue:** Lactococci impact expression and/or development of other bacteria in their biotopes, including pathogens. Several findings may prove valuable when considering the roles of cognate functions in pathogens.

V. **Applications and genetic tools of lactococci:** Lactococci are useful bacterial vectors for biomolecule delivery. Protein expression systems were developed in lactococci for applications in biotechnology, and have been used for expression in related low GC% Gram-positive bacteria
including pathogens; remarkably, some of them have stayed in use for over 25 years. As a simple non-pathogen, L. lactis is a useful host for separating potential virulence factors from pathogenic bacteria and analyzing their roles. While medical applications have not been marketed, numerous proof of concept studies show the feasibility of using L. lactis, or other lactic acid bacteria as biomolecule delivery vectors (see (15, 16) for reviews).

I. Basic features of lactoccci

What is L. lactis?

Lactic acid bacteria (LAB) are named for their ability to produce lactic acid via a fermentation metabolism. L. lactis is a mesophilic LAB with an optimal growth temperature of ~30°C. It is the most extensively characterized LAB, which comprises a highly diverse group (Table 1, Fig. 1), including various cocci and bacilli. However, the term ‘LAB’ is misleading; although LAB generally refer to bacteria used in food fermentation, lactic acid producers also include opportunists and pathogens including Streptococci and Enterococci; of note, Enterococcus faecalis, despite its reported beneficial effects for fermentation, is now clearly emerged as an important clinical antibiotic-resistant opportunist pathogen that may lead to intestinal dysbiosis after antibiotic treatments (17-19). Among sequenced relatives, L. lactis is related to Streptococcus mutans (Fig. 1). The 2.4 Mb genome of L. lactis strain IL1403 is intermediate in size between Streptococcal pathogens like Streptococcus pneumoniae or Streptococcus pyogenes (reported as 2.1 and 1.9 Mb, respectively), and E. faecalis (3.4 Mb). Although described as “coccii”, L. lactis subpopulations may differentiate to a rod shape, possibly reflecting a change in proportions of wall and division enzymes (20).

L. lactis is classified within the Streptococcaceae family, and it appears that L. lactis and the pathogenic Streptococci may have a common origin. In addition to high genetic relatedness, DNA motifs called Chi, which are required for chromosomal integrity and
are overrepresented in L. lactis, are identical to Chi sites present in both pathogenic and food-derived Streptococcaceae, giving evidence that common core genome organization preceeded divergence of streptococcal pathogens and food bacteria (21). Relatedness between L. lactis and streptococci surpasses by far that with many other LAB (including Lactobacillus species; Fig. 1). Streptococci also produce lactic acid, and so may be considered as a branch of lactic acid bacteria. The Streptococcaceae family diverged by unknown selective pressures to generate groups of pathogens, colonizers, and food bacteria, which seem to have diversified to adapt to their preferred biotopes.

Varied lifestyle of LAB

L. lactis and LAB in general seem to have a varied lifestyle. Lactococci are isolated from plants, and are likely to be ingested by grazing animals, together with milk, in the case of calves. Co-ingestion could explain how lactococci ended up in milk. The need for several plasmid- or transposon-encoded characteristics for growth in milk (e.g., enzymes for sugar and protein metabolism, DNA restriction and abortive phage infection functions, and bacteriocins) supports the hypothesis that milk is not the original habitat of LAB.

Genome plasticity

Studies of lactococci have focused on two organisms, IL1403, a subsp. lactis strain, and MG1363, a L. lactis subsp. cremoris strain. Despite ~80% sequence identity between these genomes, they differ by a large chromosomal inversion (22). Even closely related isolates of MG1363 show considerable polymorphism, corresponding to large rearrangements (23) that might be mediated by mobile elements. Studies of artificial chromosomal rearrangements suggest that lactococci tolerate certain large genomic inversions if the origin and terminus regions are not disturbed (24).

Genome transfer and rearrangements may occur in lactococci via conjugation, transposition or insertion sequences, and phage transduction, as suggested from identification of
the required elements in different species, or by experimental systems in which natural DNA transfer occurs (see (25-28) for examples). DNA transfer occurs in streptococci by natural competence, as first discovered and then characterized in *S. pneumoniae* (29). Since then, competence among LABs was shown for *Leuconostoc mesenteroides* and *Streptococcus thermophilus* (30-33). *L. lactis* strains encode homologs of all late competence genes that are regulated by ComX; *comX* overexpression led to induction of several genes linked to late competence (e.g., *comGA, GB, GC, GD, EA, EC, FA, FC, C, dprA, coiA*), but these bacteria seemed to be defective for competence (34). Nevertheless, overexpression of ComX did lead to competence in a subgroup of lactococci (35, 36), but not in *L. lactis* model strains IL1403 and MG1363. In *L. lactis* strains KW2 and KF147, *comX* overexpression led to a transformation rate of $10^{-5}$ - $10^{-7}$ transformants/total cell number/µg of plasmid. These rates are similar to those obtained by electroporation in glycine-treated cells (37).

*L. lactis* IL1403 contains 40 reported insertion elements, 14 of which correspond to an IS element similar to IS1070 from *Leuconostoc lactis*, as well as at least four prophage elements (38, 39). *L. lactis* and *S. thermophilus* share common integrative conjugative elements, as well as highly conserved regions coding, for example, exopolysaccharide synthesis enzymes (26). Existence of functional genes that have been transferred to *L. lactis* is suggested by the presence of atypical regions, i.e., containing DNA that differs structurally from its context, or unexpected Orfs, e.g., a hemolysin-like protein. Similarly, an *S. thermophilus* isolate was shown to encode a pigment that was suspected to be transferred from pathogens, and usually lost in milk bacteria (40).

Close interactions between micro-organisms, e.g., in the gastro-intestinal or vaginal mucosa of animals, or in industrial milk fermentation processes, could lead to horizontal genetic exchange. Contact between these bacteria is also suggested by the existence of nearly identical genes, e.g., in lactococci and *S. thermophilus* (26). Close physical contact, visualized
microscopically between *S. thermophilus* and *Lactobacillus bulgaricus*, may facilitate genetic exchange (41).

**II. Metabolic options for Lactococci**

Lactococcal metabolism has been intensively studied for its industrial importance in fermentation processes, with a focus on metabolic pathways and their engineering (42). Importantly, basic metabolic functions may have far-reaching effects, and as will be described below, metabolic shifts can result in dramatic changes in *L. lactis* growth characteristics and survival.

A remarkable metabolic process in lactococci, respiration, was essentially overlooked with the focus on dairy fermentation (43). Researchers confirmed and developed a 1970 study, showing that Lactococci not only ferment sugars, but are also capable of forming an active electron transport chain to generate respiration metabolism (44-47). *L. lactis* respiration requires an external heme source, due to an incomplete biosynthetic pathway (48). The respiration process and relevance for certain pathogenic Streptococci and numerous lactobacilli is discussed.

The two energy metabolism ‘options’, fermentation and aerobic/anaerobic respiration, are presented. Pathways for nucleotide metabolism are also described, including a link established between nucleotide homeostasis and cell wall regulation.

**Energy option 1: Fermentation in *L. lactis***

Simplistically, lactococci use sugars to provide energy, and amino acids to synthesize proteins (Fig. 2) Dairy lactococci have multiple nutritional requirements for amino acids and vitamins, probably resulting from their adaptation to a life in milk. Lactose is the major sugar source in milk, and through its uptake and degradation, Lactococci generate energy in glycolysis. Casein, the major protein component in milk, is degraded to provide the major carbon source for anabolism. The flow of carbon for energy production is therefore almost separable from the flow of carbon for anabolism in these bacteria, making them ideally suited for metabolic studies.
All species belonging to the genus *Lactococcus* produce acid from glucose, fructose, mannose and N-acetylglucosamine. *L. lactis* species used for dairy fermentation undergo mainly homolactic fermentation of lactose and other sugars. *L. lactis* subsp. *lactis* strains are more versatile than subsp. *cremoris* in their use of diverse sugar sources, including maltose, ribose and trehalose (reviewed in (49)). Sugars may be transported by plasmid-encoded phosphotransferase system (PTS) systems in dairy strains, or at a slower rate by a permease in non-dairy strains (50).

Carbohydrate fermentation may be shifted from homolactic (lactate production) to mixed acid fermentation (acetic acid, formic acid, CO₂, acetoin, and ethanol produced in addition to lactate; Fig. 2). Two sets of conditions accompany this change, namely, i)- altered redox state created by increased aeration during growth; or ii)- reduced entry of sugar that is used for energy production:

1)- Oxygen appears to be involved in maintaining the NADH/NAD⁺ ratio, which itself seems to regulate the switch between homolactic and mixed acid fermentation (50); aerobic conditions result in oxidation of NADH to NAD⁺ (catalyzed by NADH oxidases), thereby reducing the NADH/NAD⁺ ratio in the cell. Lactate dehydrogenase (Ldh) is active at high NADH/NAD⁺ ratios (i.e., low oxygen), while glyceraldehyde-3-phosphate dehydrogenase (GAPDH) is inhibited. Increasing the amounts of water-forming NADH oxidase from a nisin-inducible promoter was shown to decrease the NADH/NAD⁺ ratio and to decrease *in vivo* activity of Ldh. The increased pyruvate pool was directed to acetoin and the flavor compound diacetyl.

Other types of engineering (e.g., through mutations which block specific pathways) can also result in altered flavor properties in lactococcal fermentation (reviewed in (42), with descriptions of applications in (51-54)). Note that Ldh is essential in *Streptococcus mutans*, but not in *L. lactis*, possibly because *S. mutans* lacks alternative pathways for re-oxidation of NADH that are present in *L. lactis*. Among the three Ldh-like Orfs potentially present on the *L. lactis* IL1403 genome, studies suggest that at least two genes encoding functional Ldh are potentially expressed (55).
Sugar entry may be decreased if galactose is used as an energy source or when lactose is transported by a permease (56). All sugar-carbon utilization produces pyruvate regardless of the growth conditions and the pathway; activity of enzymes using pyruvate as a substrate determines if fermentation is homolactic or mixed acid. Decreased sugar flow thus favors activity of enzymes giving rise to mixed fermentation products. One hypothesis is that when carbon fluxes are high, GAPDH is a bottleneck in glycolysis, resulting in high pools of upstream intermediates. These pools would then inhibit pyruvate formate-lyase in one of the fermentation pathways from pyruvate, resulting in homolactic fermentation under anaerobic conditions. The extent to which GAPDH is a bottleneck in lactococci was examined by constructing strains with GAPDH activities ranging from 13% to 210% of the normal activity, and measuring metabolic fluxes (57, 58). Surprisingly, GAPDH was in large excess even when carbon fluxes are high; in terms of flux control, GAPDH would thus not be the controlling factor. Moreover, the fermentation pattern remains homolactic even after four-fold reduction in GAPDH activity, which shows that GAPDH has no control on mixed acid flux either. Similar studies were carried out to determine the importance of other glycolytic enzymes on the fermentation pattern. Phosphofructokinase had no control on glycolytic flux, nor on mixed acid flux (59), despite the presence of this enzyme in excess (60). Ldh had no control on glycolytic flux either but did exert a strong negative control on mixed acid flux (61). An intriguing possibility is that the excess glycolytic enzymes are somehow shuttled to the surface, where they could play other roles in bacterial dissemination; glycolytic enzymes, like GAPDH, have been reported to be present on the surface of numerous Streptococcusaceae (62-64).

The genetic organization of enzymes involved in sugar utilization may reveal regulation at the transcriptional level. L. lactis appears to coordinate the expression of 3 genes, each involved in key but distinct steps in fermentation, pfk, pyk (encoding phosphofructokinase and, pyruvate kinase, respectively) and ldh genes by having them present in one operon (called las).
The operon organization may prevent unwanted accumulation of glycolytic intermediates. In *S. pyogenes* and *S. pneumoniae*, the *ldh* gene is not within this operon, suggesting that a common regulation of the three genes may be unique to lactococci. The *las* operon is induced in the presence of glucose via the catabolite control protein CcpA (65). Expression of the entire *las* operon was modulated to determine its importance for metabolic flux; flux was highest when expression was at its normal level, and decreased rapidly when expression was reduced or increased (59).

In *Escherichia coli*, glycolytic metabolic flux is controlled almost exclusively by the demand for ATP. In *L. lactis*, the ATP demand also controls flux in slow or non-growing cells but not in fast-growing cells (59, 66-68).

The importance of individual enzymes for system properties such as fluxes, metabolite concentrations, frequencies of infection, survival rates, etc., can be accessed by modulating the activities of the respective components. Using synthetic promoters has proven useful for obtaining accurate tuning of gene expression and optimizing metabolic fluxes (57, 58). Moreover, this technique allows numerous genes to be modulated simultaneously and at differential levels in a cell; in one example an *L. lactis* strain was constructed in which all 10 genes encoding the enzymes of the glycolytic pathways were up-regulated using synthetic promoter libraries (69).

**Energy option 2: Respiration in *L. lactis*.**

*Components of the electron transport chain:* An active respiration chain comprises three elements: i) an electron donor, supplied by NADH dehydrogenases, ii) quinones (a non-protein component of the respiration chain), which deliver electrons to the terminal oxidoreductases, and iii) terminal oxidoreductases which contain heme as an essential cofactor and use oxygen as final electron acceptor (70, 71). Although *L. lactis* undergoes fermentation in a rich medium, addition
of a heme source to aerated medium activated a respiration metabolism in *L. lactis* (44, 46). Demonstration of the existence of an electron transport chain and heme-dependent membrane NADH oxidase activity, and identification of factors implicated in respiration, all point to the major impact of this metabolic mode on bacterial growth and survival (Fig. 3; (45-47, 72-75)).

Heme-stimulated growth leads to an approximate doubling of cell biomass (in keeping with greater energy production by respiration activity), increased pH, and massive production of acetoin rather than lactic acid (46, 76). In combination with sugar-limiting conditions, as occurs during growth on maltose, the biomass yield increased further and indicated a reversal of the function of the H+-ATPase towards the direction of ATP synthesis (77). An important feature of respiration metabolism is decreased oxidation in the cell, due to respiration-dependent oxygen elimination. This leads to a more reduced intracellular environment and consequently less oxygen-induced damage. This feature, together with increased pH, may explain the extraordinary increase in bacterial survival during respiration growth (46, 72).

Like other Gram-positive bacteria that grow via a respiratory metabolism, *L. lactis* utilizes only menaquinones as electron carrier to the terminal oxidoreductase. Inactivation of *menB*, which encodes a menaquinone biosynthesis enzyme (dihydroxynaphthonic acid synthase), totally abolished both quinone production and respiration in *L. lactis* (78). However, while other respiring bacteria have several oxidoreductases that assure respiration under different conditions, *L. lactis* uses a single enzyme, the cytochrome *bd* quinol oxidase (encoded by *cydAB*) (45, 46). This kind of oxidase has a high affinity for oxygen (79). Moreover, as shown for *S. agalactiae*, cytochrome *bd* quinol oxidase contributes to both virulence and colonization ((80, 81); YY, AG, and PG unpublished data).

Respiration in *L. lactis* requires uptake of heme (iron is insufficient), possibly suggesting the presence of a heme transporter. Genetic studies suggested that the *fhu* operon mediates heme uptake (75). However, more recent studies suggest that heme might diffuse into *L. lactis*
membranes and then be recovered by cytoplasmic heme-binding factors. Heme reduction by menaquinones appears to favor membrane incorporation (82). Other factors that facilitate heme entry via diffusion are being investigated.

Based on the dramatic differences in growth, survival, and metabolite production, we had anticipated that respiration growth would induce massive changes in gene expression when compared to fermentation. Surprisingly, expression of very few genes was specifically altered by respiration, as evaluated by proteomic (21 proteins) and transcriptomic (11 transcripts) analyses of late exponential phase cells (73, 83). These studies led to the hypothesis that NADH/NAD balance, which is shifted towards NAD by respiration chain activity, is responsible for modulating enzyme activities that accompany respiration growth (Fig. 3). Significant changes in expression were observed for a single operon, as discussed below (see section on heme homeostasis) (73).

**Role of CcpA in respiration:** The regulator CcpA, controlling catabolite carbon repression, imposes a hierarchical use of carbohydrates (84). CcpA controls carbohydrate catabolism (glycolysis) and positively controls production of Ldh (65), which converts pyruvate to lactate. CcpA seems to also coordinate respiration metabolism at different levels: The first control concerns sugar uptake, as *L. lactis* does not use amino acids or glycerol as carbon source (85). CcpA also controls the transporter *fhu*, which appears to contribute to heme uptake (75). Third, CcpA regulates expression of genes encoding protein components of the respiratory chain.

In *L. lactis*, *ndh* (encoding NADH dehydrogenase; Ndh) and *cydAB* are present on distinct operons (85, 86), while in other LAB they appear to be organized in a single operon (80, 87). The latter organization suggests coordinate expression of *ndh* and *cydAB*, and may indicate that the respiratory chain is fueled by just one NADH dehydrogenase. In *L. lactis*, two putative NADH dehydrogenase genes (*noxA llmg_1735; noxB, llmg_1734*) are present in the genome and both enzymes could drive electrons to menaquinones. Though *noxA* is adjacent to *noxB*, RNAseq
analyses indicated that they are not in an operon (86). In the case of noxB, direct control by CcpA is suggested by the presence of a CcpA binding motif CRE (catabolite response element) in the promoter region. Altogether, noxB seems to be repressed by CcpA while noxA and cydAB are induced (88). Deletion of noxA abolished respiration activity on glucose growth, while the role of noxB is unknown (89). As noxB is likely induced late in growth (when CcpA is less active) it may drive electrons to menaquinone as does NoxA, but in late growth.

Interestingly, CcpA does not control respiration-induced acetoin production, as gene expression of this pathway was not affected by ccpA deletion (88). A challenging question concerns regulation of the pyruvate-acetoin pathway, which is functional late in respiration growth, as evidenced by acetoin accumulation in stationary phase cells; ~70% of the pyruvate pool is converted to acetoin (46, 73, 76) (Fig. 3). Interestingly, this pathway produces not only acetoin, but also carbon dioxide, which may contribute to lowering medium acidification in respiration growth (pH 6, compared to pH 4.5 in fermentation; both media contain 1% glucose as energy source) (PG and colleagues, submitted). The search for regulators and/or cofactors of the pyruvate-acetoin pathway, and for regulatory checkpoints of L. lactis respiration is ongoing.

Although L. lactis is only equipped for aerobic respiration, other LAB can also perform anaerobic respiration. In Lactobacillus plantarum, nitrate can be used as electron acceptor instead of oxygen. Reduction of nitrate into nitrite is performed by a quinone-nitrate reductase (NarG). Nitrate respiration is repressed by glucose suggesting potential participation of CcpA in anaerobic metabolism in this organism (89).

Heme homeostasis is a key to L. lactis respiration: Transcriptome studies of L. lactis in respiration versus fermentation conditions during exponential growth revealed strong up-regulation of components of a single operon ygfCBA encoding a putative transcriptional regulator (YgfC; also detected by proteomics (83)), a predicted permease (YgfB) and an ATPase (YgfA) (73). YgfB and YgfA are HrtB and HrtA orthologs, hrtBA for “heme-regulated
transporter”, encoding HrtBA, an ABC transporter that was found in parallel in Staphylococcus aureus (90). Its function was subsequently clarified as a dedicated heme-efflux pump, which was necessary and sufficient to maintain intracellular heme concentrations at subtoxic levels while allowing sufficient intracellular amounts for functional respiration (Fig. 4). The predicted TetR family repressor YgfC, renamed HrtR, regulates expression of hrtBA in L. lactis. HrtR shows high-affinity for heme through a non-covalent hexacoordinated interaction with heme iron (74). HrtR binds a 15-nt palindromic sequence in the hrtRBA promoter region, which is needed for repression (74). Heme binding to HrtR modifies its conformation, releasing it from its target promoter and alleviating transcriptional repression of hrtRBA (74, 91).

The use of HrtR as an intracellular heme sensor and hrtBA regulator appears to be conserved among commensal bacteria. In contrast, numerous Gram-positive pathogens use an extracellular two-component system, hssRS, to regulate hrtBA (92). These findings point to an essential role of efflux for heme homeostasis in L. lactis. In contrast, heme import mechanisms remain elusive, despite a partial role for fhu (75). Membrane-associated menaquinones (MK) were shown to favor the accumulation of reduced heme in membranes (82). An oxidative environment, provided by oxygen, prevents and reverses hemin reduction by MK and thus limits heme accumulation in membranes (82). HrtBA counteracts MK-dependent membrane retention of excess heme in membrane, suggesting direct efflux from this compartment. Moreover, both HrtBA and MK-mediated reduction have a strong impact on heme intracellular pools, detected as induction of a HrtR heme sensor. This indicates that intracellular heme acquisition is controlled at the membrane level without the need for dedicated import systems, at least at high heme concentrations (82). While pathogens may have access to blood heme during infection, the existence of heme-responsive genes in lactococci and other commensal bacteria raises questions concerning the nature of heme sources in their natural ecosystems (48).

Roles of respiratory chain components in non-respiration processes: Milk, a main industrial medium for food fermentation, lacks heme and thus cannot support L. lactis respiration
metabolism. However, respiration chain components do affect fermentation by decreasing milk redox potential of \textit{L. lactis} is one of most reducing LAB, which provides a very negative redox potential value in milk. To establish low redox potential, oxygen must be removed and oxidized compounds should be reduced. During fermentation, water-forming NADH oxidase (NoxE) eliminates oxygen, and respiration chain components (NoxAB and MenC) provide lower redox potential by reducing oxidants in milk. Redox potential of milk is known to affect the microbiota and sensorial quality of fermented dairy products. Thus, although the \textit{L. lactis} respiration chain is incomplete in milk due to the absence of a heme source, its reducing ability plays a significant role during fermentation (93).

\textit{L. lactis} is a respiration prototype: Lactococci, unlike \textit{E. coli} or \textit{Bacillus subtilis}, respire using a restricted electron transport chain, and only if heme is provided. Comparative studies reveal that certain, but not all \textit{Streptococcaceae}, and certain other Firmicutes have the capacity to respire when provided with heme and a menaquinone source (45) (Fig. 1). They include \textit{S. agalactiae} (80, 94), \textit{E. faecalis} (95, 96), and \textit{L. plantarum} (89). Since the first characterization in \textit{L. lactis}, a far wider group of LAB are revealed to adopt respiration metabolism when heme, or heme and menaquinones, are added (see (48) for review).

\textbf{Nitrogen metabolism}

In a milk medium, lactococci derive amino acids from casein, via hydrolysis by the extracellular protease PrtP, transport of the generated peptides, and further degradation by a multitude of intracellular and envelope proteases and peptidases (97, 98). Amino acids readily available in milk are used both directly as amino acid building blocks, and also as a general carbon supply for other forms of anabolism in lactococci. Extracellular proteases, plus at least 14 intracellular peptidases are of key importance for amino acid utilization. Lactococci grow poorly or die in milk fermentation conditions in mutants that are devoid of different combinations of these peptidases (99).
Dairy lactococci differ from plant lactococci in that they require several amino acids for growth. Surprisingly, strains of both origins appear to have the necessary genes for biosynthesis. Nevertheless, dairy lactococci require Ile, Leu, Val and His, and sometimes Arg, Met, Pro and/or Glu (100). These amino acid requirements in dairy strains appear to result from multiple mutations rather than deletions in the structural genes (39). This may suggest that mutations accumulated as an economic measure in strains maintained in a dairy environment. Similar results are reported for *S. thermophilus*, a dairy bacterium related to *S. pyogenes*.

**Nucleotide metabolism**

Nucleotides are not only substrates for DNA and RNA polymerases, but are also substrates or allosteric effectors for many enzymes, and furthermore constitute parts of different coenzymes. Thus, mutants in nucleotide metabolism may display numerous phenotypes. Cell nucleotide pools are also influenced by the presence of exogenous nucleobases or nucleosides in the medium. Pathways of uptake and utilization of these compounds (the so-called salvage pathways, which vary in different organisms) are key contributors to bacterial responses to changes in the medium and to increased intracellular degradation of nucleic acids as illustrated below.

As seen above for operons involved in carbon metabolism, gene organization has unique characteristics in lactococci, which may impact regulation. For example, genes encoding pyrimidine (thymine, cytosine, and uracil) biosynthesis pathway leading to the formation of UMP are organized in 5 different operons in *L. lactis*, and in a single operon in most investigated Gram-positive bacteria. Similarly, purine (adenosine, guanine, and xanthine) biosynthesis genes involved in the first ten steps leading to IMP production are located in five separate operons in *L. lactis*, and in a single operon in *B. subtilis*.

Pyrimidine biosynthesis is regulated by attenuation and antitermination by PyrR. The PyrR protein binds to PyrR ‘boxes’ (5’-UCCAGAGGCUNGCAAG-3’) present on the 5’ ends of the untranslated mRNA’s on four of the identified pyrimidine biosynthetic operons (101, 102).
Mutation of the pyrR gene results in constitutively increased levels of the pyrimidine biosynthetic enzymes (101). The active RNA binding form of PyrR in Bacillus is a dimer which is formed when UMP, UTP or PRPP (5-phosphoribosyl-1-pyrophosphate) is bound to PyrR; GTP counteracts UTP binding and thus promotes transcriptional read-through (103). The regulatory mechanisms seem to be the same in L. lactis (104). Interestingly several pyr operons in Mycobacterium smegmatis have been found to be regulated by PyrR by translational repression (105).

Purine biosynthesis in L. lactis is positively controlled by PurR, and purR mutants are purine auxotrophs. PurR binds to a Pur box consensus sequence (5’ ANNNCCGAACAAT 3’) (106-108) to activate transcription of the purC and purD operons. Data show that the PurR activating effector is PRPP. As PRPP synthetase is inhibited by ADP and purine addition leads to increased purine pools (104), purine addition may indirectly decrease gene expression (108, 109). Interestingly, riboswitch control of the PurR-regulated xpt operon is modulated by purine pools (104). In operons activated by PurR, the distance between the pur box and the –10 region was 57-58 bp. Mutational studies of pur boxes supported the importance of the central CCGAAC sequence. Interestingly, while L. lactis PurR often acts an activator, the homologous PurR in B. subtilis usually works as a repressor. In addition to the importance of PurR binding sites, the two PurR proteins also differ in their PRPP binding properties. The two types of PurR proteins are related, and the Bacillus type seems to have evolved early on from the activator type of PurR (109).

The purR regulon includes purine biosynthetic genes, but also genes involved in purine uptake and conversion into purine monophosphates. Genes involved in C1 carbon metabolism, ppGpp metabolism, phosphonate transport and pyrophosphatase activity were also identified. Interestingly, a Pur box was also identified upstream of two ribosomal RNA operons, which
could imply an interconnection between the purine pathway and translation, but this link needs to be confirmed (104, 106).

*L. lactis* can utilize exogenous nucleobases or nucleosides present in the medium or formed from intracellular degradation of nucleic acids *via* the salvage pathways. Nucleobases uracil, guanine and adenine, as well as hypoxanthine and xanthine, are taken up and converted to nucleotide monophosphates, while cytosine is not utilized by lactococci (110). Orotate may be incorporated in plasmid-bearing *L. lactis* strains that express an orotate transporter, and may be used for UMP biosynthesis, since OMP is a normal intermediate (111, 112). Thymine may be incorporated via pyrimidine phosphorylase to thymidine and further phosphorylated by thymidine kinase (*tdk*) to the corresponding monophosphate (110). Thus all common nucleobases except cytosine can be transported and incorporated into nucleotides. The ability of *L. lactis* to scavenge nucleobases or nucleosides can confer a survival advantage in stress conditions.

In *L. lactis*, all nucleosides except cytidine can be degraded to the corresponding nucleobase. However cytidine may first be deaminated to uridine and further to uracil. Pyrimidine nucleosides may also be directly converted to nucleotides since the corresponding nucleoside kinases (*udk* and *tdk*) are functional in lactococi. Two high affinity nucleoside transport systems were identified: UriP is specific for uridine and deoxyuridine, while BmpA-NupABC takes up cytidine and purine nucleosides, and the corresponding deoxyribonucleosides (113).

### III. Lactococcal stress responses

Stress response is the adaptation to homeostatic changes due to the environment, and is generally modulated by sensors and regulators that sense the state and respond to signals. Major signals for *L. lactis* include metabolites like nucleotides, acid (self-induced), oxygen, salt, and metals such as heme (natural or industrially provoked), and toxic products (*e.g.*, disinfectants). We refer
readers to recent reviews on LAB stress responses (114) and on regulatory signals in *L. lactis* (115). Here we focus on specific effectors and responses that were not previously considered, *i.e.*, nucleotides, Spx, cyclic-di-AMP, and heme and respiration stress responses.

**Regulation by nucleotide pools**

Low nucleotide pool sizes may serve as internal stress signals that provoke expression of stress response genes in *L. lactis* (116, 117). An important link between nucleotide pools and cell wall synthesis was uncovered: As part of the uridine synthesis pathway, PyrB converts l-aspartate (l-Asp) to N-carbamoyl-l-aspartate. PyrB competes with cell wall enzymes for l-Asp; thus, l-Asp consumption during rapid growth favors wall flexibility, while its accumulation in stationary phase may lead to a more rigid cell envelope (118). Additionally, high UTP pools in the human pathogen *S. pneumoniae* correlated with more capsule formation and lower biomass yield (119). These examples suggest that nucleotide pools connect nucleotide metabolism to cell envelope structure.

**The Spx family of regulators**

A hot spot for *L. lactis* mutants that compensated stress sensitivity of a *recA* mutant, thermosensitivity of a *clpP* mutant, metal toxicity, and oxidative stress, all mapped to the *spx* gene (117, 120, 121). Spx is conserved in numerous Firmicutes and was shown in *B. subtilis* to be involved in oxidative stress regulation via thioredoxin gene expression (122). Seven *spx* homologues exist in *L. lactis* MG1363, five of which contain redox-reactive C-X-X-C regions (121). Exploration of other Spx proteins revealed that SpxB, a non-redox sensitive Spx, as a key regulator of cell wall resistance to lysozyme by activating O-acetylation of peptidoglycan (123). The Spx family is intrinsic to cell responses to oxidative stress and cell wall damage conditions.

**Cyclic-di-AMP as second messenger in bacterial adaptation and potential role in heme sensitivity**
Nucleotide pools are required for DNA synthesis, but are also involved in numerous stress responses, including amino acid starvation and acid stress (e.g., via ppGpp) (124). More recently, cyclic di-AMP was identified as a second messenger in stress adaptation (125). In *L. lactis*, cyclic-di-AMP homeostasis is a balance between synthesis via adenylate cyclase enzyme CdaA (Llmg_0448) and degradation by cyclic-di-AMP phosphodiesterase enzyme (GdpP, Llmg_1816). Increasing the cyclic di-AMP pool by deletion of *gdpP* increased heat resistance and salt hypersensitivity in *L. lactis* (126). These phenotypes might be due to a change in peptidoglycan (PG) synthesis (127), as i) salt sensibility in the *gdpP* mutant is suppressed by deletion of *cdaA* or *glmM* encoding a phosphoglutaminase enzyme involved in PG integrity; ii) *cdaA* is in an operon with *glmM*; iii) GlmM interacts with CdaA to modulate its activity; iv) *L. lactis* cells spontaneously lyse when cyclic-di-AMP pools are decreased. The exact role of cyclic di-AMP in PG synthesis in *L. lactis*, though unknown, may be to modulate enzymatic activity to control cell wall plasticity. In contrast to CdaA, small molecules might interact with GdpP to modulate its activity. In *B. subtilis*, the GdpP homolog (YybT) harbors a PAS domain, which contains a heme binding site (128). Interestingly, *L. lactis* GdpP also harbors this domain and deletion of *L. lactis gdpP* gene increased sensitivity of cells to heme (128). These observations suggest a dialogue/link between heme homeostasis and cyclic-di-AMP pool enabling bacteria to adapt their physiology in response to exogenous heme.

In addition to its involvement in PG synthesis, cyclic di-AMP changed pyruvate carboxylase (*pycA*) activity in *L. lactis* (129). PycA is required for aspartate synthesis. Milk acidification is delayed in a *pycA* mutant compared to the wild type strain indicating that cyclic-di-AMP might modulate the fermentation process in some conditions.

**Strategies for survival in stress conditions**

*L. lactis* is equipped to deal with several stress conditions confronted in nature or in industrial settings, as summarized below. Salt and acid induce expression of GadB and GadC, which are
putatively involved in glutamate transport by an antiporter; glutamate transport presumably involves efflux of H⁺, thereby maintaining intracellular pH (130, 131).

Toxic products such as bile, quaternary compounds, and antibiotics may be actively pumped out of the cell by specialized transport functions. Among the numerous transport systems that shuttle metabolites in and out of the cell, some mediate drug expulsion, and consequentially, can confer drug tolerance (132). In *L. lactis*, one multi-drug pump having specificity for a wide range of amphiphilic, cationic drugs (including antibiotics, quaternary ammonium compounds, aromatic dyes, and phosphonium ions) is LmrA. The *lmrA* gene encodes an efflux pump which is responsible for export of toxic molecules such as ethidium, in exchange for H⁺ influx. LmrA (590 amino acids) is similar to the human multidrug resistance P-glycoprotein, thus raising questions about the origins of the pump. Judging from sequence analyses, an LmrA dimer would be the functional equivalent of the p-glycoprotein (LmrA is 32% identical to half of the P-glycoprotein particularly within known functional domains). Remarkably, LmrA is functional in eukaryotic cells, and is able to replace p-glycoprotein defects, thus making *L. lactis* an excellent model to study drug extrusion (132). Note that sequence comparisons predict an LmrA homologue in *S. pneumoniae* (an Orf with ~30% identity over 539 amino acids is present). The CmbT transporter (Llmg_1104), described initially as a cysteine and methionine biosynthesis transporter, was subsequently identified as a multi-drug efflux pump that effluxes a wide range of antibiotics and toxic drugs (133).

Other efflux systems protect *L. lactis* from toxic metabolites: The dedicated heme-specific HrtBA efflux pump (see above) protects cells from heme toxicity. *L. lactis* homologs of heme and protoporphyrin IX efflux pumps identified in *S. agalactiae* may suggest that backup systems are available (134).

A series of exciting reports reveal that there is more to the envelope than what was previously thought: First, pili discovered in *L. lactis* led to the idea that these bacteria might...
persist in the gut upon ingestion of fermented food products (135, 136). Second, a pellicle-like carbohydrate structure was visualized on the *L. lactis* MG1363 surface and characterized biochemically (137). A third structure, described as trapped within the peptidoglycan, corresponds to another pellicle-like carbohydrate thin outer layer surrounding *L. lactis* strain MG1363 (138).

Finally, basic cell metabolism can determine how well the bacterium copes with oxidative conditions. Respiration metabolism presents a clear advantage to lactococci in an aerobic environment for both growth and long term survival (Fig. 7; (46, 72)).

**Selections leading to improved adaptation to environmental stress situations**

Oxygen is a ubiquitous stress. If not eliminated, reactive oxygen derivatives provoke cell damage that can be lethal. One means of reducing oxygen-related damage is by removing oxygen. In fermenting cells, H_{2}O-forming NADH oxidases do eliminate oxygen during growth (139), although cells are sensitive to oxygen-related damage in stationary phase (140). Overproduction of H_{2}O-forming NADH oxidase could not only change metabolic end products (139), but also might improve survival in an oxidizing environment, due to oxygen removal. An alternative means of creating a more reducing environment is by adding glutathione, a redox peptide (141), or dithiothreitol, a reducing agent (75), to the medium. Lactococci lack catalase, which eliminates hydrogen peroxide in many aerobic bacteria. Hydrogen peroxide has been effectively removed by cloning catalase in *L. lactis* (142). Respiration metabolism in lactococci is a “natural” and efficient means of eliminating oxygen, compared to fermentation, leading to good survival in stationary phase (46, 72).

While acidification is generated by the fermentation growth process itself, and as such is self-inflicted, cell survival is handicapped by acid accumulation (143). Significantly, acidification may be more severe if cells are immobilized, as acid diffusion may be slower. This situation may provide a natural selection for strains to escape from a constrained environment,
and was shown experimentally to generate multi-stress-resistant mutants (116). Interestingly, acid resistant mutants evoked changes linking nucleotide pools (ppGpp) and cell wall alterations (116, 118). Studies using a semi-liquid medium have been used to impose a natural selection for mutants that can more readily escape a constraining environment (144). In one case, bacteria that make chains diffuse more slowly than single cells (Fig. 8). ‘De-chaining’ mutants, affected in penicillin binding protein and cell wall-synthesis enzyme PBP1A were isolated; the mutants no longer formed chains, and were able to disperse more readily in the semi-liquid medium; these mutants display greater permeability, probably due to increased cell wall breaks (144). In S. agalactiae, interruption of ponA results in reduced virulence that was linked to host antimicrobial peptide susceptibility (145, 146). In view of the identified role of PBP1A in lactococci, it is tempting to speculate that the ponA S. agalactiae mutant may also be defective in cell wall permeability and its chain-forming ability, thus impacting its in vivo localization. Use of semi-liquid medium for selections might prove effective in examining factors that are at work when bacteria are immobilized in their host, and has also proven useful in other types of selections in which surface properties are altered (137).

Transposon insertional mutagenesis was also used to select for stress-resistant strains of lactococci. Three examples are given: i) A combination of stress conditions is lethal for lactococci (and possibly for other organisms), although each condition alone may be non-lethal. Simultaneous high temperature (37°C), oxygen, and either low pH (i.e., similar to conditions in the stomach) or a recA background, give rise to mutants of which many seem to affect intracellular metabolic pools of guanosine-phosphate and phosphate in stress response (116, 117). Low intracellular levels of these metabolite pools in the mutants may constitute a starvation signal to induce a stress response. These mutant strains show better long term survival than their non-mutated parents. It thus appears that a general stress response is induced in L. lactis when intracellular guanosine-phosphate and phosphate pools are low. In accordance with this, L. lactis
showed resistance to acid and heat stress when grown in milk or a synthetic medium without
purines (147). Hydrogen peroxide is toxic to lactococci. An H$_2$O$_2$-resistant mutant was
isolated at high temperature (37°C), and although its resistance was 1000-fold greater than the
parental strain, it displayed no other stress resistance phenotypes (148). One interest of this type
of strain is its capacity to live in co-culture with strains producing mM amounts of H$_2$O$_2$, e.g.,
some Lactobacilli (149), and Streptococci (150). In the former case, use of H$_2$O$_2$-resistant
lactococci could lead to development of new fermented products; in the latter, more efficient
growth of lactococci could improve the hygiene of food products. Mutants were selected
for increased resistance to dithiothreitol (DTT), a reducing agent, at elevated temperature. By
preventing the formation of disulfide bonds, DTT disables a part of the oxidative stress response
pathway (numerous lactococcal stress-response proteins contain one or more CXXC motifs). All
18 DTT-resistant mutants mapped within a single operon, $pst$, involved in phosphate transport.
Greater oxidative stress tolerance of $pst$ was linked to its effects on copper and zinc homeostasis
(151).

Stress resistant lactococci have several potential uses. First, such strains are potentially
valuable in dairy fermentation. Their greater resistance to stress may overcome survival
variability as seen in conventional strains. The specifically acid-resistant strains may provide
resistance to extreme acid pH conditions, or may be better at maintaining a neutral internal pH.
Stress resistant strains may survive longer in fermentation, and may also be more resistant to
harsh storage conditions (like freezing and lyophilization). Second, as such strains may survive
better to the harsh environments in the gut, they may be attractive for probiotic uses. In a proof
of concept mouse inflammatory gut model, evidence was given that antioxidant superoxide
dismutase produced by wild type $L.$ lactis may improve gut integrity (4, 152, 153). Third, lactic
acid bacteria are potentially valuable candidates for production of molecules with medical or
biotechnological uses; Lactococci are non-toxic and have documented potential for expressing

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and exporting proteins or other molecules of interest, either for industrial production, or in the gut.

IV. L. lactis and inter-bacterial dialogue

Several L. lactis properties were shown to either stimulate or inhibit bacterial growth, in some cases of opportunist pathogens, in unique and interesting ways. Three examples are described:

Interbacterial cross-feeding

Via menaquinones: Other than Lactococcus, Leuconostoc, and Enterococcus species, most known LAB (pathogen or not) lack a complete menaquinone biosynthesis pathway, and require exogenous menaquinone or precursor, in addition to heme, for respiration growth (48, 89, 154, 155). Unexpectedly, menaquinone-deficient strains can overcome this deficiency via contacts with menaquinone producer species. For example, respiration of S. agalactiae, an opportunistic pathogen, is activated in the presence of L. lactis via menaquinone donation, and a heme source (78) (Fig. 5). Remarkably, E. coli expulses dihydroxynapthoate acid (DHNA), which is used by S. agalactiae to synthesize its own menaquinone (155). DHNA is coupled to an isoprenyl chain by menaquinone prenyltransferase, encoded by menA. In Lb plantarum, a menA ortholog is adjacent to the cydAB locus, suggesting that this organism can also use DHNA to synthesize menaquinones. Free menaquinones and DHNA are likely present in host ecosystems and can be used to activate metabolic pathways in commensal and pathogen bacteria as well. Thus, environmental non-pathogens might contribute to fitness and potentially virulence, of neighboring pathogens.

Via respiration: Respiration growth leads to higher pH and efficient oxygen consumption. The presence of a respiration-positive L. lactis strain (grown aerobically with heme) rescued growth and survival of a respiration-negative mutant strain (tested with cydA) (Fig. 6).

Interbacterial cross-inhibition
Via secreted products: Lactococci, like other Streptococcusae, produce toxic hydrogen peroxide under aerobic fermentation conditions, which may inhibit growth of peroxide-sensitive bacteria (72). Numerous LAB produce bacteriocins, which have widely different host spectra (see (156) for review); lysis of heterologous bacteria by bacteriocins may provide needed nutrients for the producer cells. Production of lysins may act to regulate bacterial cell wall properties in trans (157).

Via regulation of expression in trans: Studies in mixed cultures give evidence for inhibitory effects of L. lactis on S. aureus growth and/or virulence gene expression. Tests conducted in either food or mammary gland cell lines indicate that i) L. lactis reducing activity contributes to inhibition of a major virulence gene regulator (agr), and ii) L. lactis at high bacterial concentrations inhibits S. aureus internalization in a mammary epithelial cell line (158, 159). These findings open prospects for antibiotic alternatives for prevention of infection, although their applicability to food industry and in vivo situations remain to be confirmed.

V. Applications and tools of Lactococci

The use of lactococci in ‘bioprotein’ delivery, i.e., for antigen or enzyme delivery in vivo, is based on the idea that L. lactis can act as an effective non-pathogenic carrier, which can be administered orally or nasally without provoking a reaction other than that induced by the bioprotein of interest. Prior to this update, numerous reports gave encouraging results for using lactococci for: i) prophylaxis to prevent bacterial infection (160), ii) treatment of inflammatory bowel disease (3), iii) prophylaxis and/or treatment of virally-induced tumors (161), and iv) allergy prevention (162, 163). Other properties of lactococci, including the expression of pili that might mediate bacterial adhesion in situ (135, 136), and novel applications were since reported (164), which support their uses in biomolecule delivery. Despite a strong start, pharmaceutical drug development using L. lactis has not been reported, possibly due to the cost of development.
and the existence of industrially-backed alternatives. Reported success with bioactive molecules might change this picture in the future.

Numerous applications using lactococci result from the development of genetic tools. Interestingly, a thermosensitive plasmid developed in *L. lactis* over 20 years ago, pG+host, is still the gene replacement and transposition tool of choice in numerous Firmicutes including pathogenic bacteria (165, 166). pG+host is a derivative of broad host-range plasmid pWV01, and is non-replicative in *L. lactis*, streptococci, staphylococci, and in some lactobacilli at temperatures of 37.5°C or above. The thermosensitive replication protein has been retooled to produce numerous derivatives. Furthermore, pG+host is replicative in *E. coli*. The existence of an *E. coli* strain bearing a chromosomal copy of the non-temperature-sensitive *rep* gene (167) facilitates gene cloning, as well as characterization of pG+host insertions.

Another tool that has withstood time is the nisin-inducible expression system. Nisin is a bacteriocin encoded by a conjugative transposon (168-170). The promoter for the nisin biosynthesis gene, *nisA*, is regulated by *nisR* and *nisK* gene products. In the absence of nisin, or of *nisR* and *nisK* genes, promoter activity is very low. Addition of sublethal amounts of nisin results in strong induction of promoter activity, as demonstrated in innumerable applications of this system. This system has been shown to be functional in other Firmicutes (see (171) for review). Expression systems have also been developed for wider use among Firmicutes, and may be advantageous in some cases over the nisin system. Controlled expression systems have arisen from studies of pH, salt, metal, heme, chloride, or sugar- regulated promoters (see (115) for review); some systems may overcome two potentially important limitations of the nisin system: i) The requirement for either a specific host strain or a second plasmid (to provide *nisR* and *nisK* genes), and ii) effects of nisin on the membrane, which may in particular be undesirable in studies of membrane proteins.
Highly regulated promoters can also be turned around for use as specific sensors. An interesting example concerns the HrtR repressor, which regulates heme efflux (Fig. 4). This, and other heme-responsive promoters have been developed as sensitive heme sensors, with proof of activity \textit{in vivo} ((81), DL unpublished data).

Constitutive expression of promoters at fixed levels can be valuable for quantitative physiological studies or for fine-tuning of gene expression in biotechnology. \textit{L. lactis} promoters P45 and P32 are commonly used, and strong constitutive promoters have been described (172). A set of synthetic promoters that differ by the sequence and length of spacers between the consensus sequences allow a broad range of constitutive activities (57, 173). A high expression promoter based on the PTS system is further enhanced by cellobiose and shown to be active in \textit{B. subtilis} (174).

**Site specific single copy integration**

Lactococcal bacteriophages were initially studied with the goal of controlling starter culture lysis during fermentations. These phage studies led to the characterization of numerous bacterial strategies to abort phage activity. Phage studies have also been exploited to develop a site specific integration system. Integration of temperate phages makes use of a phage-specified integrase that catalyzes insertion of the phage at a specific bacterial target, which is often localized at, or near a tRNA gene. Using elements of lactococcal bacteriophage TP901-1, a site specific integrative vector was designed to obtain chromosomal single-copy integration (175). This system should allow stable insertion and expression of foreign genes, and can also be used to study expression of genes in single copy under different growth conditions. The TP901-1 integration system has also been used in human cells (176). Another integration system based on a lactococcal intron led to efficient and stable insertion of genes without the need for selection (177).

**Immune system against phages: CRISPR-Cas system**
Chromosomal manipulation was upstaged by a mechanistic breakthrough made in *S. thermophilus* on phage resistance via the CRISPR-Cas system (CRISPR: clustered regulatory interspaced short palindromic repeat, Cas: CRISPR-associated), which comprises a natural immune system against phage infections. (178). During phage DNA infection, a bacteriophage DNA fragment is integrated in a specific chromosomal CRISPR locus. During a second attack with the same bacteriophage, small RNA expressed from the CRISPR system directs adjacently expressed Cas nuclease to cleave phage DNA, which prevents new infection. CRISPR-Cas is present in various LAB including *S. thermophilus* and lactobacilli, but is uncommon in lactococci (179), although applications of the system are seen as ubiquitous (180).

**Protein export and display systems**

Protein export reporters were developed to probe membrane protein structure and identify export signals (181) (182, 183). One such reporter, the nuclease of *S. aureus*, is a stable, well characterized protein which is active when present as an amino- or carboxy-terminal fusion to other peptides, and is faithful in reporting export events and in determining membrane protein topology. It is used in *L. lactis* and other Firmicutes to follow expression of exported proteins under different environmental conditions. The major advantages of using the nuclease over previously described export reporters are that it rapidly assumes its conformation and so avoids degradation, and as few as ~300 nuclease molecules per cell can be detected in colony assays (182-185).

The use of *L. lactis* as a cell-bacterial factory has been developed to deliver proteins of interest in the extracellular medium, or on the bacterial surface. The export signal of Usp45, a secreted *L. lactis* protein, or signals from identified secreted native or heterologous proteins are routinely used. Secretion efficiency can be improved by introducing (if necessary) an overall negative charge at the N-terminal end of the mature, translocated secreted protein (184). Advantages of using LAB other than *L. lactis* concern transit time in the intestinal tract; systems
are thus being transposed to lactobacilli, such as *Lactobacillus plantarum* to improve efficacy (see (164) for review).

Anchoring of exported proteins *via* C-terminal LPXTG motifs is widely documented, particularly with respect to virulence factors, such as *S. pyogenes* M-protein (186-188). *L. lactis* encodes several anchored proteins, and at least two sortase-like proteins (189). Expression of the anchoring motif of the *S. pyogenes* M6 protein has been adopted to express recombinant proteins on the lactococcal surface (190). The AcmA autolysin is also cell wall-associated, due to the presence of a three-times repeated LysM motif (191). Anchoring *via* the AcmA binding motif was found to effectively present antigens at the lactococcal surface; a spacer between the anchoring domain and the protein of interest (a fimbrial protein) facilitates its access to target epithelial cells (192).

**Expression strains**

Lactococci are remarkable for their simple genomes, compared to complex bacterial models, such as *B. subtilis* or *E. coli*. An interesting example is HtrA, which is the only surface protease in *L. lactis*. In contrast, *B. subtilis* and *E. coli* both encode numerous exported proteases (193). An *htra* mutation leads to temperature sensitivity; at 30°C, exported proteins showed increased stability, although activity was not necessarily improved (193).

**Cell lysis systems**

Controlled cell lysis is a potentially powerful means of arresting cellular and metabolic activity; in fermentation, it may additionally result in a coordinated release of enzymes which could accelerate product maturation. The host autolysin, AcmA (194), or bacteriophage encoded lysins and holins (which allow lysin release) are good candidates for this purpose. This application is potentially useful in controlling cell growth in fermented dairy products, as well as for enzyme release (195, 196). Expression of lysin and holin by a nisin-induced promoter does indeed appear to accelerate cheese ripening (197). Lysins, derived from bacteriophage specific to several
pathogens, effectively prevented or treated infections by streptococci, bacilli, and staphylococci in mice (198-200).

The cell envelope is an important barrier protecting the cell from stress situations. Cell wall damage via autolytic enzymes can render cells more sensitive to environmental conditions. Bacteria that have undergone even partial cell wall damage are permeable to small labeled probes used in standard in situ hybridization methods, while undamaged cells are not (201). As mentioned above, the SpxB protein mediates regulation of cell wall integrity (123). Rather than affect autolysin activity, SpxB is induced by lysozyme, and or peptidoglycan (PG) digestion products leading to PG O-acetylation, which renders it lysozyme-resistant. Such changes could have profound effects on bacterial state, fitness, and resistance to autolysins, with applications for probiotic uses.

**Containment and food-grade strains**

Inactivation of the *L. lactis* thymidylate synthase thyA gene results in a requirement for thymine or thymidine (66, 202). In *L. lactis*, a thyA mutant was exploited as a means of strain containment, i.e., strains can grow in a thymine-containing environment, such as the gut, but not in more limiting environments (202).

Nonsense suppressor strains, used in *E. coli* genetics to analyze phenotypes of point mutations. Plasmids carrying the suppressor genes could suppress an otherwise lethal nonsense mutation in the cell. This property of suppressors was exploited in *L. lactis* to construct and establish a food-grade plasmid (i.e., no foreign DNA) containing the suppressor genes in a suppressible purine auxotroph; this plasmid is stable in a milk media which cannot sustain growth of a purine auxotroph (203).

**Higher antigenicity**

Studies have compared effectiveness of presentation of different antigens as bacterial cytoplasmic, surface-anchored, or secreted proteins. Cell wall associated antigens seem to induce
greater immune response than secreted or cytoplasmically expressed proteins (204, 205). Furthermore, immune response was reportedly enhanced in mutant strains with cell wall defects due to mutation of the alanine racemase (206).

Conclusions

*L. lactis* is likely the microorganism most eaten by man. It belongs to a family comprised of pathogens (e.g., *S. pneumonaie*, *S. pyogenes*), commensal microorganisms (e.g., *S. gordonii*, *S. mutans*), and food microorganisms (e.g., *S. thermophilus* and *L. lactis*). Studies of *L. lactis* reveal that differences between pathogens and non-pathogens are limited. As a bacterium that acidifies its own medium, *L. lactis* may have a high capacity for stress resistance when pre-adapted; stress-resistant mutants with constitutive stress-resistance can be selected. Its metabolic flexibility by shifting to respiration metabolism can dispense with most acid production. As a food microorganism, *L. lactis* comes into close contact with, and may cross-feed other bacteria in both the food environment and in the gut; as such, it may impact behavior of other bacteria in complex ecosystems. As a non-toxic bacterium that secretes relatively few proteins in quantity, *L. lactis* may also be an organism of choice for oral vaccine or protein delivery design and biotechnological uses (see (164, 207, 208) for reviews). As a simple organism with diverse biotopes, including plants, milk products, and the gut, *L. lactis* may be a good choice for studies on the influence of environmental stress on evolution.

Acknowledgements

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<table>
<thead>
<tr>
<th><strong>Classification</strong></th>
<th>Gram-positive, 38% G+C genome, 2.4 Mb, non-pathogenic food microorganism (referred to as a Lactic acid bacterium).</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Close neighbors</strong></td>
<td>Streptococci (food, commensal, pathogen)</td>
</tr>
<tr>
<td><strong>Optimal growth temperature</strong></td>
<td>30°C</td>
</tr>
<tr>
<td><strong>Growth medium</strong></td>
<td>Plants, milk, food silage, farm animal and human gut constituents(?)</td>
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<tr>
<td><strong>Environmental contacts</strong></td>
<td>Plants (environmental niche)</td>
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<tr>
<td></td>
<td>Farm animals</td>
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<tr>
<td></td>
<td>Milk and other foods (environmental niche may rely on plasmid-encoded factors)</td>
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<td></td>
<td>Gastrointestinal tract.</td>
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<tr>
<td><strong>Metabolism</strong></td>
<td>Fermentation.</td>
</tr>
<tr>
<td></td>
<td>Respiration in aerobic, heme-containing medium.</td>
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<td><strong>Survival</strong></td>
<td>Usually poor after fermentation growth.</td>
</tr>
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<td></td>
<td>Good long term survival after growth by respiration.</td>
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</tbody>
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Figure Legends

Fig. 1. Phylogenetic tree reveals similarities between lactococci and streptococcal pathogens. A phylogenetic tree built on 16S sequences was constructed directly on the Ribosomal Database interface (209). Branches with a bootstrap value below 60% are indicated with an asterisk. Respiration capacity (see text) is indicated by a red “R”. Rc indicates conditional respiration: for *L. lactis*, *E. faecalis*, and leuconostoc sp., aerobic respiration is activated by exogenous heme. For all lactobacillus species, and *S. agalactiae*, respiration is activated by exogenous heme and menaquinone. Opportunist pathogens are indicated in bold. This figure is based on (154).

Fig. 2. Basics of *L. lactis* fermentation. The NADH/NAD+ ratio is placed as a central determinant of carbon metabolic choice in *L. lactis* (56). 1. Sugar fermentation generates ATP, which is used for amino acid anabolism. In anaerobic conditions and rapid sugar flux essentially all sugar is converted to lactate (homolactic fermentation) from pyruvate (glycolysis). When sugar flux is slower, or in the presence of sugars other than glucose or lactose, or in aerobic growth, mixed acid fermentation may occur. The latter conditions are characterized by lower NADH/NAD⁺ ratios than those found during homolactic fermentations. Beside NADH, Glycolysis generates ATP and pyruvate from sugar degradation. 2. Pyruvate dehydrogenase (Pdh) provides extra NADH from pyruvate when oxygen is present. 3. Lactate dehydrogenase (Ldh) oxidizes NADH into NAD⁺ by conversion of pyruvate into lactate thus maintaining glycolytic activity during fermentation. 4. When oxygen is present, NADH can be oxidized by the cytoplasmic H₂O-forming NADH oxidase (NoxE), generating NAD⁺+. 5. The ATPase expels H⁺ at the expense of ATP to avoid acidification due to glycolysis. Pyruvate build-up leads to synthesis of acetate or the neutral acetoin and diacetyl (also see Fig 4). This figure is modified from (154).
Fig. 3. Basics of *L. lactis* respiration. Refer to Fig. 2 for reactions 1, 2, and 3, which are common to fermentation and respiration (numbering is the same). 1. The membrane respiration chain (RC) comprises an electron donor (putatively encoded by *noxAnoxB*; (45, 89)), menaquinones (MKs) (encoded by *men* operon genes, or provided exogenously; (80)), and a terminal electron acceptor (the cytochrome oxidase encoded by *cydAcydB*; (46, 210)). Heme (red star) must be added exogenously (red arrow) to activate cytochrome oxidase. *S. agalactiae* and lactobacilli with respiration capacity (see Fig. 1 legend) require MKs (schematic molecule with green center) and heme to activate respiration. Respiration chain activity results in H⁺ expulsion. 2. The ATPase might import H⁺, which generates ATP, but with low efficiency (47, 77). *L. lactis* lacks a complete Krebs cycle. Thus, NADH, which is needed for the respiratory chain, is produced by carbon catabolism. 3. Once phosphorylated, sugar is catabolized to pyruvate via glycolysis with production of ATP and NADH. As the respiration chain consumes NADH, Ldh activity decreases and pyruvic acid accumulates. Pyruvic acid dissociates to pyruvate and a proton, decreasing the internal pH. 4. To avoid acidification, pyruvate/pyruvic acid is converted to acetolactate via acetolactate synthase (Als), then to the neutral compound acetoin with production of CO₂. Diacetyl is produced by spontaneous oxidation of acetolactate. This pathway raises the pH and improves cell survival. Some LAB convert acetoin to 2,3 butanediol. 5. Pyruvate may also be converted to acetyl-CoA via Pyruvate dehydrogenase (Pdh), providing extra NADH and CO₂. Acetyl-CoA is further converted to acetate with production of ATP, promoting higher cell density. Acetate, acetoin, diacetyl, and 2,3-butanediol diffuse or are secreted into the medium. This figure is modified from (154).
Fig. 4. Schematic representation of *L. lactis* heme-sensing and HrtBA-mediated efflux, which regulate heme homeostasis. In *L. lactis* and numerous commensal bacteria, heme is suggested to be taken up by *fhuDBA* gene products (green ovals) and/or by diffusion through membranes (75, 82). Internalized heme binds to the HrtR repressor, which releases binding to the *hrtRBA* operon. Consequent activation of *hrtBA* results in heme efflux (74). Red squares, heme.

Fig. 5. *L. lactis* produces menaquinones that cross-feed the opportunistic pathogen *S. agalactiae*. Heme is present in the solid medium. A broad horizontal streak of an *S. agalactiae* strain (NEM316) is shown. Spots of cultures of *L. lactis* wild type (left) or *menB* that is defective for menaquinone synthesis (right) are deposited directly over the streaks. A stimulated growth zone is observed directly surrounding the WT *L. lactis* strain but not the *menB* mutant. From Mol. Microbiology (78).

Fig. 6. Respiring *L. lactis* can improve survival of non-respiring bacteria in co-culture. Differentially marked WT and *cydA* (non-respiring mutant) *L. lactis* strains were grown separately or together in co-culture. Non-respiring *cydA* grew less well, and showed poor survival when maintained in aerobic medium with heme over a 3-day period. In contrast, the respiring WT strain thrived. In contrast, the *cydA* strain fared much better when grown in co-culture with the WT strain, as determined by cell count determinations. From Mol. Microbiology (72).

Fig. 7. Respiration metabolism increases survival capacity of lactococci. When supplemented with hemin, aerobically grown lactococci can undergo respiration metabolism. As a result, cells
stored at 4°C show a markedly better survival, as compared to cells grown aerobically in the absence of hemin, or in static conditions. Improved survival was also observed when cells are maintained at 30°C. Experiment shown was performed by Karin Vido in authors’ laboratory.

**Fig. 8. Bacterial root formation in semi-liquid medium.** Bacterial chains (here, an acmA mutant of *L. lactis*; ‘parental strain’), diffuse slowly in a semi-liquid (0.035% agar) medium. Bacterial ‘de-chained’ mutants diffuse more quickly to form ‘roots’. In this experiment, all the roots corresponded to independent mutants in the same gene, *ponA*, encoding PBP1A (reproduced from Kulakauskas and coworkers, reference (144)). Note that a similar strategy of semi-liquid medium selection was used to uncover the existence of a cell-surface carbohydrate pellicle in *L. lactis*; the system is readily applied to other bacteria (137).
References


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