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Metabolic engineering of the *Lactococcus lactis* diacetyl pathway

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**Summary** — Inactivation of the gene for lactate dehydrogenase caused a significant change to metabolic flux with the elimination of lactic acid as a metabolic end product and production of ethanol, formate and acetoin. Acetoin production by *Idh* defective *L. lactis* growing on sugar was comparable with that by a *diacetylactis* biovar growing on both sugar and citrate. The overexpression of biosynthetic α-acetolactate synthetase encoded by *i1vBN* genes also led to increased acetoin production in *L. lactis*. The inactivation of the *aldB* gene encoding α-acetolactate decarboxylase increased the production of α-acetolactate and diacetyl at the expense of acetoin.

*Lactococcus lactis* / genetic engineering / diacetyl / lactate dehydrogenase / α-acetolactate

**INTRODUCTION**

Lactic acid bacteria convert sugars to lactic acid via the intermediate pyruvate and certain strains, including *diacetylactis* biovars of *Lactococcus lactis* subsp *lactis*, ferment citrate that is converted to pyruvate. These latter strains metabolize the excess pyruvate by additional pathways and this can lead to formation of the flavour compound diacetyl (fig 1). The enzyme α-acetolactate synthase (ALS; EC 4.1.3.18) converts pyruvate to α-acetolactate and this unstable intermediate is converted to acetoin by α-acetolactate decarboxylase (ALD; EC 4.1.1.5; Hugenholtz, 1993) or by chemical decarboxylation (Hugenholtz, 1993). In addi-
tion, the \(\alpha\)-acetolactate can undergo oxidative decarboxylation to form diacetyl. Increased levels of acetoin and diacetyl are found under aerated conditions (Bassit et al, 1993) and this may be due in part to the increased activities of \(\alpha\)-acetolactate synthase and NADH oxidase (Bassit et al, 1993) or to decreased pyruvate-formate lyase activity (Starrenburg and Hugenholtz, 1991). The enzyme diacetyl-acetoin reductase catalyses the reduction of vicinal diketones and the subsequent reversible reduction of the monohydroxy carbonyl products. This enzyme converts diacetyl to acetoin and acetoin to 2,3-butandiol.

Diacetyl is a desirable flavour component in dairy products and there is interest in the construction of \(L\) lactis strains which generate increased yields of diacetyl. This might be achieved by a metabolic engineering approach and in general this lactococcal pathway provides a good model system with which to analyse metabolic flux and the consequences of changed levels of enzyme activity. We, in collaboration with others (W DeVos,NIZO and P Renault, Inra) have initiated genetic manipulation of the biochemical pathway involved in diacetyl production focussing on enzymes that might be expected to influence the yield of diacetyl. Here we summarize the work conducted in our laboratory. A number of genes encoding key enzymes were targeted for cloning, sequencing and manipulation. These included the \(ldh\) gene, encoding lactate dehydrogenase; the \(ilvBN\) genes encoding an \(\alpha\)-acetolactate synthase; the \(aldB\) gene encoding acetolactate decarboxylase; the \(dar\) gene encoding diacetyl-acetoin reductase; the \(oad\) gene encoding oxaloacetate decarboxylase; and the \(pfl\) gene encoding pyruvate formate lyase. The genes \(ldh\), \(aldB\) and \(ilvBN\) have been used in metabolic engineering experiments.

RESULTS AND DISCUSSION

The \(ldh\) gene encoding lactate dehydrogenase

In \(L\) lactis pyruvate is converted to lactate by the enzyme lactate dehydrogenase. It was anticipated that eliminating or decreasing the activity of this enzyme would elevate the amount of pyruvate present in the cell, mimicking the situation in citrate fermenting strains such as \(L\) lactis subsp lactis biovar diacetylactis. The increased pyruvate pool...
would then be available for metabolism by additional pathways resulting in acetoin and diacetyl production.

The N-terminal amino acid sequence and the sequence of a tryptic peptide from the *L. lactis* subsp. *cremoris* lactate dehydrogenase enzyme were used to design oligo primers. These primers were used to PCR amplify an 800-bp fragment from total genomic DNA of *L. lactis* subsp. *cremoris* MG1363. This fragment showed DNA sequence homology with the *ldh* genes of other organisms and was subsequently cloned into the Smal site of pUC13 to yield plasmid pF1514.

The 800-bp fragment was used to probe a lambda library of *L. lactis* and DNA from several plaques which gave positive signals was isolated. The DNA from one PCR positive clone was digested with a range of restriction enzymes and electrophoresed on an agarose gel. A Southern hybridization experiment showed that the 800-bp probe hybridized to a single 1.3 kbp *HindIII* band and a single 6.0 kbp *EcoRV* band. These fragments were subcloned into the *HindIII* site of pUC18 (to yield plasmid pFI515) and the *Smal* site of pUC13 (to yield pFI516) respectively. The plasmids were used to determine the DNA sequence of a 1.2 kbp region containing the entire *L. lactis* *ldh* gene (Swindell et al., 1994).

In order to inactivate the gene for lactate dehydrogenase a truncated *ldh* gene (missing both the 5' and the 3' regions) was integrated into the chromosome of MG1363 (Gasson, 1983) by a single cross-over event. This was achieved by transformation of *L. lactis* subsp. *cremoris* MG1363 with plasmid pFI658, a construct which has an *E. coli* replicon precluding replication in *L. lactis*, an erythromycin resistance gene that is selectable in *L. lactis* and an internal fragment of the *ldh* gene that provides homology for chromosomal integration. The derived strain, FI7851, possesses two dysfunctional copies of the *ldh* gene. This *ldh* mutant strain lacked lactate dehydrogenase activity and during growth on glucose produced very low levels of lactate but increased levels of acetoin (table I). This strain generated a similar amount of acetoin during sugar fermentation to that produced by *L. lactis* subsp. *lactis* biovar *diacetylactis* strains growing on both sugar and citrate.

**The aldB gene encoding α-acetolactate decarboxylase**

Inactivation of the gene encoding α-acetolactate decarboxylase should prevent the enzymatic conversion of α-acetolactate to acetoin, thereby increasing the opportunity for its oxidative decarboxylation to form diacetyl. Oligonucleotide primers for use in PCR amplification were designed from the sequence of the *aldB* gene of *L. lactis* IL1403 (JJ Godon, SD Ehrlich, personal communication). These primers were subsequently used to identify individual clones containing the *aldB* gene from a lambda library of *L.

**Table I.** The effect of *ldh* gene inactivation on *L. lactis* fermentation.

<table>
<thead>
<tr>
<th>Strain</th>
<th><em>ldh</em></th>
<th>LDH activity (U μg⁻¹)</th>
<th>Lactate (mmol/l)</th>
<th>Acetoin (mmol/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MG 1363</td>
<td>+</td>
<td>1.157 x 10⁻¹</td>
<td>53</td>
<td>4.0</td>
</tr>
<tr>
<td>FI17851</td>
<td>-</td>
<td>7.8 x 10⁻³</td>
<td>2.2</td>
<td>17.0</td>
</tr>
</tbody>
</table>
lactis subsp cremoris MG1363 using the direct PCR screening method (Griffin et al., 1993; Griffin, 1994). Positive clones were identified in this way and an insert fragment was excised and maintained as plasmid pF1937. The DNA sequence of the \textit{aldB} gene of MG1363 was determined and has been submitted to the DNA databank (accession number X82620). The \textit{aldB} DNA sequence from \textit{L. lactis} subsp \textit{cremoris} MG1363 shows 89% identity with the equivalent sequence from IL1403. The \textit{aldB} gene of \textit{L. lactis} subsp \textit{cremoris} MG1363 encodes a predicted protein sequence of 236 amino acids with a molecular mass of 26 kDa. The predicted amino acid sequence shows 97.5% identity with the equivalent sequence from IL1403; 36% identity (63% similarity) with the \textit{Bacillus brevis} sequence (Diderichsen et al., 1990); 34% identity (60% similarity) with the \textit{B. subtilis} sequence (Renna et al., 1993); 36% identity (58% similarity) with the \textit{Enterobacter aerogenes} sequence (Sone et al., 1988); 34% identity (58% similarity) with the \textit{Klebsiella pneumoniae} sequence (Blomqvist et al., 1993); and 29% identity (56% similarity) with the \textit{Coxiella burnetii} sequence (Blomqvist et al., 1993). A multiple alignment of all seven sequences shows two highly conserved regions (amino acids 34–49 and 171–188 on the MG1363 sequence) that probably represent the active site, and substrate or co-factor binding site.

PCR primers were designed to amplify a 1.4 kbp DNA fragment from the region of pF1937 immediately upstream of the start site of \textit{aldB}. The amplified fragment was cloned into the cloning vector pCRII to generate plasmid pF1777. A combined 2.8 kbp insert was subcloned into the \textit{BamHI} site of pG+host6 to form pFI805. The insert of plasmid pFI805 is, in effect, a region of lactococcal chromosomal DNA lacking the entire \textit{aldB} gene and it was used to delete the chromosomal \textit{aldB} gene of \textit{L. lactis} by a double cross-over recombination event using the temperature sensitive replicon of pG+host6. \textit{L. lactis} subsp \textit{cremoris} MG1363 was transformed with plasmid pFI805 at the permissive temperature (28°C) with selection for erythromycin. The temperature was then shifted to the restrictive temperature (37°C) at which plasmid replication was disabled. Subsequent growth on erythromycin enabled the selection of cells in which the plasmid had integrated into the chromosome (single cross-over integration). Further growth of these strains at 28°C enabled the detection of erythromycin-sensitive cells in which the second cross-over (excision) event had occurred. PCR analysis was employed to identify double cross-over recombinants containing an \textit{aldB} deletion and to confirm the chromosomal structure. One of the recombinants, strain F18076, was chosen for further study.

MG1363 was found to express a similar level of \textit{\alpha}-acetolactate decarboxylase activity to the levels expressed by other \textit{L. lactis} strains (Monnet et al., 1994; Phalip et al., 1994). \textit{\alpha}-Acetolactate decarboxylase activity was stimulated ten-fold by the addition of leucine, as observed previously (Monnet et al., 1994; Phalip et al., 1994). No \textit{\alpha}-acetolactate decarboxylase activity was detected, in the absence or presence of leucine, in F18076 cultures. Deletion of the \textit{aldB} gene had a significant effect on the pattern of fermentation end-products. In contrast to MG1363, the \textit{aldB} deletion strain F18076 produced increased quantities of \textit{\alpha}-acetolactate and diacetyl at the expense of acetoin (table II).

The \textit{ilvBN} genes encoding \textit{\alpha}-acetolactate synthase

The \textit{\alpha}-acetolactate synthase normally active in the diacetyl production pathway has a low affinity for pyruvate (Snoep et al., 1992; Monnet et al., 1994) and pyruvate is not
diverted into this pathway unless it is present in excess as for example during citrate fermentation by \textit{L. lactis} subsp. \textit{lactis} biovar \textit{diacetylactis} (Hugenholtz, 1993). The \textit{als} gene encoding this enzyme has been characterized by Marugg et al. (1994). Recently the genes \textit{ilvBN} encoding another lactococcal acetohydroxy acid synthase \textit{IlvBN} have been described (Godon et al., 1992). Enzymes homologous to \textit{IlvBN} convert pyruvate to \(\alpha\)-acetolactate during branched-chain amino acid biosynthesis (Umbarger, 1987) and transcript analysis has shown that the \textit{ilvBN} operon is not transcribed when branched-chain amino acids are supplied in the medium for example during growth in milk (Godon et al., 1993). \textit{IlvBN} consists of two subunits and has a higher affinity for pyruvate (Godon et al., 1992; K Benson, H Griffin, M Gasson, unpublished). In order to increase flux of pyruvate into the diacetyl production pathway the \textit{ilvBN} genes were expressed from the heterologous lactococcal promoter P32 (Van der Guchte, 1991; Van der Guchte et al., 1992; Van der Vossen et al., 1987). Such constitutive \textit{IlvBN} expression allowed \(\alpha\)-acetolactate synthase activity in the presence of branched-chain amino acids and increased the generation of diacetyl and acetoin in several \textit{L. lactis} strains.

A DNA fragment encoding the \textit{ilvBN} genes was constructed using a combination of gene cloning and PCR amplification. The 5' region was redesigned for expression under the control of a heterologous promoter and the fragment was inserted between the \textit{SstI} and \textit{SaiI} sites of the vector pMG36e, downstream of promoter P32 to yield plasmid pFI749. Plasmid pFI749 was used to express the \textit{ilvBN} genes in \textit{L. lactis} strains MG1363, IL1403, FI8076 and JIM4882. The latter two strains lack \(\alpha\)-acetolactate decarboxylase activity. \textit{L. lactis} FI8076 is described above and \textit{L. lactis} JIM4882 is a derivative of IL1403 from which the chromosomal \textit{ilv} and the flanking \textit{leu} and \textit{ald} operons have been deleted (JJ Godon, P Renault, unpublished results). Strains harboring plasmid pFI749 showed increased production of \(\alpha\)-acetolactate, acetoin and diacetyl (table II).

\textbf{Other genes}

The multiple alignment of amino acid sequences from databases together with

\begin{table}[h]
\centering
\begin{tabular}{|l|c|c|c|c|c|}
\hline
Strain & Subspecies & Genotype & \(\alpha\)-acetolactate & Acetoin & Diacetyl \\
& & chromosomal & plasmid & (mmol/l) & (mmol/l) & (mmol/l) \\
& & \textit{aldB} & \textit{ilvBN} & & & \\
\hline
MG1363 & \textit{cremoris} & + & - & 0 & 0.85 & 0 \\
FI8001 & \textit{cremoris} & + & + & 0 & 3.08 & 0.02 \\
FI8076 & \textit{cremoris} & - & + & 0.52 & 0.30 & 0.32 \\
FI8191 & \textit{cremoris} & + & + & 1.48 & 0.73 & 0.53 \\
IL1403 & \textit{lactis} & + & - & 0 & 1.28 & 0.05 \\
FI8046 & \textit{lactis} & + & + & 0 & 3.25 & 0.08 \\
JIM4882 & \textit{lactis} & - & - & 0 & 0 & 0 \\
FI7979 & \textit{lactis} & - & + & 0.41 & 0.21 & 0.10 \\
\hline
\end{tabular}
\caption{The effect of \textit{ilvBN} overexpression and \textit{aldB} gene inactivation on \textit{L. lactis} fermentation.}
\end{table}
redundant primer design and PCR amplification have been used to identify further genes that are relevant to diacetyl production. The oxaloacetate decarboxylase gene involved in the conversion of citrate to pyruvate and the diacetyl-acetoin reductase gene involved in the interconversion of diacetyl, acetoin, and 2,4-butanediol have been isolated. Work is currently under way to analyse these genes and exploit them in additional metabolic engineering strategies. The gene coding for pyruvate formate lyase has also been targeted for cloning and manipulation as inactivation of this gene could lead to increased availability of pyruvate for conversion to α-acetolactate.

An important aspect of diacetyl production is the use of aeration to inhibit pyruvate formate lyase and increase the pyruvate pool that is available for conversion to α-acetolactate. There is thus interest in the influence of oxygen on gene expression in L. lactis. In Gram-negative bacteria FNR-type regulators function as redox responsive control proteins in a range of different systems. These proteins are related to the cyclic AMP receptor protein (CRP or CAP) which activates catabolite-sensitive genes in response to intracellular cAMP and the CRP-FNR family contains at least 17 structurally-related transcriptional regulators. They are site-specific DNA-binding proteins possessing C-terminal helix-turn-helix motifs in their DNA-binding domains and β-roll structures in their nucleotide-binding or sensory domains and they activate or repress the transcription of target genes in response to metabolic or environmental stimuli. A multiple alignment was produced of the amino acid sequences of CRP-FNR type proteins from various bacteria and PCR primers were designed from areas of conservation based on the lactococcal codon usage table. These degenerate primers were used in a PCR reaction to amplify DNA fragments from the chromosome of MG1363 and subsequently clones containing genes for FNR-like proteins were isolated. DNA sequence analysis of two such genes has been performed. The predicted amino acid sequences of the two flp genes show approximately 60% homology (D Gostick, H Griffin, M Gasson, unpublished data).

DISCUSSION

Diacetyl is an important flavour compound in the dairy industry, providing the characteristic 'buttery' flavour of many fermented milk products. It is normally produced by a limited number of strains of lactic acid bacteria, such as L. lactis subsp. lactis biovar diacetylactis and its generation depends on citrate utilization (Hugenholtz, 1993). Here we describe a metabolic engineering approach to alter the activities of some key enzymes involved in the generation of diacetyl and demonstrate that changes in metabolic flux leading to altered patterns of fermentation end-products can be achieved.

As already described diacetyl production depends on citrate utilization leading to an increased pyruvate pool and its dissipation via several metabolic routes including conversion to the diacetyl precursor α-acetolactate. Deletion of the Idh gene encoding lactate dehydrogenase facilitated conversion of pyruvate to α-acetolactate when L. lactis is grown on sugar in the absence of citrate. The yields of α-acetolactate derivatives were equivalent to those obtained for L. lactis subsp. lactis biovar diacetylactis strains utilizing citrate as well as sugar. The fact that higher levels were not found when the pyruvate pool was significantly increased in the Idh mutants suggests that the low affinity of α-acetolactate synthase for pyruvate is a rate limiting step. This has been addressed by the constitutive expression of an anabolic α-acetolactate synthase IlvBN that is normally
involved in amino acid biosynthesis and subject to transcriptional regulation. Expression of the *ilvBN* genes under the control of a constitutive promoter in *L. lactis* led to increased conversion of pyruvate to α-acetolactate and this led to changed fermentation end products. This effect is probably due to the higher affinity of the *IlvBN* α-acetolactate synthase for pyruvate and in the longer term it is conceivable that a protein engineering approach might be used to improve the affinity of α-acetolactate synthases for pyruvate and this might further increase the entry of pyruvate into the diacetyl generation pathway. In industry the production of diacetyl depends on a natural mutant of *L. lactis* subsp *lactis* biovar *diacetylactis* that lacks α-acetolactate decarboxylase activity. The block in the enzymatic conversion of α-acetolactate into acetoin facilitates its chemical conversion into diacetyl by oxidative decarboxylation. This effect was reproduced by the use of a genetic engineering strategy in which the *aldB* gene was deleted from the chromosome of *L. lactis*.

The experiments described here clearly illustrate the potential of a metabolic engineering approach to influence the yield of a desirable metabolite. There remains considerable potential to further develop this approach. It would be interesting to stack the genetic changes that have been described individually and experiments are in progress to construct a *ldh− aldB−* double mutant and to overexpress *ilvBN* in a *ldh−* mutant. Genes for additional enzymes might also be exploited and to this end the gene for diacetyl reductase has already been cloned and that for pyruvate formate lyase is being sought. Attention might also be given to cofactor regeneration and the control of gene expression in response to oxygen. With respect to this last point it is of interest that lactococcal genes homologous to FNR-like regulators have recently been characterized.

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