Lactobacilli are frequently involved in food fermentation processes, many of which are applied on an industrial scale for the production of such diverse products as sausages, sauerkraut, olives, pickled vegetables, cheese, or yoghurt. Although Lactobacillus casei may not always be the dominant organism, it influences the final texture and organoleptic properties of the product (e.g., by the production of exopolysaccharides or efficient acidification of the substrate).

Recent developments in the application of molecular biology to lactic acid bacteria have shown that it could be feasible to engineer metabolic pathways to either enhance specific metabolic fluxes or to divert metabolites for the production of different or new end products. However, this engineering requires detailed knowledge of metabolism and regulation within the targeted organism. We have chosen to investigate the complex of lactose and lactose metabolism. Lactose is initially the only carbohydrate available in milk fermentations, while galactose would, e.g., be derived from the fermentation of lactose by streptococci, which can expel the galactose moiety of lactose by using the galactose-lactose antiport mode of the lactose transporter (36).

Galactose metabolism via the Leloir pathway is a ubiquitous trait in bacterial cells. It can be used as a catabolic pathway for the degradation of galactose as an energy and carbon source while it links as an anabolic pathway the metabolism of carbohydrates, e.g., to the synthesis of lipopolysaccharides, of cell wall components, and of exopolysaccharides, for which galactosides are frequently required as building blocks. The Leloir pathway represents in the absence of an external galactose source the only means to provide this carbohydrate by biological interconversion from glucose to galactose (14).

Galactose can enter the cells by several types of transport systems. In the gram-negative bacterium Escherichia coli K-12, at least seven different systems have been characterized (18, 37), all of which release unmodified galactose into the cytoplasm. Galactose is also released as a hydrolysis product of internalized galactosides, e.g., from lactose, melibiose, or raffinose. The cleavage of β-galactosides generates β-D-galactose, which is converted into α-D-galactose by an aldose 1-epimerase (mutarotase) prior to phosphorylation (5). The free α-D-galactose moiety can be activated by an ATP-dependent galactokinase (GalK) to initiate further metabolism via the Leloir pathway. Galactose 1-phosphate, the product of this reaction, is subsequently transferred to UDP-glucose in exchange with glucose 1-phosphate by galactose 1-phosphate-uridylyltransferase (GaIT). The resulting UDP-galactose is a substrate for the reaction catalyzed by UDP-galactose 4-epimerase (GalE), resulting in UDP-galactose. Recently, gal genes have been cloned from several lactic acid bacteria. They have been found to either constitute operons of their own, as is the case for the galKTM operon of Lactobacillus helveticus (32), or they are linked to genes of lactose metabolism like those in Streptococcus thermophilus (35). Galactose metabolism in L. casei 64H is mediated by two alternative pathways: the Leloir pathway and the tagatose 1,6-bisphosphate pathway (4, 11). The latter pathway catalyzes galactose that has been transported via a galactose-specific phosphotransferase system (Gal-PTS) which releases galactose 6-P into the cytoplasm. Galactose 6-phosphate results also from the metabolism of lactose, which in L. casei 64H is exclusively transported via the Lac-PTS (8, 9).

We report here the results of our investigations of the Leloir pathway from L. casei 64H. The gal genes of L. casei 64H are the first ones to be described that are derived from a facultative heterofermentative lactobacillus that has two pathways for the degradation of galactose.

MATERIALS AND METHODS

Bacterial strains and media. L. casei subsp. casei 64H (15) was maintained at 4°C in calcium carbonate-fortified litmus milk containing 1% glucose (12). Experimental cultures were grown in lactobacillus carrying medium (LCM) supplemented with 0.5% carbohydrate (13). Plating of lactobacilli was performed on LCM media solidified with 14 g of agar per liter. E. coli strains are listed in Table 1. They were grown in LB, or plated on LB, solidified with 12 g of agar per liter. LB contains 10 g of Bacto tryptone, 10 g of yeast extract, and 5 g of NaCl per liter (26). Selection for E. coli cells transformed with the appropriate plasmids was performed by using 100 μg of ampicillin per ml or 25 μg of chloramphenicol
Where required, 50 μg of 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside per ml was used for blue-white selection in subcloning procedures. Marker tests with *E. coli* were performed on MacConkey agar-base indicator plates supplied with 1% of the carbohydrate to be tested (27). For T7 overexpression, cells were grown in phosphate-buffered minimal medium (47).

**DNA isolation and manipulation.** Chromosomal DNA was isolated from *L. casei* by the method of Neumann et al. (33). The precipitated DNA was resuspended in Tris-EDTA-RNase (10 mM Tris-HCl [pH 8.0], 0.1 mM EDTA, 20 μg of RNase H per ml) and incubated for several hours at 37°C before an additional extraction by phenol-chloroform and a final ethanol precipitation. Other DNA manipulations and plasmid preparations were performed according to standard procedures as described by Sambrook et al. (40). Restriction enzyme digests and ligations were performed as recommended by the suppliers. The sources for enzymes and other materials used for molecular biological procedures were Boehringer Mannheim (Mannheim, Germany), GIBCO BRL (Gaithersburg, Md.), New England Biolabs (Beverly, Mass.), Pharmacia Biotech Europe (Freiburg, Germany), and Qiagen (Hilden, Germany).

**Construction of pKBL1 and subclones.** Chromosomal DNA of *L. casei* 64H was digested with HindIII, EcoRI, BamHI, PstI, or XbaI. The digested DNA was size fractionated on 0.7% agarose gels, and fragments of 2 to 20 kb were isolated with the Qiaquick Gel Extraction Kit from Qiagen. Fragments were ligated to dephosphorylated pUC19 vector (53).

For T7 overexpression, the HindIII insert of pKBL1 was cloned into pHEX3 (20) in the direction with the T7p promoter (pKBL384) or against the T7p promoter (pKBL385). For overexpression of GalR, the EcoRI-HindIII fragment of pKBL1 containing the complete galR gene was cloned into pHEX3 in the direction of T7p, giving pKBL393, and in pHEX5 against T7p, giving pKBL392. For sequencing, subclones were constructed with EcoRI and the KpnI restriction sites on pKBL1. Additional subclones were obtained by creating deletion clones as described by S. Henikoff (19). Graphical representations of relevant plasmids are given in Fig. 1.

**TABLE 1. E. coli strains used in this study**

<table>
<thead>
<tr>
<th>Strain</th>
<th>Relevant genotype or phenotype</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>DH5α</td>
<td>ΔlacU169 (f80 ΔlacZΔM15) Gal⁺</td>
<td>17</td>
</tr>
<tr>
<td>HB101</td>
<td>galK2 mutant</td>
<td>6</td>
</tr>
<tr>
<td>JWL184-1</td>
<td>galT6 galP63</td>
<td>25</td>
</tr>
<tr>
<td>JWL191</td>
<td>galT6 galP63 ptsI191</td>
<td>24</td>
</tr>
<tr>
<td>S165</td>
<td>Δ(galE-TK)</td>
<td>42</td>
</tr>
<tr>
<td>JM1100</td>
<td>galK2 mgl-50 galP64 ptsG23 ptsM8 fruA10</td>
<td>18</td>
</tr>
<tr>
<td>BL21(lDE3)F' hsdS (rK⁺ mK⁺) gal [lDE3(BamHI) int::lacUV5r T7 gene 1.0]</td>
<td>45</td>
<td></td>
</tr>
</tbody>
</table>

**FIG. 1.** Schematic representation of relevant plasmids. Bent arrows indicate the direction of promoters present on the vectors. The restriction sites indicated are H, HindIII; E, EcoRI; K, KpnI; S, SalI.
Cloning of the 5' and 3' regions of the gal genes. Cloning of the 5' region was achieved by inverse PCR (49). Approximately 50 μg of chromosomal DNA from *L. casei* 64H was digested with *Kpn*I. After inactivation of the restriction endonuclease by extraction with phenol, the fragments were diluted to approximately 0.05 μg of DNA per μl and ligated overnight. The ligation mixture was inactivated by phenol treatment, precipitated with ethanol, and resuspended in 50 μl of H2O. One microliter of the ligation mixture was used as a template in a 10-μl PCR mixture by the method of Saiki et al. (39) in an Air Thermo Cycler from Idaho Technologies (Idaho Falls, Idaho) according to the protocol supplied with the cycler. A product of approximately 600 bp from a reaction with primers *PgalK* (priming site nucleotides [nt] 550 to 530 [nucleotide numbers refer throughout the paper to GenBank accession no. AF005933]) and *PgalT* (priming site nt 2170 to 2191) was cloned into vector pUC19. After the sequence was determined (2), or the DNAsis for Windows DNA and protein analysis system (Hitachi Software Engineering America, South San Francisco, Calif.).

### RESULTS

**Cloning the gal genes.** In order to isolate the Leloir pathway genes of *L. casei* 64H, the galK2 mutation of *E. coli* HB101 was used. The labeled oligonucleotides were purified with MicroSpin Sepharose HR resins from Pharmacia.

### Enzyme assays.

For measuring the enzymatic activities, extracts were prepared from *E. coli* K-12 strains carrying pKB1 or its pSU19 derivative (3). pKB275, with the complete *HindIII* insert from pKB1. For some tests, extracts were prepared from *E. coli* DH5α or *L. casei* 64H which had been induced with 0.2 or 0.5% galactose, respectively, for 3 h. Cells were harvested in mid-logarithmic phase and broken by shaking with zirconia beads in a Retsch mill MM2 (Retsch, Haan, Germany). Extracts were centrifuged at 15,000 *g* for 15 min. Protein concentrations were determined by the bichinchoninic acid method of Smith et al. (44). Galactokinase activity was determined according to the method of Sherman (43) and a protocol described by Lengeler et al. (23). Probes were taken at 30, 60, and 90 s after the start of the reaction.

Galactose 1-phosphate-uridylytransferase activity was determined by the method of Kurahashi and Anderson (22) as described in Methods of enzymatic analysis by Isselbacher (21). Concentrations of phosphoglucomutase and of glucose-6-phosphate dehydrogenase had to be increased by a factor of 10 to give detectable enzyme activities. UDP-galactose 4-epimerase activity was determined by the method of Wilson and Hogness (52).

### Nucleotide sequence accession number.

The nucleotide sequence reported in this paper has been assigned GenBank accession no. AF005933.

**TABLE 2.** GalK, GalT, and GalE activities encoded by pKB275 in the *Gal* mutant S165 of *E. coli* K-12 and in control strains.

<table>
<thead>
<tr>
<th>Strain/plasmid</th>
<th>Activity (nmol/mg/min) of:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>GalK</td>
</tr>
<tr>
<td><em>L. casei</em> 64H (Gal+)</td>
<td>83</td>
</tr>
<tr>
<td><em>E. coli</em> DH5α (Gal-)</td>
<td>24</td>
</tr>
<tr>
<td>S165/pSU19</td>
<td>0</td>
</tr>
<tr>
<td>S165/pNP5 (galKTM')</td>
<td>45</td>
</tr>
<tr>
<td>S165/pKB275 (galKETRM')</td>
<td>66</td>
</tr>
</tbody>
</table>

* ND not determined.

**FIG. 2.** Arrangement of the gal operon of *L. casei* 64H. Shaded grey arrows indicate the location and length of the genes. Black lines show fragments that were used as probes in the Northern hybridization studies. The sizes of the detected mRNAs are schematically shown as wavy lines. Restriction sites used for the construction of subclones are given.
Complementation with shotgun clones carrying fragments from chromosomal DNA of *L. casei* 64H in pUC19. One colony of approximately 5,000 transformants carrying insert sizes larger than 2 kb showed a Gal' phenotype when tested on MacConkey galactose plates. This clone contained a 5.8-kb insert of *Hind*III-digested DNA and was designated pKBL1.

**Complementation analysis with pKBL1.** In order to identify in vivo further functions encoded by pKBL1, several *E. coli* Gal' mutants were tested for complementation with either pKBL1 or the reference plasmid pNP5 carrying *galKTM* from *L. helveticus* (32). pKBL1 conferred a Gal' phenotype to strains *JWL*184-1 (*galT*6 *galP*63) and *S165* (agalETK), indicating that the insert codes for the enzymes of the Leloir pathway. Complementation was also possible in the *pts* background of *JWL*191 (*pts191 galT6 galP63*), indicating that the Gal' phenotype was not caused by the enzymes of the tagatose 1,6-bisphosphate pathway, which depend on a functional PTS. Interestingly the GalK' mutant *JM1100*, which supposedly lacks the known galactose transport systems (14), is also complemented by pKBL1 and, to a lesser extent, by pNP5. Because pNP5 and pKBL1 do not carry any transport genes in addition to the Leloir pathway genes (reference 32 and see below), strain *JM1100* must contain an as yet unidentified galactose transport system.

**Determination of enzymatic activities encoded by pKBL1.** The phenotypic characterization of pKBL1 is supported by the in vitro analysis of enzymatic activities. GalK, GalE, and GalT activities are expressed from pKBL275, the pSU19 derivative of pKBL1 (Table 2). Comparison of the activity levels of enzymes is problematic if they are expressed from different plasmids or from the chromosome and in different organisms. Host-specific factors such as codon usage or ribosome binding site specificity can further affect the results. However, the expression of active Leloir pathway enzymes is unequivocally shown by the data given.

**Cloning of the regions flanking the insert from pKBL1.** Initial sequencing data indicated the presence of an incomplete open reading frame (galM') at the 3' end of the cloned *Hind*III fragment. They also indicated that the promoter region upstream of *galK* was not cloned into pKBL1. The chromosomal regions flanking the *Hind*III fragment were amplified by inverse PCR and cloned into pUC19 as described under Materials and Methods. Using a 0.8-kb *Hind*III-*Eco*RI fragment (Fig. 2) as a probe for the 5' end, a *KpnI* fragment of 2.6 kb was detected, indicating the presence of a *KpnI* site approximately 0.6 kb upstream of the *Hind*III site. A 0.6-kb product was obtained after inverted PCR (49) and cloned into pUC18. Sequencing the ends of this fragment revealed known sequences up to the *Hind*III and *KpnI* sites respectively. A PCR with primers PgalK1 and PgalPr1 with uncut chromosomal DNA as a template gave a product of 0.6 kbp, thus proving the continuity at the level of chromosomal DNA.

A similar procedure was used in order to clone the 3' end of the *galM* gene. Southern hybridization indicated that there should be an extension of 3 kbp beyond the 3' *Hind*III site of the *gal* genes when *Eco*RI-digested chromosomal DNA was used for the inverse PCR technique. A PCR product of the expected size of 3 kbp was obtained, but various attempts to clone it into different vectors were unsuccessful. However, a smaller fragment comprising some 180 nt up to a *KpnI* site could be added by this technique to the 3' end of *galM*. Based on comparison with known sequences, there are still about 50 codons missing in the *galM* reading frame and possibly the whole *gal* gene cluster (see also below). The total established DNA sequence of the cloned region is 6,168 bp. It contains the genes *galKETR* and the incomplete gene, galM'.

**T7 overexpression of the *gal* gene products.** The enzymatic assays indicated that the *galK*, *galE*, and *galT* genes are translated into active proteins, but the expression of *galR* cannot easily be tested. In order to demonstrate its synthesis, a labeling experiment with the T7 expression system of Tabor and Richardson (46) was performed with either the complete *Hind*III fragment of pKBL1 or with the 2.3-kbp *Eco*RI-*Hind*III fragment carrying *galR* (Fig. 2). There appear three additional bands upon induction of the system when clone pKBL384 is used (Fig. 3). A prominent band of 36 kDa also present in the controls is probably a protein encoded by the vector or the host strain. The insert-dependent bands with apparent molecular masses of 52, 42, and 34 kDa correspond very well to the DNA-derived molecular masses of the encoded proteins. GalE and GalR probably run as a double band with similar masses, although it cannot be excluded that one product was not labeled in the experiment. The presence and expression of GalR are clearly demonstrated in lanes 3 and 4 of Fig. 3, since pKBL395 codes only for GalR and parts of GalT and GalM. The 34-kDa band in these lanes should therefore represent GalR and not GalE.

**Analysis of *gal* gene expression and primer extension analysis.** The hypothesis that the *gal* genes are organized as an operon was tested by Northern hybridization with probes against various parts of the *Hind*III insert from pKBL1. Two sizes of mRNAs could be detected (Fig. 4). (i) A signal at 6.5 kb appeared with all probes in RNA from galactose-induced cells. It was not present when RNA from glucose-grown cells was tested. This signal corresponds to a polycistronic mRNA, expressing *galKETRM*. (ii) A shorter mRNA of 5.6 kb was present in both samples. It could not be detected with a probe against *galK*, suggesting that this messenger carries *galETR*. The signal was weaker in RNA from glucose-grown cells than in RNA from galactose-grown cells, indicating a semiconstitut-
tive expression. Differential expression of the genes as indicated by the Northern hybridization experiment would require the activity of two promoters, \textit{galKp} and \textit{galEp}. Both transcription start sites were analyzed by primer extension and mapped to nt 334 for \textit{galKp} and to nt 1208 for \textit{galEp} (data not shown). As expected, the reactions for the \textit{galKp} extension were only positive with RNA isolated from induced cells, while the reaction for \textit{galEp} was positive with RNA from induced and uninduced cells. Potential \textit{2}10 and \textit{2}35 regions could be located at nt 1188 to 1193 and 1162 to 1167, respectively. The \textit{galEp} promoter shows only a low degree of similarity with other known promoters from lactobacilli (10, 31). This could be due to the fact that \textit{galEp} is a semiconstitutive promoter.

\textit{GalK, GalT, and GalE activities during growth on different carbohydrates.} The activities of \textit{GalK}, \textit{GalT}, and \textit{GalE} of \textit{L. casei} 64H were measured during growth on different carbohydrates and during growth in LCM in the absence of carbohydrates (Table 3). All activities are induced about 10-fold during growth with galactose, and \textit{GalK} activity is slightly repressed in the presence of glucose. As expected, growth on lactose has no inducing effect, since lactose transport and metabolism in \textit{L. casei} 64H are exclusively mediated by a lactose PTS and the tagatose 1,6-bisphosphate pathway. In order to study the induction by galactose in more detail, we measured the activity of the galactokinase after growth in LCM without additional carbohydrate or with galactose or lactose. The \textit{GalK} activity reached a maximum after approximately 3 h and then reproducibly dropped off sharply (Fig. 5).

\textbf{DISCUSSION}

We report the cloning, sequencing, and analysis of the major part of the \textit{gal} operon of \textit{L. casei} 64H. Analysis of the DNA sequence supports this hypothesis, because the deduced amino acid sequences of the five open reading frames show significant similarities to the known enzymes of the Leloir pathway from different organisms. \textit{GalK} (386 amino acids, molecular mass of 42.3 kDa) shows 57\% identical amino acids to \textit{Streptococcus mutans} and is also highly similar to other \textit{GalK} proteins. \textit{GalE} (331 amino acids, molecular mass of 36.3 kDa) shows high similarity to different \textit{GalE} proteins (e.g., 68\% identical amino acids to \textit{Streptococcus mutans} and 41\% identical amino acids to \textit{E. coli}). \textit{GalT} (486 amino acids, molecular mass of 54.1 kDa) exhibits similarities to \textit{GalT} of several gram-positive bacteria (55\% identical residues compared to \textit{GalT} of \textit{L. helveticus} and \textit{S. mutans}) but not to those of \textit{E. coli} or other gram-negative bacteria. At present, \textit{GalT} of this type seems therefore to be limited to gram-positive bacteria. However, the complementation studies and enzyme tests clearly show that this type of transferase is also active in \textit{E. coli}. The proposed start codon of \textit{galT} is a TTG and overlaps the stop codon TGA of \textit{galE}. The use of the TTG start codon is supported by the low expression of \textit{galT} in the \textit{T7} overexpression experiment. TTG as a start codon, as it is suggested here, is seldom used in \textit{E. coli} and could therefore be the reason for the poor expression of \textit{galT} in this organism. However, TTG as a start codon has been observed in \textit{L. casei} (1, 16).

\textit{GalR} (331 amino acids, molecular mass of 36.5 kDa) is a member of the LacI/GalR family of repressor proteins, because it shows up to 41\% identical amino acids with repressors of this type. It resembles, e.g., \textit{GalR} and \textit{GalS}, the repressors of the \textit{gal} regulon of \textit{E. coli} (7, 28, 38, 51). Because of its similarity to other repressors and because of its location...
within the gal gene cluster, we propose that GalR could be the repressor of the L. casei 64H gal operon, although the experiments presented provide no direct proof for this hypothesis. Galactose seems the most likely inducer, as indicated by the induction studies. Our Northern blot analysis clearly shows that galR is transcribed on polycistronic messengers which carry galKETRM or galKETR. The operon may therefore be autoregulated. Although gal operons are frequently regulated by repressor proteins of the LacI/GalR type, autoregulation has not been reported before for any of these operons. Repressors of the LacI/GalR type bind to palindromic operator sequences. Two nearly perfect inverted repeats are found in front of the operon. They are the palindromic sequences TTTTAGTA AA centered around −15 and −115 which could serve as operators.

Downstream of galR, there is a fifth partial open reading frame of 290 codons (32.1 kDa). This reading frame extends beyond the KpnI site at the end of the sequenced region. Its derived amino acid sequence shows similarities to different aldose 1-epimerases (mutarotases). Three different start codons are possible for this open reading frame, which are in frame with the proposed reading frame starting at nt 5299. The use of this start site is most probable, since there are no potential ribosome binding sites upstream of the alternative start sites. Also, the predicted additional N-terminal sequence of the longer proteins does not exhibit any similarity to other known sequences, while the protein translated from the proposed start site shares significant similarity to the so far identified GalM proteins. Like the enzymes of L. helveticus (32), E. coli (5, 29), Haemophilus influenzae (30), and S. thermophilus (35), the protein does not contain a leader peptide, indicating that it is localized within the cell. This gene is not completely cloned on pKBL1. Judging from amino acid sequence alignment, there are still about 50 codons missing to complete the reading frame of galM. It is likely that the addition of the missing codons of galM would also complete the gal operon, because the deduced length of a polycistronic messenger transcribing galKETRM corresponds to the length of the messenger detected by the Northern blot experiments.

The derived gene order for the gal region in L. casei 64H is galKETRM and represents a new arrangement of gal genes. As has been shown by the induction studies, the operon is clearly inducible by galactose in the medium. A slight repression of GalK and GalT activities is observed with glucose. In contrast to this, GalE is not repressed by glucose. This difference could result from the transcription of the operon by two different mRNAs.

The gal genes of L. casei 64H are the first ones to be cloned from a lactobacillus which can metabolize galactose by two distinct pathways. The gal genes of L. casei 64H resemble those previously cloned from other organisms. However, a feature which distinguishes the gal operon of L. casei 64H from other known gal operons is that it could be autoregulated. Autoregulation may be a requirement for the coordination of the activity of the two metabolic pathways. It will be interesting to study the regulation of the tagatose 1,6-bisphosphate pathway too, to find out whether there is any higher control mechanism in order to coordinate the two pathways.

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gal K is homologous with gal M from other organisms. Mol. Microbiol. 6:2211.


