Powerful methods to establish chromosomal markers in *Lactococcus lactis*: an analysis of pyrimidine salvage pathway mutants obtained by positive selections

Jan Martinussen and Karin Hammer

Using different 5-fluoropyrimidine analogues, positive selection procedures for obtaining mutants blocked in pyrimidine and purine salvage genes of *Lactococcus lactis* were established. Strains lacking the following enzyme activities due to mutations in the corresponding genes were isolated: uracil phosphoribosyltransferase (*upp*), uridine/cytidine kinase (*udk*), pyrimidine nucleoside phosphorylase (*pdp*), cytidine/deoxycytidine deaminase (*cdd*), thymidine kinase (*tdk*) and purine nucleoside phosphorylase (*pup*). Based on an analysis of the mutants obtained, the pathways by which *L. lactis* metabolizes uracil and the different pyrimidine nucleosides were verified. The substrate specificities of the different enzymes were determined. It was demonstrated that a single pyrimidine nucleoside phosphorylase accounts for the phosphorolytical cleavage of uridine, deoxyuridine and thymidine, and a single purine nucleoside phosphorylase has activity towards both the ribonucleoside and deoxyribonucleoside derivatives of adenine, guanine and hypoxanthine. No phosphorylase activity towards xanthosine appeared to be present. The selection procedures developed during this work may be employed in establishing markers on the chromosome of many related lactic acid bacteria.

**Keywords**: *Lactococcus lactis*, chromosomal markers, nucleotide metabolism, pyrimidine salvage pathway mutants, positive selections

**INTRODUCTION**

*Lactococcus lactis* has been shown to grow on a defined medium without nucleosides or nucleobases added, i.e. it contains the pathways for *de novo* synthesis of nucleotides (Jensen & Hammer, 1993). In contrast to the *de novo* pathways which, when present, seem to contain the same enzymic steps in all organisms, the salvage pathways may vary considerably in different bacteria. This provides the basis for differential usage of toxic nucleoside or nucleobase analogues to inhibit some bacteria. Furthermore, knowledge of the salvage pathways of a given bacterium is very important for the experimental use of radioactive nucleosides or nucleobases for *in vivo* labelling of RNA and DNA.

In lactococci, part of the purine salvage pathways have been studied and the *bpt* gene encoding hypoxanthine guanine phosphoribosyltransferase has been cloned and sequenced (Nilsson & Lauridsen, 1992). Recently, the nature of the pyrimidine salvage pathways of *L. lactis* have been elaborated by investigating the effect of 5-fluoropyrimidines on growth, by testing the ability of different pyrimidine derivatives to facilitate growth of a pyrimidine-requiring strain, and by assaying the presence or absence of enzyme activities in cell extracts from wild-type strains (Martinussen et al., 1994). Furthermore, the uracil phosphoribosyltransferase gene (*upp*) has been cloned and sequenced in our laboratory (Martinussen & Hammer, 1994).

In the work described in this report, we utilized the sensitivity of *L. lactis* to most 5-fluoropyrimidine deriva-
tives to define conditions for positive selection of mutants in the salvage pathways since these analogues are only toxic to the bacterial cell after being converted by the corresponding salvage enzymes. The most toxic metabolite formed inside the cell has been identified as 5-fluorodeoxyuridine monophosphate (FdUMP), which covalently binds to thymidylate synthase, the enzyme catalysing the formation of dTMP from dUMP (Heidelberger et al., 1983).

We describe positive selections for upp, udk, tdk and cdd mutants in wild-type L. lactis by modification of the procedures developed for Salmonella typhimurium by Beck and co-workers (Beck & Ingraham, 1971; Beck et al., 1972). The selections developed for pdp and pdp mutants were performed in upp or upp tdk parental strains as originally proposed for Escherichia coli by Ahmad & Pritchard (1969). Altogether, ten different conditions are described which can be used for selection of spontaneous mutants with frequencies from $3 \times 10^{-4}$ to $2 \times 10^{-7}$. The corresponding salvage genes are expected to be located on the bacterial chromosome as found for $bpt$ (Nilsson & Lauridsen, 1992) and $app$ (Martinussen & Hammer, 1994). Thus, the selections described are useful for establishment of chromosomal markers for genetic experiments in L. lactis. Previously, a positive selection procedure exploiting toxic L-alanine analogues has been described for isolating mutants in the oligopeptide transport system of L. lactis (Kunji et al., 1993). Also, selections for resistance to antibiotics such as rifampicin (Lautier et al., 1988) and streptomycin (Gasson, 1983), have been developed.

The enzymic analysis of the mutants obtained has extended the knowledge of the salvage pathways in L. lactis, particularly with respect to the number of enzymes involved. Only one pyrimidine ribonucleoside kinase ($udh$), one pyrimidine nucleoside phosphorylase ($pdp$) and one purine nucleoside phosphorylase ($pdp$) were found in this organism, and cytidine deaminase ($cdd$) was found to use both cytidine and deoxyuridine as substrates.

### METHODS

**Bacterial strains and growth procedure.** The strains used in this work are all derivatives of L. lactis subsp. cremoris MG1363 (Gasson, 1983). This laboratory strain has been cured of plasmids and prophages. The genotypes of the strains are presented in Table 1. Cultures were grown on SAG medium consisting of 1% glucose in the MOPS-based, defined medium SA containing 7 vitamins and 19 amino acids (Jensen & Hammer, 1993). When needed, 5-fluorouracil and 5-fluorouracil (5 µg ml$^{-1}$), other fluoro analogues (10 µg ml$^{-1}$), uracil (20 µg ml$^{-1}$) and uridine and cytidine (40 µg ml$^{-1}$) were added to the medium. Alternatively, the cells were grown in M17 broth (Terzaghi & Sandine, 1975) supplied with 1% glucose (GM17) or 0.5% xanthosine (XM17). For plating medium, agar was added at 15 g l$^{-1}$. In liquid media, the cells were grown without shaking. All incubations were at 30 °C in the presence of oxygen.

**Enzyme assays.** The cells were grown overnight in SAG medium, harvested, washed and resuspended in 50 mM Tris/HCl pH 7.8, 1 mM EDTA and 1 mM dithiothreitol, resulting in a 100-fold concentration of the cells. They were lysed using a French pressure cell at 20000 p.s.i. (138 MPa). Cell debris was removed by centrifugation and the supernatant was used directly as enzyme source in the assays. The uracil phosphoribosyltransferase assay was performed according to Rasmussen et al. (1986). Thymidine and uridine phosphorylases, uridine and cytidine kinases, and cytidine and deoxyuridine deaminases were assayed as described previously (Martinussen & Hammer, 1994). Thymidine kinase was assayed as described by Martinussen & Hammer (1994), with the modification that phosphonopyruvate (84 µM) and pyruvate kinase (20 units ml$^{-1}$) were included. Purine nucleoside phosphorylase activity was assayed using either the thiobarbituric acid method (deoxyadenosine and deoxyguanosine) or the xanthine oxidase coupled assay (inosine, deoxynosine and xanthosine) as described by Jensen & Nygaard (1975). Units are defined as nanomoles substrate transformed per minute at 30 °C and enzyme activities are expressed in units per mg of total protein in the crude extracts. Protein determination was performed according to the Lowry method.

**Selection procedures.** In all selections, six cultures of each strain were inoculated from different single colonies and grown overnight in GM17 medium. Cells were washed and resuspended in an equal volume of 0.9% NaCl, and 2 x 10$^7$ or 2 x 10$^8$ cells as indicated in Table 2 were spread on minimal SA plates supplied with different 5-fluoropyrimidines. The cells were not subjected to mutagenesis. The genetic background and the conditions used in the different positive selection procedures are presented in Table 2. In each selection, at least six resistant colonies from the six independently inoculated cultures were screened on plates containing the different 5-fluoropyrimidines.

### RESULTS

The use of 5-fluoropyrimidine derivatives for selection of mutants requires that the chosen component can be transported and converted to nucleotides by the cell. If the normal pathway is mutated or the uptake is impaired,
isolate was recently shown that FU is toxic to L. lactis (Hammer, 1994). In order to investigate whether it was possible to isolate upp mutants directly, wild-type cells were spread on plates containing 10 μg FU ml⁻¹. Six independently isolated resistant colonies had the same 5-fluoropyrimidine sensitivity pattern, and all had lost UPRTase activity (Table 2).

### Selection of upp mutants

The main pathway of uracil incorporation is the reaction of uracil with phosphoribosyl pyrophosphate (PRPP) to form UMP. This reaction is catalysed by uracil phosphoribosyltransferase (UPRTase) encoded by the upp gene. The toxic analogue 5-fluorouracil (FU) has been used to isolate upp mutants in other bacteria (Neuhard, 1983). It was recently shown that FU is toxic to L. lactis (Martinussen et al., 1994), indicating that uracil is being metabolized. By reverse genetics, a upp strain was constructed, and surprisingly it was only resistant to very low concentrations of FU, and for loss of uracil phosphoribosyltransferase (UPRTase) activity. As shown in Table 2, although upp mutants were obtained, the majority of clones selected on 0.3 μg FU ml⁻¹ did not lose their UPRTase activity, suggesting that mutations in other loci can confer resistance to FU at low concentrations. Resistant mutants were also obtained when FU concentrations of 1, 2, 5, 10 and 25 μg ml⁻¹ were used for the selection, but none of these had lost their UPRTase activity. The nature of mutants resistant to high FU concentrations was not determined. The frequency of FU-resistant cells decreased with increasing FU concentration (Martinussen & Hammer, 1994). As described below, tdk mutants were readily obtained. In order to isolate upp mutants in this strain, tdk cells were spread on plates containing 10 μg FU ml⁻¹. Six independently isolated resistant colonies had the same 5-fluoropyrimidine sensitivity pattern, and all had lost UPRTase activity (Table 2).

### Selection of tdk mutants

In order to isolate mutants in the tdk gene encoding thymidine kinase, wild-type cells were spread on minimal medium containing FUdR (5-fluorodeoxyuridine)

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**Table 2. Positive selections used**

<table>
<thead>
<tr>
<th>Mutant obtained</th>
<th>Corresponding enzyme</th>
<th>Genetic background</th>
<th>Selection procedure†‡</th>
<th>Frequency of resistance</th>
<th>Screened phenotype‡§</th>
<th>Frequency of analogue sensitivity (%)¶</th>
<th>Frequency in assay¶</th>
</tr>
</thead>
<tbody>
<tr>
<td>upp</td>
<td>UPRTase</td>
<td>Wild-type</td>
<td>FU₉₀</td>
<td>1·1 × 10⁻⁵</td>
<td>FU₉₀</td>
<td>81</td>
<td>2/15</td>
</tr>
<tr>
<td>tdk</td>
<td></td>
<td>FU₉₀</td>
<td>1·5 × 10⁻⁷</td>
<td></td>
<td>FU₉₀</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>upp tdk</td>
<td></td>
<td></td>
<td>4·4 × 10⁻⁸</td>
<td></td>
<td>FU₉₀</td>
<td>75</td>
<td>1/8</td>
</tr>
<tr>
<td>upp</td>
<td>Upp</td>
<td>Wild-type</td>
<td>FU₉₀ + U₅₀</td>
<td>2·3 × 10⁻⁸</td>
<td>FU₉₀</td>
<td>44</td>
<td>7/7</td>
</tr>
<tr>
<td>tdk</td>
<td>Thymidine kinase</td>
<td>Wild-type</td>
<td>U₃₀</td>
<td>2·3 × 10⁻⁷</td>
<td>U₃₀</td>
<td>100</td>
<td>5/6</td>
</tr>
<tr>
<td>tdk</td>
<td></td>
<td></td>
<td>FU₉₀ + GA₅₀</td>
<td>2·5 × 10⁻⁶</td>
<td>FU₉₀</td>
<td>33</td>
<td>4/4</td>
</tr>
<tr>
<td>pdp</td>
<td>Pyrimidine nucleoside phosphorylase</td>
<td>Wild-type</td>
<td></td>
<td>3·2 × 10⁻⁴</td>
<td>FU₉₀</td>
<td>10</td>
<td>4/4</td>
</tr>
<tr>
<td>cdd</td>
<td>Cytidine deaminase</td>
<td>Wild-type</td>
<td>FC₃₀ + U₅₀</td>
<td>3·8 × 10⁻⁴</td>
<td>FC₃₀ + U₅₀</td>
<td>92</td>
<td>6/6</td>
</tr>
</tbody>
</table>

† The number of cells plated was 2 × 10⁷ or, where indicated by an asterisk, 2 × 10⁶.
‡ Subscripts indicate the concentrations of the metabolites added to the medium in μg ml⁻¹.
§ s, sensitive; r, resistant.
¶ Frequency of mutants with the expected 5-fluoropyrimidine-resistant phenotypes as judged by screening relative to the total number of mutants.
¶ Number of mutants determined by enzyme assay among the 5-fluoropyrimidine resistant cells with the expected sensitivity pattern.
The FU moiety may then subsequently be phosphorylated by the upp gene product to give FUMP (5'-fluoro-UMP). The presence of excess uracil was expected to inhibit the formation of FUMP, thus suppressing the toxic effect of FU. The procedure was successful: five of six resistant mutants obtained lacked thymidine kinase activity.

**Selection of udk mutants**

Using the same rationale as used for selection of tdk mutants in a wild-type background, cells were plated on FUR (5-fluorouridine) (5 µg ml⁻¹) and uracil (30 µg ml⁻¹) in order to obtain mutants lacking uridine kinase activity (udk). As shown in Table 2, among eight independently isolated strains, only a single mutant in the udk gene was obtained. Using the same selection in MB112 (upp), udk mutants were readily obtained: all mutants assayed were shown to have lost uridine kinase activity. The presence of cytidine kinase activity in *L. lactis* has previously been demonstrated (Martinussen et al., 1994). No such activity could be detected in any of the udk mutants, indicating that uridine and cytidine are phosphorylated to UMP and CMP by the same enzyme (Table 3).

**Selection of cdd mutants**

5-Fluorodeoxycytidine (FCdR) is deaminated to FUdR by a deaminase encoded by the cdd gene. In order to obtain cdd mutants, wild-type cells were plated in the presence of FCdR at concentrations of 0.3, 0.5 and 1 µg ml⁻¹. The frequency of resistant mutants decreased with increasing FCdR concentration, and from 0.5 to 1 µg FCdR ml⁻¹ the frequency dropped by a factor of approximately ten.

### Table 3. Enzymic analysis of mutants and substrate specificity of the individual enzymes

<table>
<thead>
<tr>
<th>Mutant strain</th>
<th>Gene</th>
<th>Encoded enzyme</th>
<th>Substrate</th>
<th>Specific activity*</th>
<th>Wild-type</th>
<th>Mutant</th>
</tr>
</thead>
<tbody>
<tr>
<td>MB112</td>
<td>upp</td>
<td>Uracil phosphoribosyltransferase</td>
<td>Uracil</td>
<td>6.0</td>
<td>&lt; 0.1</td>
<td></td>
</tr>
<tr>
<td>MB105</td>
<td>udk</td>
<td>Uridine kinase</td>
<td>Uridine</td>
<td>4.2</td>
<td>&lt; 1.2</td>
<td></td>
</tr>
<tr>
<td>MB103</td>
<td>tdk</td>
<td>Thymidine kinase</td>
<td>Thymidine</td>
<td>0.5</td>
<td>&lt; 0.1</td>
<td></td>
</tr>
<tr>
<td>MB139</td>
<td>pdp</td>
<td>Uridine phosphorylase</td>
<td>Uridine</td>
<td>51</td>
<td>&lt; 1</td>
<td></td>
</tr>
<tr>
<td>MB130</td>
<td>pdp</td>
<td>Thymidine phosphorylase</td>
<td>Thymidine</td>
<td>40</td>
<td>&lt; 1</td>
<td></td>
</tr>
<tr>
<td>MB109</td>
<td>cdd</td>
<td>Cytidine deaminase</td>
<td>Cytidine</td>
<td>17</td>
<td>&lt; 1</td>
<td></td>
</tr>
<tr>
<td>MB140</td>
<td>pup</td>
<td>Purine nucleoside phosphorylase</td>
<td>Deoxyguanosine</td>
<td>76</td>
<td>&lt; 9</td>
<td></td>
</tr>
</tbody>
</table>

* Specific activity is given in units (mg protein)⁻¹ at 30 °C in crude extract.
Pyrimidine metabolism in *Lactococcus lactis*

### Table 4. 5-Fluoropyrimidine sensitivity patterns of the isolated mutants

<table>
<thead>
<tr>
<th>Strain</th>
<th>Genotype</th>
<th>5-Fluoropyrimidine added*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>FU₀₃</td>
</tr>
<tr>
<td>MG1363</td>
<td>Wild-type</td>
<td></td>
</tr>
<tr>
<td>MB112</td>
<td>upp</td>
<td>+</td>
</tr>
<tr>
<td>MB105</td>
<td>udk</td>
<td></td>
</tr>
<tr>
<td>MB128</td>
<td>upp, udk</td>
<td>+</td>
</tr>
<tr>
<td>MB103</td>
<td>tdk</td>
<td></td>
</tr>
<tr>
<td>MB116</td>
<td>tdk, upp</td>
<td></td>
</tr>
<tr>
<td>MB139</td>
<td>tdk, upp, pdp</td>
<td></td>
</tr>
<tr>
<td>MB140</td>
<td>tdk, upp, pdp</td>
<td></td>
</tr>
<tr>
<td>MB117</td>
<td>upp, pdp</td>
<td></td>
</tr>
<tr>
<td>MB109</td>
<td>cdd</td>
<td></td>
</tr>
</tbody>
</table>

*5-Fluoropyrimidine derivatives were added at 10 μg ml⁻¹, except where indicated by subscripts (in μg ml⁻¹). Uracil was added at 30 μg ml⁻¹, and AR and GdR at 0.5 mg ml⁻¹.*

The different FCdR-resistant mutants were screened for sensitivity for FUdR, since a *cdd* mutant can be expected to be sensitive to this analogue (Fig. 1). Independently isolated, FUdR-sensitive mutants from each selection were assayed for deoxyxanthine deaminase activity, and only among strains selected on 0.5 μg FCdR ml⁻¹ were mutants found which had lost this enzyme activity (Table 2). These mutants were shown to be sensitive to FCdR concentrations of 1 μg ml⁻¹ and above, suggesting the existence of an alternative pathway for the metabolism of deoxyxanthine. The *cdd* mutants isolated had lost both cytidine and deoxyxanthine deaminase activity, indicating that both reactions are catalysed by the same enzyme (Table 3).

### Selection of pdp mutants

Lactococci contain gene(s) encoding enzyme activities responsible for the phosphorolytic cleavage of uridine, deoxyuridine and thymidine (Martinussen *et al.*, 1994). Positive selection procedures for obtaining mutants in these enzymes cannot be established in the wild-type background, but require a primary mutation, often a *upp* background (Neuhard, 1983). The normal pathway of uracil incorporation involves a phosphoribosylation to UMP catalysed by the *upp* gene product. If this pathway is blocked, the cells are resistant to FU. Certain conditions may allow FU to follow alternative pathways involving a reaction between FU and pentose-1-phosphate (P-1-P) catalysed by a pyrimidine nucleoside phosphorylase, resulting in the formation of the corresponding 5-fluoropyrimidine nucleoside that subsequently will lead to death of the cell (Fig. 1). A prerequisite for this reaction is the presence of a sufficient concentration of P-1-P inside the cell, resulting in the formation of the corresponding P-1-P and pyrimidine base. Therefore, it is reasonable to believe that in the *upp* strain, mutants in the alternative uracil pathway may be obtained by plating on FU in the presence of a purine nucleoside. As shown below, FU-resistant lactococcal strains are sensitized by addition of purine nucleosides. Alternatively, mutants may be obtained by plating on high concentrations of FU, since the *upp* strain is sensitive to FU concentrations above 0.3 μg ml⁻¹, whereas a *upp tdk* strain is resistant, indicating that dR-1-P is present to some extent in the cell, even in the absence of exogenous deoxyribonucleosides.

The *upp* strain MB112 was suitable for the isolation of a mutant unable to cleave thymidine, either by using FU in high concentration, or using FU in the presence of deoxyguanosine (Table 2). The presence of deoxyguanosine increases the toxicity of FU in the *upp* background, thus reducing background growth, making the selection more efficient. As expected, two classes of mutants were obtained, namely pyrimidine nucleoside phosphorylase mutants and strains lacking thymidine kinase activity. The two mutants can easily be distinguished by phenotype screening, since only the *upp tdk* mutants are resistant to FUdR (Table 4). The thymidine phosphorylase mutants obtained in these selections were shown to have lost both thymidine and uridine phosphorylase activities, suggesting that both enzyme activities are properties of the same polypeptide.

Strain MB116 (*upp tdk*) is resistant to FU in high concentrations, but becomes sensitive to FU in the presence of adenosine as R-1-P donor. Thus, mutants lacking uridine phosphorylase activity were selected in strain MB116 (*upp tdk*) on plates containing FU in the presence of adenosine. Mutations in at least three different genes will result in resistance. In addition to uridine-phosphorylase-deficient strains, mutants affected in urid-
dine kinase (udk) and adenosine phosphorylase were isolated (Table 2). udk mutants are characterized by their resistance towards FUR, whereas uridine and adenosine phosphorylase mutants have the same 5-fluoropyrimidine sensitivity pattern. By screening the enzyme activities, the identity of the FUR-sensitive mutants was established, showing that mutants in both nucleoside phosphorylases were obtained (Table 2). Furthermore, both the uridine and thymidine phosphorylase activities were shown to be absent in the strains exhibiting uridine phosphorylase deficiency, as would be expected if both activities were properties of the same enzyme (see above). These results demonstrate that *L. lactis* has a general pyrimidine nucleoside phosphorylase. The corresponding gene is designated pdp.

In other bacteria, mutants lacking uridine phosphorylase activity have been obtained in a upp background by selection on FU in the presence of adenosine, which serves as a R-1-P donor (Neuhard, 1983). Since the upp mutation in MB112 only confers resistance to FU at a very low concentration, mutations in both the U → UR → UMP and U → UdR → dUMP pathways are required if plated on high FU in the presence of adenosine (Fig. 1). This demands either double mutations, or if one enzyme is shared by both pathways, a single mutation in the corresponding gene. As indicated above, the lactococcal pyrimidine nucleoside phosphorylase is responsible for the phosphorolysis of both UR and UdR. Therefore, mutants in MB112 (upp) selected for resistance to FU in the presence of adenosine are most likely pdp mutants. Indeed, virtually all resistant mutants were shown to have lost both uridine and thymidine phosphorylase activity (Table 2).

**Selection of pup mutants**

Purine nucleoside phosphorylase catalyses the phosphorolytic cleavage of a purine nucleoside into a purine base and P-1-P. A general purine nucleoside phosphorylase using adenosine, inosine, guanosine and the corresponding deoxyribosyl derivatives has been described for *E. coli* (Jensen & Nygaard, 1975), whereas in *Bacillus subtilis* two different enzymes (adenosine phosphorylase and inosine-guanosine phosphorylase) are required for the same reactions (Jensen, 1978). As described above, mutant lactococci unable to cleave adenosine were readily isolated. The same result was obtained using inosine (Table 2). No purine nucleoside phosphorylase activity towards deoxyadenosine, deoxyguanosine, inosine or deoxyinosine as substrate could be measured in crude extracts from the different mutants isolated (Table 3). This demonstrates that only one purine nucleoside phosphorylase responsible for the cleavage of adenosine, guanosine, inosine and the corresponding deoxyribosyl derivatives is present in *L. lactis* when grown in rich glucose medium (GM17). An additional purine nucleoside phosphorylase activity responsible for the cleavage of xanthosine can be detected in *E. coli* only when grown in the presence of this substrate (Hammer-Jespersen et al., 1980). Therefore, an extract from lactococcal cells grown on rich medium supplied with xanthosine instead of glucose (XM17) was assayed for the presence of xanthosine phosphorylase activity. No activity could be measured (data not shown), suggesting that such an enzyme is absent in *L. lactis*.

**DISCUSSION**

**Mutant selections**

FU and the 5-fluoropyrimidine nucleosides FUR, FUdR and FCDR were used for the selections. When designing the selective conditions, it is important to consider the concentrations of the toxic analogues in relation to the number of cells plated. For the experiments given in Table 2, this is particularly important for the selections using FCdR and FU in wild-type strains. Selection for resistance to FU is the classical selection to obtain mutants in the upp gene. Previous work has shown that upp mutants in *L. lactis* are sensitive to concentrations above 0.3 μg ml⁻¹. Even when this concentration of FU is used for the selection, only 2 of 15 mutants isolated have lost the corresponding enzyme activity, thus being upp. The nature of the other mutants is unknown, but strains impaired in uracil uptake or overproducers of uracil may also be resistant to FU. In *B. subtilis*, an uracil-overproducing strain (pyR) is viable in the presence of 1 μg FU ml⁻¹, whereas upp mutations result in resistance up to 3 μg ml⁻¹ (Martinussen et al., 1995).

Another important consideration when using 5-fluoropyrimidine nucleosides is that FU is formed by phosphorolytic cleavage by cells harbouring pyrimidine nucleoside phosphorylase (pdp) activity (Fig. 1), and that FU may be excreted by the layer of wild-type cells on the selective medium. In the case of FUR and FUdR, concentrations of 5 and 10 μg ml⁻¹ were used for the selections. Therefore substantial amounts of FU would be present. Since FU is toxic to *L. lactis* in amounts lower than 0.3 μg ml⁻¹, a surplus of uracil (30 μg ml⁻¹) was included in the selections for udk and tdk mutants in a wild-type strain. The presence of this amount of uracil does obviate the effect of any FU formed as seen from the sensitivity of the isolated mutants to FUR and FUdR when uracil is omitted (Table 4).

A third principle was also used: the addition of a purine nucleoside together with FU results in a sensitive phenotype in a upp or a upp tdk strain. Formation of internal P-1-P by the action of purine nucleoside phosphorylase facilitates the synthesis of FUR and FUdR by the reverse action of pyrimidine nucleoside phosphorylase (pdp) (Fig. 1). The upp mutant of *L. lactis*, in contrast to that of *E. coli*, contains adequate internal dR-1-P to synthesize FUdR when the FU concentration is raised above 0.3 μg ml⁻¹ (Martinussen & Hammer, 1994). The addition of deoxyguanosine reduces background growth, thus facilitating the selection of pdp or tdk mutants (Table 2).

**Salvage pathways**

This paper describes the conditions for positive selection of mutations in six different genes encoding enzymes...

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involved in the nucleoside salvage pathways in *L. lactis.* Based on analogy with the corresponding genes in *E. coli* and/or *B. subtilis* we have designated the genes *upp*, *tdk*, *pdp*, *cdd*, and *pap.* From the enzyme analysis of the corresponding mutants (Table 3) we conclude that the genes encode uracil phosphoribosyltransferase (*upp*), uridine-cytidine kinase (*udp*), thymidine kinase (*tdk*), thymidine-uridine phosphorylase (*pdp*), cytidine-deoxycytidine deaminase (*cdd*), and deoxyadenosine-deoxyguanosine-inosine phosphorylase (*pap*). The substrate specificity of the enzymes was defined by the mutant phenotypes. Cytidine deaminase was found to use both cytidine and deoxycytidine as substrate, as found for other organisms (Neuhard, 1983). A single pyrimidine ribonucleoside kinase using both cytidine and uridine as substrate is found in *L. lactis,* as in *E. coli* (Valentin-Hansen, 1978) and *B. subtilis* (Orengo & Kobayashi, 1978). Only one pyrimidine phosphorylase enzyme with activity towards both thymidine and uridine was found in *L. lactis.* It has been shown for *B. subtilis* that both thymidine and uridine phosphorylase activity resides within one protein encoded by *pdp* (Rumyantseva et al., 1979). Consequently, the corresponding gene was named *pdp.* In *E. coli,* the two activities are mediated by separate enzymes: thymidine phosphorylase is encoded by *deoA* (Schwartz, 1978), whereas uridine phosphorylase is encoded by *udp* (Leer et al., 1977). Just as for the *deoD* gene product in *E. coli* (Jensen & Nygaard, 1975), a single purine phosphorylase enzyme with activity towards adenine, guanine and hypoxanthine nucleosides was found in *L. lactis.* In *B. subtilis,* two purine nucleoside phosphorylases are found: adenosine phosphorylase specific for adenine-containing nucleosides, and inosine/guanosine phosphorylase (Jensen, 1978). In all organisms where purine nucleoside phosphorylases occur, the enzyme does not discriminate between ribose or deoxyribose in the substrate nucleoside (Parks & Agarwal, 1972). This was also found for the enzyme encoded by *pap* in *L. lactis.*

The established salvage pathways for nucleosides and nucleobases in *L. lactis* are given in Fig. 1. They are different from those in *B. subtilis* with respect to *papA* and *papI,* and from those in *E. coli* with respect to *deoA* and *udp.* As previously described, *L. lactis,* like *B. subtilis,* does not contain a *cod* gene encoding cytosine deaminase (Martinussen et al., 1994). In contrast, *E. coli* does have the capacity to utilize this substrate (Neuhard, 1983).

The sensitivity of the *cdd* mutants of *L. lactis* for FCdR concentrations above 0.5 µg ml⁻¹ indicates that *L. lactis,* as *B. subtilis* but in contrast to *E. coli,* may contain a deoxycytidine kinase. Although no deoxycytidine kinase activity could be measured, such an activity may be present, accounting for the toxic effect of FCdR in high concentration despite the presence of a *cdd* mutation. This was further indicated by testing the sensitivity of the *upp* *tdk* mutants to FCdR. FCdR is deaminated by cytidine deaminase to create FUDR, and strain MB116 (*upp tdk*) is resistant to 10 µg FUDR ml⁻¹ (Table 4). It is, however, sensitive to 1 µg FCdR ml⁻¹, but like *cdd* mutants, this strain is resistant to 0.5 µg FCdR ml⁻¹. Further experi-

ments are required to determine unequivocally the nature of the sensitivity of *cdd* mutants to FCdR.

**REFERENCES**


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