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Abstract

Yeast strains transformed with at least one copy of a gene coding for lactic dehydrogenase (LDH) and further modified for the production of lactic acid with high yield and productivities, are described.

2 Claims, 12 Drawing Sheets
OTHER PUBLICATIONS


* cited by examiner
Figure 1

- Ethanol
- Acetaldehyde
- Acetate
- Acetyl-CoA
- CO₂
- TCA
- Glucose
- Pyruvate
- Lactic acid
- Prom-LDH

1. Glucose to Pyruvate
2. Pyruvate to Ethanol
3. Ethanol to Acetaldehyde
4. Acetaldehyde to Acetate
5. Acetate to Acetyl-CoA
6. CO₂ from Acetyl-CoA
7. NAD⁺ from NADH
8. Lactic acid from NADH
9. Prom-LDH to Lactic acid

Reactions:
- 2NAD⁺ + Glucose → 2NADH + Pyruvate
- Pyruvate → Ethanol + NAD⁺ + NADH
- Ethanol → Acetaldehyde + NAD⁺ + NADH
- Acetaldehyde → Acetate + CO₂ + NAD⁺
- Acetate → Acetyl-CoA + NAD⁺
- CO₂ + O₂ → H₂O + CO₂
- Lactic acid → NAD⁺ + NADH
- Glucose + NAD⁺ → Pyruvate + NADH
Figure 2

![Diagram of a circular molecule with various labels and annotations.](image-url)
Figure 3B

LDH-A

Hind III (1676)
Cla I (1683)
Sal I (1691)
Ava I (1697)

Xba I (4595)

Apa I (3670)

Apa I (2424)

pKSEXH/16
4595 bp
Figure 6

- ADH promoter
- LDH
- Amp^r
- URA3
- plAT-ADH 8936 bp
- HindIII (31)
- PstI (11)
- TEF promoter
- APT1, G418r
- EcoRI (1551)
- PstI (2077)
- SmaI (2308)
- EcoRI (2327)
- ARS1
- CEN4
Fig. 8.1

Kluyveromyces lactis ΔPDC

Fig. 8.2

Kluyveromyces lactis ΔPDC - buffered
**Fig. 3A**

**Kluyveromyces lactis Δ PDC, Δ PDH**

**Fig. 3B**

**Glucose Vs Lactate**

\[ y = -0.8546x + 91.989 \]

\[ R^2 = 0.997 \]
Figure 10

Torulaspora delbrueckii

Time (hours)

cells/ml

4.10E+07
2.10E+07
1.00E+06

LA, ET, pH
0 50 100 150
Figure 11

Zygosaccharomyces bailii

![Graph showing growth and pH over time for Zygosaccharomyces bailii](image)
YEAST STRAINS FOR THE PRODUCTION OF LACTIC ACID TRANSFORMED WITH A GENE CODING FOR LACTIC ACID DEHYDROGENASE

The invention refers to yeast strains transformed with at least one copy of a gene coding for lactic dehydrogenase (LDH) and further modified for the production of lactic acid with high yield and productivity.

BACKGROUND OF THE INVENTION

The applications of lactic acid and its derivatives encompass many fields of industrial activities (i.e., chemistry, cosmetic, and pharmacy), as well as important aspects of food manufacture and use. Furthermore, today there is growing interest in the production of such an organic acid to be used directly for the synthesis of biodegradable polymer materials.

Lactic acid may be produced by chemical synthesis or by fermentation of carbohydrates using microorganisms. The latter method is now commercially preferred because microorganisms have been developed that produce exclusively one isomer, as opposed to the racemic mixture generated by chemical synthesis. The most important industrial microorganisms, such as species of the genera Lactobacillus, Bacillus, and Rhizopus, produce L(+)-lactic acid. Production by fermentation of D(-)-lactic acid or mixtures of L(+) and D(-)-lactic acid are also known.

During a typical lactic acid fermentation, there is an inhibitory effect caused by lactic acid produced on the metabolic activities of the producing microorganism. Besides the presence of lactic acid, lowering the pH value also inhibits cell growth and metabolic activity. As a result, the extent of lactic acid production is greatly reduced.

Therefore, the addition of Ca(OH)₂, CaCO₃, NaOH, or NH₄OH to neutralise the lactic acid and thereby prevent the pH decrease is a conventional operation in industrial processes to counteract the negative effects of free lactic acid accumulation.

These processes allow the production of lactate(s) by maintaining the pH at a constant value of the range of about 5 to 7; this is well above the pKₐ of lactic acid, 3.86.

Major disadvantages are connected to the neutralisation of lactic acid during the fermentation. Mainly, additional operations are required to regenerate free lactic acid from its salt and to dispose of or recycle the neutralising cation; this is an expensive process. All the extra operations and expense could be eliminated if free lactic acid could be accumulated by microorganisms growing at low pH values, thus minimising the production of lactate(s).

It has been proposed the use of recombinant yeasts expressing the lactate dehydrogenase gene so as to shift the glycolytic flux towards the production of lactic acid.

However, because of the high production of ethanol, the yield in the production of lactic acid for both the processes described was not considered to be competitive with that obtainable by the use of lactic bacteria.

In the past decade, "non conventional yeasts" other than S. cerevisiae have gained considerable industrial interest as host for the expression of heterologous proteins. Examples are the methanol-utilising yeasts such as Hansenula polymorpha and Pichia Pastoris, the lactose-utilizing yeasts such as Kluyveromyces lactis. In addition to enabling the use of a wider range of substrates as carbon and energy sources, other arguments have been put forward to the industrial use of "non conventional yeasts". Generally speaking, biomass and product-yield are less affected, in some of these yeasts, by extreme conditions of the cellular environment. High sugar-tolerant (i.e., 50–80% w/v glucose medium; Torulaspora syn. Zygosaccharomyces-delerbreuckii, Zygosaccharomyces rouxii and Zygosaccharomyces bailii; Ok T and Hashinaga F., Journal of General & Applied Microbiology 43(1): 39–47, 1997) and acid- and lactose-tolerant (Zygosaccharomyces rouxii and Zygosaccharomyces bailii; Houtsma P C, et al., Journal of Food Protection 59(12), 1300–1304, 1996) "non conventional yeasts" are available. As already underlined the cost of down stream processing could be strongly reduced if the fermentation process is carried out under one or more of the above mentioned "extreme conditions".

SUMMARY OF THE INVENTION

According to a first embodiment, this invention provides yeast strains lacking ethanol production ability or having a reduced ethanol production ability and transformed with at least one copy of a gene coding for lactic dehydrogenase (LDH) functionally linked to promoter sequences allowing the expression of said gene in yeasts.

More particularly, this invention provides yeasts strains having a reduced pyruvate dehydrogenase activity and a reduced pyruvate decarboxylase activity and transformed with at least one copy of a gene coding for lactic dehydrogenase (LDH) functionally linked to promoter sequences allowing the expression of said gene in yeasts.

According to another embodiment, this invention provides yeast strains of Kluyveromyces, Torulaspora and Zygosaccharomyces species, transformed with at least one copy of a gene coding for lactic dehydrogenase (LDH) functionally linked to promoter sequences allowing the expression of the gene in said yeasts.

According to a further embodiment, the invention also provides yeast cells transformed with a heterologous LDH gene and overexpressing a lactate transporter.

Other embodiments are the expression vectors comprising a DNA sequence coding for a lactic dehydrogenase functionally linked to a yeast promoter sequence and a process for the preparation of D,L-, D- or L-lactic acid by culturing the above described metabolically engineered yeast strains in a fermentation medium containing a carbon source and recovering lactic acid from the fermentation medium.

Furthermore, the invention provides processes for improving the productivity (g/l/hr), production (g/l) and yield (g/g) on the carbon source of lactic acid by culturing said yeast strains in a manipulated fermentation medium and recovering lactic acid from the fermentation medium.

DESCRIPTION OF THE INVENTION

It has been found that production of lactic acid can be obtained by metabolically modified yeasts belonging to the
genera Kluyveromyces, Saccharomyces, Torulaspora and Zygosaccharomyces. More particularly, it has been found that very high yields in the production of lactic acid are obtained by engineered yeast strains so as to replace at least the ethanolic fermentation by lactic fermentation.

Even higher yields (>80% g/g) in the production of lactic acid may be obtained by engineered yeast strains so as to replace both the ethanolic fermentation and the use of pyruvate by the mitochondria by lactic fermentation.

To this purpose, the invention also provides transformed yeast cells having an increased LDH activity, for instance as a consequence of an increased LDH copy number per cell or of the use of stronger promoters controlling LDH expression.

An increased LDH copy number per cell means at least one copy of a nucleic acid sequence encoding for lactic dehydrogenase protein, preferably at least two copies, more preferably four copies or, even more preferably, at least 10–50 copies of said nucleic acid sequence.

In order to have the highest production of lactic acid, yeast cells transformed according to the invention preferably overexpress a lactate transporter. This can be obtained by transforming yeast cells with one or more copies of a gene required for lactate transport.

The strains according to the invention can be obtained by several methods, for instance by genetic engineering techniques aiming at the expression of a lactate dehydrogenase activity, and by inactivating or suppressing enzymatic activities involved in the production of ethanol, e.g. pyruvate decarboxylase and alcohol dehydrogenase activities, and by inactivating or suppressing enzymatic activities involved in the utilisation of pyruvate by the mitochondria.

Since pyruvate decarboxylase catalyses the first step in the alcohol pathway, yeast strains without or having a substantially reduced pyruvate decarboxylase (PDC) activity and expressing a heterologous lactate dehydrogenase gene are preferred.

Further, since pyruvate dehydrogenase catalyses the first step in the utilization of pyruvate by the mitochondria, yeast strains having no or a substantially reduced pyruvate dehydrogenase (PDH) activity and expressing a heterologous lactate dehydrogenase gene are also preferred.

Since lactate is excreted in the medium via a lactate transporter, cells producing lactic acid and overexpressing the lactate transporter are also preferred.

The expression of a LDH gene in yeast strains allows the production of lactic acid at acid pH values so that the free acid is directly obtained and the cumbersome conversion and recovery of lactate salts are minimized. In this invention, the pH of the fermentation medium may initially be higher than 4.5, but will decrease to a pH of 4.5 or less, preferably to a pH of 3 or less at the termination of the fermentation.

Any kind of yeast strain may be used according to the invention, but Kluyveromyces, Saccharomyces, Torulaspora and Zygosaccharomyces species are preferred because these strains can grow and/or metabolize at very low pH, especially in the range of pH 4.5 or less; genetic engineering methods for these strains are well-developed; and these strains are widely accepted for use in food-related applications.

Good yields of lactic acid can moreover be obtained by Kluyveromyces, Torulaspora and Zygosaccharomyces strains transformed with a gene coding for lactic dehydrogenase having a "wild-type" pyruvate decarboxylase and/or a pyruvate dehydrogenase activity.

The term "reduced pyruvate decarboxylase activity" means either a decreased concentration of enzyme in the cell or reduced or no specific catalytic activity of the enzyme.

The term "reduced pyruvate dehydrogenase activity" means either a decreased concentration of enzyme in the cell or reduced or no specific catalytic activity of the enzyme.

According to the invention, it is preferred the use of strains wherein the ethanol production is or approaches zero but a reduced production for instance at least 60% lower, preferably at least 80% lower and even more preferably at least 90% lower than the normal of wild-type strains is acceptable.

According to the invention, it is preferred the use of strains wherein the pyruvate decarboxylase and/or pyruvate dehydrogenase activities are or approach zero but a reduced activity for instance at least 60% lower, preferably at least 80% lower and even more preferably at least 90% lower than the normal of wild-type strains is acceptable.

An example of K. lactis having no PDC activity has been disclosed in Mol. Microbiol. 19 (1), 27–36, 1996.

Examples of Saccharomyces strains having a reduced PDC activity are available from ATCC under Acc. No. 200027 and 200028. A further example of a Saccharomyces strain having a reduced PDC activity as a consequence of the deletion of the regulatory PDC2 gene has been described in Holmann S (1993) (Mol Gen Genet 241:657–666).

An example of a Saccharomyces strain having no PDC activity has been described in Flikweert M. T. et al. (Yeast, 12:247–257, 1996). In S. cerevisiae reduction of the PDC activity can be obtained either by deletion of the structural genes (PDC1, PDC5, PDC6) or deletion of the regulatory gene (PDC2).

An example of Kluyveromyces strain having no PDC activity has been described in Zeeman et al. (Genes involved in pyruvate metabolism in K. lactis; Yeast, vol 13 Special Issue April 1997, Eighteenth International Conference on Yeast Genetics and Molecular Biology, p143).

An example of Saccharomyces strain having no PDH activity has been described in Pronk J T. et al. (Microbiology. 140 (Pt 3):601–10, 1994).

PDC genes are highly conserved among different yeast genera (Bianchi et al., Molecular Microbiology, 19(1): 27–36, 1996; Lu P. et al., Applied & Environmental Microbiology, 64(1):94–7, 1998). Therefore it can be easily anticipated that following classical molecular approaches, as reported by Lu P. et al. (supra), it is possible to identify, to clone and to disrupt the gene(s) required for a pyruvate decarboxylase activity from both Torulaspora and Zygosaccharomyces yeast species. Further, it can be also anticipated that following the same classical approaches, as reported by Neveling U. et al. (1998, Journal of Bacteriology, 180(6): 1540–8, 1998), it is possible to isolate, to clone and to disrupt the gene(s) required for the PDH activity in both Torulaspora and Zygosaccharomyces yeast species.


The pyruvate dehydrogenase activity can be measured by known methods, e.g. according to Neveling U. et al. (supra). Suitable strains can be obtained by selecting mutants and/or engineering of wild-type or collection strains. Hundreds of mutants could be selected by "high throughput screen" approaches. The modulation of pyruvate decarboxylase...
lase activity by using nutrients supporting different glycolytic flow rates (Biotechnol. Prog. 11, 294–298, 1995) did not prove to be satisfactory.

A preferred method for decreasing or destroying the pyruvate decarboxylase activity and/or pyruvate dehydrogenase activity in a yeast strain according to the invention consists in the deletion of the corresponding gene or genes.

These deletions can be carried out by known methods, such as that disclosed in Bianchi et al., (Molecular Microbiol. 19 (1), 27–36, 1996; Flickweert M. T. et al., Yeast, 12:247–257, 1996 and Pronk J T. et al., Microbiology 140 (Pt 3):601–10, 1994), by deletion or insertion by means of selectable markers, for instance the UR3 marker, preferably the UR3 marker from Saccharomyces cerevisiae.

Alternatively, deletions, point-mutations and/or frame-shift mutations can be introduced into the functional promoters and genes required for the PDC and/or PDH activities. These techniques are disclosed, for instance, in Nature, 305, 391–397, 1983. An addition method to reduce these activities could be the introduction of STOP codons in the genes sequences or expression of antisense mRNAs to inhibit translation of PDC and PDH mRNAs.

A Kluyveromyces lactis strain wherein the PDC gene has been replaced by the UR3 gene of S. cerevisiae has already been described in Molecular Microbiology 19(1), 27–36, 1996.

The gene coding for lactate dehydrogenase may be of any species (e.g. mammalian, such as bovine, or bacterial), and it may code for the L(-)-LDH or the D(-)-LDH.

Alternatively, both types of LDH genes may be expressed simultaneously. Further, any natural or synthetic variants of LDH DNA sequences, any DNA sequence with high identity to a wild-type LDH gene, any DNA sequence complementing the normal LDH activity may be used.

As transporter gene, for example the JEN1 gene, encoding for the lactate transporter of S. cerevisiae, can be used.

The transformation of the yeast strains can be carried out by means of either integrative or replicative vectors, linear or plasmidial.

The recombinant cells of the invention can be obtained by any method allowing a foreign DNA to be introduced into a cell (Spencer J F et al., Journal of Basic Microbiology 28(5): 321–333, 1988), for instance transformation, electroporation, conjugation, fusion of protoplasts or any other known technique. Concerning transformation, various protocols have been described: in particular, it can be carried out by treating the whole cells in the presence of lithium acetate and of polyethylene glycol according to Ito H. et al. (J. Bacteriol., 153:163, 1983), or in the presence of ethylene glycol and dimethyl sulphoxide according to Durrens P. et al. (Curr. Genet., 18:7, 1990). An alternative protocol has also been described in EP 361991. Electroporation can be carried out according to Becker D. M. and Guarente L. (Methods in Enzymology, 194:18, 1991).

The use of non-bacteriial integrative vectors may be preferred when the yeast biomass is used, at the end of the fermentation process, as stock fodder or for other breeding, agricultural or alimentary purposes.

In a particular embodiment of the invention, the recombinant DNA is part of an expression plasmid which can be of autonomous or integrative replication.

In particular, for both S. cerevisiae and K. lactis, autonomous replication vectors can be obtained by using autonomous replication sequences in the chosen host. Especially, in yeasts, they can be replication origins derived from plasmids (2μ, pKD1, etc.) or even chromosomal sequence (ARS).

The integrative vectors can be obtained by using homologous DNA sequences in certain regions of the host genome, allowing, by homologous recombination, integration of the vector.

Genetic tools for gene expression are very well developed for S. cerevisiae and described in Romanos, M. A. et al. Yeast, 8:423, 1992. Genetic tools have been also developed to allow the use of the yeasts Kluyveromyces and Torulaspora spendes host cells for production of recombinant proteins (Spencer J F, et al., supra; Reiser J. et al., Advances in Biochemical Engineering/Biotechnology, 43, 75–102, 1990). Some examples of vectors autonomously replicating in K. lactis are reported, either based on the linear plasmid pKGl of K. lactis (de Lovencourt L. et al. J. Bacteriol., 154:737, 1982), or containing a chromosomal sequence of K. lactis itself (KARS), conferring to the vector the ability of self replication and correct segregation (Das S., Hollenberg C. P., Curr. Genet., 6:123, 1982). Moreover, the recognition of a 2μ-like plasmid native to K. drosophilanum (plasmid pKD1-U.S. Pat. No. 5,166,070) has allowed a very efficient host/vector system for the production of recombinant proteins to be established (EP-A-361 991). Recombinant pKGl-based vectors contain the entire original sequence, fused to appropriate yeast and bacterial markers. Alternatively, it is possible to combine part of pKGl, with common S. cerevisiae expression vectors (Romanos M. A. et al. Yeast, 8:423, 1992) (Chen et al., Curr. Genet. 16: 95, 1989).

It is known that the 2μ plasmid from S. cerevisiae replicates and is stably maintained in Torulaspora. In this yeast the expression of heterologous protein(s) has been obtained by a co-transformation procedure, i.e. the simultaneous presence of an expression vector for S. cerevisiae and of the whole 2μ plasmid. (Compagno C. et al., Mol. Microbiol., 3:1003–1010, 1989). As a result of inter and intra molecular recombinations, it is possible to isolate a hybrid plasmid, bearing the complete 2μ sequence and the heterologous gene; such a plasmid is in principle able to directly transform Torulaspora.

Moreover, an episomal plasmid based on S. cerevisiae AR1 sequence has also been described, but the stability of this plasmid is very low, Compagno et al. (supra).

Recently, an endogenous, 2μ-like plasmid named pTD1 has been isolated in Torulaspora (Blaisonneau J. et al., Plasmid, 38:202–209, 1997); the genetic tools currently available for S. cerevisiae can be transferred to the new plasmid, thus obtaining expression vectors dedicated to Torulaspora yeast species.


2μ-like plasmids from Zygosaccharomyces species are known and have been isolated in Z. rouxii (pSR1), in Z. bisporus (pSB3), in Z. fermentati (pSM1) and in Z. bailii (pSB2) (Spencer J E et al., supra).


Any yeast promoter, either inducible or constitutive, may be used according to the invention. To date, promoters used for the expression of proteins in *S. cerevisiae* are well described by Romanos et al. (supra). Promoters commonly used in foreign protein expression in *K. lactis* are *S. cerevisiae* PGK and PHOS (Romanos et al., supra), or homologous promoters, such as LAC4 (van den Berg J. A. et al., BioTechnology, 8:135, 1990) and KIPDC (U.S. Pat. No. 5,631,143). The promoter of pyruvate decarboxylase gene of *K. lactis* (KIPDC) is particularly preferred.

Vectors for the expression of heterologous genes which are particularly efficient for the transformation of *Kluyveromyces lactis* strains are disclosed in U.S. Pat. No. 5,166,070, which is herein incorporated by reference.

Pyruvate decarboxylase gene promoters, preferably from *Kluyveromyces lactis* species and even more preferably from *Kluyveromyces lactis*, disclosed in Molecular Microbiol. 19(1), 27–36, 1996, are particularly preferred. Triose phosphate isomerase and alcohol dehydrogenase promoters, preferably from Saccharomyces species and even more preferably from Saccharomyces *cerevisiae*, are also preferred (Romanos et al., supra).

For the production of lactic acid, the yeast strains of the invention are cultured in a medium containing a carbon source and other essential nutrients, and the lactic acid is recovered at a pH of 7 or less, preferably at a pH of 4.5 or less, and even more preferably at a pH of 3 or less. Since the pH of the culture medium is reduced, a lower amount of neutralizing agent is necessary. The formation of lactate salt is correspondingly reduced and proportionally less regeneration of free acid is required in order to recover lactic acid. The recovery process may employ any of the known methods (T. B. Vickroy, Volume 3, Chapter 38 of “Comprehensive Biotechnology,” (editor: M. Moo-Young), Pergamon, Oxford, 1985.) (R. Datta et al., FEMS Microbiology Reviews 16, 221–231, 1995). Typically, the microorganisms are removed by filtration or centrifugation prior to the lactic acid recovery. Known methods for lactic acid recovery include, for instance, the extraction of lactic acid into an immiscible solvent phase or the distillation of lactic acid or an ester thereof. Higher yields with respect to the carbon source (g of lactic acid/g of glucose consumed) and higher productivity (g of lactic acid/L/h) are obtained by growing yeast strains, particularly Saccharomyces strains, in media lacking Mg**+** and Zn**+** ions or having a reduced availability of said ions. Preferably, culture media will contain less than 5 mM of Mg**+** and/or less than 0.02 mM of Zn**+**.

The present invention offers the following advantages in the production of lactic acid:

1. When the fermentation is carried out at pH 4.5 or less, there is less danger of contamination by foreign microorganisms, as compared with the conventional process. Further, the fermentation facility can be simplified and the fermentation control can be facilitated.

2. Since less neutralizing agent is added to the culture medium for neutralization, there is correspondingly less need to use mineral acids or other regenerating agents for conversion of the lactate salt to free lactic acid. Therefore, the production cost can be reduced.

3. Since less neutralizing agent is added to the culture medium, the viscosity of the culture broth is reduced. Consequently, the broth is easier to process.

4. The cells separated in accordance with the present invention can be utilized again as seed microorganisms for a fresh lactic acid fermentation.

5. The cells can be continuously separated and recovered during the lactic acid fermentation, in accordance with the present invention, and hence, the fermentation can be carried out continuously.

6. Since the recombinant yeast strains lack ethanol production ability and pyruvate dehydrogenase activity or have both a reduced ethanol production and a reduced pyruvate dehydrogenase activity, the production of lactic acid can be carried out with higher yield in comparison to yeast strains having both a wild-type ability to produce ethanol and a wild-type ability for the utilization of pyruvate by the mitochondria.

7. The production of lactic acid by metabolically engineered non-conventional yeasts belonging to the Kluyveromyces, Torulaspora and Zygospaeromyces species can be obtained from non conventional carbon sources (i.e., galactose-lactose-sucrose-raffinose-maltose-cellobiose-arabinose-xylene, to give some examples), growing the cells in high-sugar medium and growing the cells in presence of high concentration of lactic acid.

**BRIEF DESCRIPTION OF THE FIGURES**

FIG. 1. Cloning of the lactate dehydrogenase gene shifts the glycolytic flux towards the production of lactic acid.

Key enzymatic reactions at the pyruvate branch-point are catalysed by the following enzymes: (1): pyruvate decarboxylase; (2): alcohol dehydrogenase; (3): acetaldehyde dehydrogenase; (4): acetyl-CoA synthetase; (5): acetyl-CoA shuttle from the cytosol to mitochondria; (6): acetyl-CoA shuttle from mitochondria to the cytosol; (7): heterologous lactate dehydrogenase. (8): pyruvate dehydrogenase. Enzymatic reactions involved in anaerobic syntheses have been omitted.

**FIG. 2.** Diagram of the plasmid pVC1.

**FIG. 3A, 3B.** Diagram of the plasmid pKSMD8/7 and pKSEXH16, respectively.

**FIG. 4.** Diagram of the plasmid pEPL2.

**FIG. 5.** Diagram of the plasmid pLC5.

**FIG. 6.** Diagram of the plasmid pLAT-ADH.

**FIG. 7A.** L(+)-Lactic acid production from the transformed *Kluyveromyces lactis* PM6-7a/pEPL2 during growth on Glu-YNB based media. The residual glucose concentration at T=49 was not detectable. Production of D(−)-lactic acid was not detectable too. The LDH specific activity was higher than 3 U/mg of total cell protein along all the experiment.

Similar results have been obtained using the bacterial *L. casei* LDH (data not shown). ((A) cells/ml; (−) pH value; (−) Ethanol production, g/l ([(E)]) L(+)-Lactic acid production, g/l)

**FIG. 7B.** L(+)-Lactic acid production from the transformed *Kluyveromyces lactis* PM6-7a/pEPL2 during growth on Glu-YNB based media. Medium was buffered at time T=0 (pH=5.6) using 200 mM phosphate buffer. In this text batch, the pH value decreases much later than during the text batch shown in FIG. 7A. The residual glucose concentration at T=49 was not detectable. The LDH specific activity was higher than 3 U/mg of total cell protein along all the experiment.

Similar results have been obtained using the bacterial *L. casei* LDH (data not shown). ((A) cells/ml; (−) pH value; (−) Ethanol production, g/l ([(E)]) L(+)-Lactic acid production, g/l)

**FIG. 8A.** L(+)-Lactic acid production from the transformed *Kluyveromyces lactis* PM1/C1[pEPL2] during
growth on Glu-YNB based media. The residual glucose concentration at T=60 was 12.01 g/l. Longer incubation times did not yield higher productions of both biomass and L(+)-Lactic acid. The LDH specific activity was higher than 3 U/mg of total cell protein along all the experiment.

Similar results have been obtained using the bacterial L. casei LDH (data not shown). ((A)) cells/ml; (-) pH value; (\(\cdot\)) Ethanol production, g/l ((\(\cdot\))\() L(+)-Lactic acid production, g/l

FIG. 8B. L(+)-Lactic acid production from the transformed Klyuvoromeces lactis PM1-Cl[pPE1.2] during growth on Glu-YNB based media. Medium was buffered at time T=0 (pH=5.6) using 200 mM phosphate buffer. In this text batch, the pH value decreases much later than during the text batch shown in FIG. 8A. The residual glucose concentration at T=87 was zero. The LDH specific activity was higher than 3 U/mg of total cell protein along all the experiment. ((A)) cells/ml; (-) pH value; (\(\cdot\)) Ethanol production, g/l ((\(\cdot\))\() L(+)-Lactic acid production, g/l

Similar results have been obtained using the bacterial L. casei LDH (data not shown).

FIG. 9A. L(+)-Lactic acid production from the transformed Klyuvoromeces BM3-12[pLAZ10] cells in stirred tank bioreactor (see also text) ((A)) cells/ml; (\(\cdot\)) Glucose concentration, g/l ((\(\cdot\))\() L(+)-Lactic acid production, g/l

FIG. 9B. L(+)-Lactic acid yield from the transformed Klyuvoromeces BM3-12[pLAZ10] cells in stirred tank bioreactor.

Glucose Vs lactic acid production. The yield (g/g) is 85.46%.

FIG. 10. L(+)-Lactic acid production from the transformed Torulaspora (syn. Zygossaccharomyces) debrueckii CBS817[pLAT-ADH] during growth on Glu-YNB based media. The residual glucose concentration at T=130 was 3 g/l. Longer incubation times did not yield higher productions of both biomass and L(+)-Lactic acid. The LDH specific activity was higher than 0.5 U/mg of total cell protein along all the experiment. ((A)) cells/ml; (-) pH value; (\(\cdot\)) Ethanol production, g/l ((\(\cdot\))\() L(+)-Lactic acid production, g/l

FIG. 11. L(+)-Lactic acid production from the transformed Zygossaccharomyces bailii ATCC60483[pLATADH] during growth on Glu-YNB based media. The residual glucose concentration at T=60 was 8 g/l. Longer incubation times did not yield higher productions of both biomass and L(+)-Lactic acid. The LDH specific activity was higher than 0.5 U/mg of total cell protein along all the experiment. Similar results were obtained using a different strain (ATCC36947, data not shown) ((A)) cells/ml; (-) pH value; (\(\cdot\)) Ethanol production, g/l ((\(\cdot\))\() L(+)-Lactic acid production, g/l

**DETAILED DESCRIPTION OF THE INVENTION**

**Definitions**

The following definitions are provided in order to aid those skilled in the art in understanding the detailed description of the present invention.

"Amplification" refers to increasing the number of copies of a desired nucleic acid molecule.

"Codon" refers to a sequence of three nucleotides that specify a particular amino acid.

"Deletion" refers to a mutation removing one or more nucleotides from a nucleic acid sequence.

"DNA ligase" refers to an enzyme that covalently joins two pieces of double-stranded DNA.

"Electroporation" refers to a method of introducing foreign DNA into cells that uses a brief, high voltage electric charge to permeabilize the host cells, causing them to take up extra-chromosomal DNA.

The term "endogenous" refers to materials originating from within the organism or cell.

"Endonuclease" refers to an enzyme that hydrolyzes double stranded DNA at internal locations.

The term "expression" refers to the transcription of a gene to produce the corresponding mRNA and translation of this mRNA to produce the corresponding gene product, i.e., a protein, polypeptide, or protein.

The term "expression of antisense RNA" refers to the transcription of a DNA to produce a first RNA molecule capable of hybridizing to a second RNA molecule encoding a gene product, e.g., a protein. Formation of the DNA-RNA hybrid inhibits translation of the second RNA molecule to produce the gene product.

The phrase "functionally linked" refers to a promoter or promoter region and a coding or structural sequence in such an orientation and distance that transcription of the coding or structural sequence may be directed by the promoter or promoter region.

The term "gene" refers to chromosomal DNA, plasmid DNA, cDNA, synthetic DNA, or other DNA that encodes a peptide, polypeptide, protein, or RNA molecule, and regions flanking the coding sequence involved in the regulation of expression.

The term "genome" encompasses both the chromosome and plasmids within a host cell. Encoding DNAs of the present invention introduced into host cells can therefore be either chromosomally-integrated or plasmid-localized.

"Heterologous DNA" refers to DNA from a source different than that of the recipient cell.

"Homologous DNA" refers to DNA from the same source as that of the recipient cell.

"Hybridization" refers to the ability of a strand of nucleic acid to join with a complementary strand via base pairing. Hybridization occurs when complementary sequences in the two nucleic acid strands bind to one another.

"Lactate dehydrogenase" (LDH) refers to a protein that catalyzes the conversion of pyruvate and NADH to lactic acid and AND. L(+)-LDH produces L(+)-lactic acid; D(-)-LDH produces D(-)-lactic acid.

The term "lactate transporter" refers to a protein that allows the transport of lactate from inside to outside the cell.

"Mutation" refers to any change or alteration in a nucleic acid sequence. Several types exist, including point, frame shift, and splicing. Mutation may be performed specifically (e.g., site directed mutagenesis) or randomly (e.g., via chemical agents, passage through repair minus bacterial strains).

Nucleic acid codes: A=adenosine; C=cytosine; G=guanosine; T/thymidine; N=equimolar A, C, G, and T; I=inosine; K=equimolar G and T; R=equimolar A and G; S=equimolar C and G; W=equimolar A and T; Y=equimolar C and T.

"Open reading frame (ORF)" refers to a region of DNA or RNA encoding a peptide, polypeptide, or protein.

"Pyruvate decarboxylase" (PDC) refers to a protein which catalyzes the conversion of pyruvate to acetaldehyde.
“Pyruvate dehydrogenase” (PDH) refers to a protein complex which catalyzes the conversion of pyruvate to acetyl-CoA.

“Plasmid” refers to a circular, extrachromosomal, self-replicating piece of DNA.

“Point mutation” refers to an alteration of a single nucleotide in a nucleic acid sequence.

“Polymerase chain reaction (PCR)” refers to an enzymatic technique to create multiple copies of one sequence of nucleic acid. Copies of DNA sequence are prepared by shutting a DNA polymerase between two amplifiers. The basis of this amplification method is multiple cycles of temperature changes to denature, then re-anneal amplifiers, followed by extension to synthesize new DNA strands in the region located between the flanking amplifiers.

The term “promoter” or “promoter region” refers to a DNA sequence, that includes elements controlling the production of messenger RNA (mRNA) by providing the recognition site for RNA polymerase and/or other factors necessary for start of transcription at the correct site.

A “recombinant cell” or “transformed cell” is a cell whose DNA has been altered by the introduction of an exogenous nucleic acid molecule into that cell.

The term “recombinant DNA construct” or “recombinant vector” refers to any agent such as a plasmid, cosmid, virus, autonomously replicating sequence, phage, or linear or circular single-stranded or double-stranded DNA or RNA nucleotide sequence, derived from any source, capable of genomic integration or autonomous replication, comprising a DNA molecule in which one or more DNA sequences have been linked in a functionally operative manner. Such recombinant DNA constructs or vectors can be capable of introducing a 5’ regulatory sequence or promoter region and a DNA sequence for a selected gene product into a cell in such a manner that the DNA sequence is transcribed into a functional mRNA which is translated and therefore expressed. Recombinant DNA constructs or recombinant vectors may alternatively be constructed to be capable of expressing antisense RNAs, in order to inhibit translation of a specific RNA of interest.

“Reduced (enzymatic) activity” refers to lower measured enzymatic activity isolated from a transformed or mutagenized strain as compared to the measured enzymatic activity isolated from a wild type strain of the same species. Reduced enzymatic activity may be the result of lowered concentrations of the enzyme, lowered specific activity of the enzyme, or a combination thereof.

“Repair minus” or “repair deficient” strains refer to organisms having reduced or eliminated DNA repair pathways. Such strains demonstrate increased mutation rates as compared to the rates of wild type strains of the same species. Propagation of a nucleic acid sequence through a repair minus strain results in the incorporation of random mutations throughout the nucleic acid sequence.

“Restriction enzyme” refers to an enzyme that recognizes a specific palindromic sequence of nucleotides in double stranded DNA and cleaves both strands, also called a restriction endonuclease. Cleavage typically occurs within the restriction site.

“Selectable marker” refers to a nucleic acid sequence whose expression confers a phenotype facilitating identification of cells containing the nucleic acid sequence. Selectable markers include those which confer resistance to toxic chemicals (e.g. ampicillin resistance, kanamycin resistance), complement a nutritional deficiency (e.g. uracil, histidine, leucine), or impart a visually distinguishing characteristic (e.g. color change, fluorescence).

“Transcription” refers to the process of producing an RNA copy from a DNA template.

“Transformation” refers to a process of introducing an exogenous nucleic acid sequence (e.g., a vector, plasmid, recombinant nucleic acid molecule) into a cell in which the exogenous nucleic acid is incorporated into a chromosome or is capable of autonomous replication.

“Translation” refers to the production of protein from messenger RNA.

The term “yield” refers to the amount of lactic acid produced (g/l) divided by the amount of glucose consumed (g/l).

“Unit” of enzyme refers to the enzymatic activity and indicates the amount of micromoles of substrate converted per mg of total cell proteins per minute.

“Vector” refers to a plasmid, cosmid, bacteriophage, or virus that carries nucleic acid sequences into a host organism.

Site-directed Mutagenesis of the Bovine Bacterial Dehydrogenase Gene (LDH-A)

In order to isolate from the full length cDNA the coding sequence of the bovine enzyme LDH-A (EC 1.1.1.27), a classical site directed mutagenesis, (J. Biol. Chem. 253:6551, 1978, Meth. Enzymol. 154:329, 1987), was performed. Oligonucleotide-driven site-specific mutagenesis is based on the in vitro hybridization of a single strand DNA fragment with a synthetic oligonucleotide, which is complementary to the DNA fragment except for a central mismatching region in correspondence of the DNA sequence that must to be mutagenized.

In order to introduce a Xba I restriction enzyme site 11 bp before the ATG codon, the 1743 bp bovine LDH cDNA was cloned from the plasmid pDH12 (Ishiguro et al., Gene, 91 281–285, 1991) by digestion with Eco RI and Hind III restriction enzymes (New England Biolabs, Beverly, Mass.). The isolated DNA fragment was then inserted in the pALTER-1 (Promega, cat #96210; lot #48645, 1996) expression vector.

Such vector contains M13 and R408 bacteriophages origin of replication and two genes for antibiotic resistance. One of these genes, for tetracycline resistance, is functional. The other (i.e., ampicillin resistance) has been inactivated. An oligonucleotide is provided which restores ampicillin resistance to the mutant strain during mutagenesis reaction (Sequence Id. No. 1, oligoAMP; Promega, Madison, Wis. Tab. 1). This oligonucleotide is annealed to the single-stranded DNA (ssDNA) template. At the same time the mutagenic oligonucleotide (Sequence Id. No. 2, oligoLDH, Madison Wis.) has been annealed as well. Following DNA synthesis and ligation, the DNA is transformed into a repair minus strain of E. coli (BMH 71-18 mutS; Kit Promega).

Selection was performed on LB+ampicillin (Molecular Cloning a laboratory manual, edited by Sambrook et al., Cold Spring Harbor Laboratory Press). A second round of transformation in JM 109 (kit Promega) E. coli strain ensured proper segregation of mutant and wild type plasmids.
Table 1: Nucleotide sequence of the synthetic oligonucleotides used for the site-directed mutagenesis. The underlined sequence in the oligoLDH show the Xba I restriction site introduced by mutagenesis.

Further details of the technique and material used (with the exception of the oligoLDH) can be found in the dataset.

The plasmid obtained, containing the mutated cDNA for the bovine LDH, was denominated pVC1 (FIG. 2).

PCR Mutagenesis of the Bacterial Lactate Dehydrogenase Gene (LDH) from Lactobacillus casei, Bacillus megaterium and Bacillus steaamorrhous.


We obtained plasmid pST2 and LDH sequence from Hutsolts Robert, University of Nebraska, USA). pST2 is based on pUC19 vector (Boehringer Mannheim GmbH, Mannheim, Germany, cat.885827) and contains a BamHI-SphI LDH-cDNA fragment amplified from the L. casei 686 (Culture collection of the University of Nebraska).

In order to obtain a coding sequence starting with the usual eukaryotic first codon (i.e. ATG), the LDH sequence has been mutagenised via PCR.

The introduction of a Neo I restriction enzyme site at position 163 of the LDH sequence (accession n. M76708 of the GenBank Sequence Database, supra), allows the concomitant change of the original GTG codon into ATG. PCR reaction (Mascycler 5330, Eppendorf, Hamburg, Germany) was performed starting from the plasmid pLC1, based on pGEM7Z f(+) (Promega corporation, Madison Wis., USA, cat.P2251) vector, and containing the L. casei's gene (fragment BamHI-SphI excided from pST2). The sequences of the oligonucleotides used as primers of the reaction are reported in Table 2 (Sequence Id. No. 3 and Sequence Id. No. 4).

At the end of the reaction, a single band, corresponding to the amplified and mutated gene, was isolated. The DNA fragment was then inserted at the EcoRV site of pMOSBlue.

Table 2: Nucleotide sequence of the synthetic oligonucleotides used for the PCR amplification. The underlined sequences in the oligoATG shows the NcoI restriction site introduced by mutagenesis, and the resulting ATG starting codon obtained.


Construction of the pEPL2 Replicative Vector Containing the KIPDCA Promoter and the Bovine LDH cDNA.

The KIPDCA promoter and the coding sequence were subcloned as a 4 Kbp HindIII fragment from a K. lactis genomic library clone complementing the ragA mutation of K. lactis (Bianchi et al., Mol. Microbiol., 19: 27-36, 1990). The promoter region was subcloned into Sal I and Xba I sites of the vector pBleuScript II KS (Stratagene, LaJolla, Calif. #212205) with T4 DNA ligase using molecular cloning standard procedure (Sambrook et al., Molecular Cloning, supra). The bovine LDH sequence, isolated as a Xba I-Hind III fragment of 1675 bp from the pVC1 vector, was cloned in the corresponding cloning sites of the vector pBleuScript II KS. GM288 E. coli strain (dam- dcm-) (available from ATCC or COSC collections) was transformed with the two new vectors, called respectively pKSMDS8/7 and pKSEXH/16 (FIG. 3A and 3B).

KIPDCA promoter and bovine LDH sequence, isolated as Sal I-Xba I fragments, respectively from pKSMDS8/7 and PKSEXH16, were ligated in vitro with T4 DNA ligase at room temperature in the presence of Sal I endonuclease in order to allow the ligation at Xba I ends. The ligation product was cloned in Sal I cloning site of pE1 vector (Bianchi M.et al., Curr. Genet. 12: 185-192, 1987; Chen X. J. et al., Curr. Genet. 16: 95-98, 1989 and U.S. Pat. No. 5,166,070). This plasmid is based on the YIp5 integrative plasmid containing the Saccharomyces cerevisiae genetic marker URA3 and on the pKD1 plasmid (U.S. Pat. No. 5,166,070), isolated from Kluyveromyces drosophilarum.
The plasmid pE1 has a functional organization similar to the S. cerevisiae 2μ DNA and is able to replicate in a stable way in K. lactis cells (Chen X. J. et al., supra). The URA3 marker on the plasmid allows the complementation of the K. lactis uraA1-1 mutation (de Louvencourt et al., J. Bacteriol. 154: 737–742 (1982)), and therefore growth of transformed cells in selective medium without uracil.

The vector obtained was called pEPL2 (FIG. 4) and used to transform E. coli DH5-α (α strain (Life Technologies Inc., Gaithersburg, Md.). Construction of the pEPL4 Replicative Vector Containing the KIPDC A Promoter and the Bacterial LDH Gene.

The bovine LDH gene described for the pEPL2 μ lasmid was substituted with the LDH DNA sequence from the bacterial Lactobacillus casei gene (see above), following classical molecular approaches described throughout the text, yielding the plasmid pEPL4. Transformed K. lactis yeast cells bearing the bovine or bacterial LDHs gave similar results.

Construction of the pLAZ10 Replicative Vector Containing the KIPDCA Promoter and the Bovine LDH cDNA.

Vector pLAZ10 was obtained by cloning the Sall fragment of pEPL2, bearing the KIPDCA1 promoter and the bovine LDH coding sequence, into the unique Sall site of vector pSK31. Vector pSK31 is composed of the commercial vector pUC19 and the G418 resistance cassette of vector pkan707 (Fluck et al. Bio/technology 9: 928–974, 1991) inserted in the unique SplI site of plasmid pKD1.


The L. casei LDH gene was excised from pLC3 described above) with a NcoI-Sall digestion, and ligated into pYX012 or pYX022 integrative vectors (R&D System Europe Ltd, Abingdon, England).

The two plasmids obtained, containing the mutated DNA for the bacterial LDH gene under the control of the TPI promoter, and carrying the auxotrophic markers URA3 or HIS3, were denominated respectively pLC5 (FIG. 5) and pLC7. For the construction of pB1, pB2M and pBST2 we used an approach similar to that described for the construction of pLC5; however, we used the bovine LDH, the B. megaterium LDH and the B. stearothermophilus LDH (Biolog Chem. Hoppe-Seyler, 1987, 368: 1391) (Biolog Chem. Hoppe-Seyler, 1987, 368: 1167), respectively. Finally, plasmid pFA6a-KanMX (Wach et al., Yeast, 10, 1994, 10:1793–1808) was digested with SacI and Smal and the resulting fragment was ligated into pLC5 cut with the same enzymes yielding the plasmid pLC5-KanMX. On the plasmids, the LDH gene is under the control of the TPI promoter.

The DNA sequence of JEN1 (the DNA sequence is available at the accession n. U24155 of the Genbank Sequence Database provided by the National Center of Biotechnology-NCBI-retrieved [online] [retrieved on Jul. 18, 1997] from the Internet: <URL: http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=Nucleotide&list_uids=7878081&dopt=GenBank>, encoding for the lactate transporter of S. cerevisiae (Davis E. S., Thesis, 1994-Laboratory of Eukaryotic Gene Expression, Advanced Bioscience Laboratories) (Davis, E. S. et al., Proc. Natl. Acad. Sci. U.S.A. 89 (23), 11169, 1992) (Andre, B. Yeast (11), 1575, 1995), has been obtained from E. S. Davis (University of Maryland, USA). The JEN1 coding sequence has been amplified by classical PCR approach described throughout the text and cloned into the plasmid pYX022 (see above). On the integrative plasmid, JEN1 overexpression is under the control of the TPI promoter.

Construction of the pLAT-ADH Replicative Vector Containing the ADH1 Promoter and the Bovine LDH cDNA.

First, the pLDH-Kan plasmid was constructed, cloning at EcoRV site of the pBlueScript II KS (Promega corporation, Madison Wis., USA, cat. 212028) cloning vector the APT1 gene, conferring geneticin (G418) resistance, derived from a Smal/EcoRV digestion of pFA6-KanMX4 vector (Wach et al. Yeast 10:1793–1808 (1994)).

Secondly, the coding region of bovine LDH gene was cloned under the control of S. cerevisiae’s ADH1 promoter and terminator sequences by subcloning a XbaI/HindIII fragment, from the previously described pVC1 plasmid into pVT102-U vector (Vernet et al. Gene 52:225–233 (1987)).

Finally, the whole expression cassette (ADH1 promoter—LDH gene—ADH1 terminator) was excised with a SphI digestion and ligated with pLDH-Kan, linearized with SphI, obtaining pLAT-ADH vector (FIG. 6).

Isolation of the K. lactis PMI/C1 Strain

Deletion of the KIPDCA gene in the PM6-7A yeast strain (MATa, adeI-F-600, uraA1-1) (Wesolowski et al., Yeast 1992, 8: 711) yielded the strain PMI. Deletion has been carried out by insertion of the URA3 marker of S. cerevisiae. The strain PMI grows on glucose containing media; PDC activity is not detectable and the strain does not produce ethanol (Bianchi M. M., et al.,(1996), supra). It is important to underline that S. cerevisiae cells without any detectable PDC activity do not grow on glucose mineral media (Flikweert M. T. et al., Yeast, 12:247–257, 1996).

1x10^7–3x10^7 cells from a stationary culture of PMI yeast cells were plated on synthetic medium containing 5-fluoroorotic acid. Growth of yeast cells in media containing 5-fluoroorotic acid allows the selection of cells impaired in uracil synthesis (McCusker and Davis, Yeast 7: 607–608 (1991)). After 5 days incubation at 28° C. some ura-mutants were isolated. One of these mutants obtained, called PMI/C1, resulted mutated in the URA3 gene previously introduced by integrative transformation, as it resulted from a complementation test by transformation with an URA3 gene-containing plasmid (Kep6 vector; Chen et al., J. Basic Microbiol. 28: 211–220 (1988)).

The genotype of PMI/C1 is the following: MAIa, adeI-F-600. uraA1-1, pdcA:ura3.

Isolation of the CENPK113APDC1APDC5APDC6 CENPK113APDC2 and GRF181APDC2 strain

The general strategy was to generate first single deletion mutants of each of the PDC genes (PDC1, PDC2, PDC5 and PDC6). The gene deletions were performed by integration of a loxP-Kan<sup>′</sup>-<sup>′</sup>-loxP cassette by homologous recombination at the locus of the corresponding PDC gene using the short flanking homology (SFi) PCR method described by Wach et al. (1994; Yeast 10, 1793–1808) and Guldener et al. (1996; Nucleic Acids Res. 24, 2519–2524). Subsequently the deletion cassette was removed by expressing the cre recombinase leading behind a single copy of the loxP site at the deletion locus. The pdc1 pdc5 pdc6 triple deletion mutant was created by subsequently crossing the single haploid deletion strains.
The PCR reaction was carried out on a DNA-template containing the gene for the kanamycin resistance (open reading frame of the E. coli transposon Tn903) fused to control sequences (promotor/terminator) of a certain <i>Schwannomyces occidentalis</i> gene (confidential). This selection cassette is flanked on both ends by a loxP sequence (loxP-Kan<sup>R</sup>-loxP) and was developed by SRD (Scientific Research and Development GmbH). The used primer to amplify the loxP-Kan<sup>R</sup>-loxP cassette are designed so that the DNA sequence of the sense primer is homologous to the 5'-end of the selection cassette sequence and so that the primer presents in addition at its 5'-end a region 40 nucleotides, which corresponds to the 5'-terminal sequence of the certain <i>Sacharomyces cerevisiae</i> PDC gene. The antisense primer is constructed in an analogous manner, it is complementary to the 3'-end of the selection cassette, wherein this primer contains at its 5'-end a region of also preferably 40 nucleotides, which corresponds to the 3'-terminal sequence of the certain <i>Sacharomyces cerevisiae</i> PDC gene.

The following table shows the primers (Sequence Id. Nos. 5–12) used for gene deletion of the corresponding PDC genes by SFH PCR method. Sequences underlined are homologous to the corresponding PDC gene and sequences complementary to the loxP-Kan<sup>R</sup>-loxP cassette are in bold letters.

<table>
<thead>
<tr>
<th>Reference Number</th>
<th>Primer Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sequence Id. No. 5</td>
<td>TCC TTA CTA TAA CCT CAC GCA AAA TAA CAC ACTCAA ATG ACA GCT GAA GCT TGT TAC GC</td>
</tr>
<tr>
<td>Sequence Id. No. 6</td>
<td>ATT GCT TAT AAA ACT TTA ACT ATT ATG AGA TTA ACC CAC ATA GCA CAC TAT TGC ATT TC</td>
</tr>
<tr>
<td>Sequence Id. No. 7</td>
<td>TCC ATT GCT AAA GAC TAC ACA AAG ACA ATA CAA TCA CAA CCA GCA GAA GCT TCC TGC CAC GC</td>
</tr>
<tr>
<td>Sequence Id. No. 8</td>
<td>AAA ATA CAC AAA CTT TTA ATT ATG ATT TTT ATT TTA ATT AGC ATA GCA CAC TAC TAG TGT ATC TG</td>
</tr>
<tr>
<td>Sequence Id. No. 9</td>
<td>TAA AAT AAA AAC CCA CTT ATT ATA GCA AAA ACA TAT TGC CCA GCT GAA GCT TCG TAC GC</td>
</tr>
<tr>
<td>Sequence Id. No. 10</td>
<td>TTT ATT TGC AAC RAT ATT TGG ATT TAC ACT ACT ATT GCC ATA GCA CAC TAC TAG TGG ATC TG</td>
</tr>
<tr>
<td>Sequence Id. No. 11</td>
<td>ACG CAA CTT GAA TGG GCA AAA TGG GCT TAT GAG ACG TTC CCA GCT GAA GCT TCG TAC GC</td>
</tr>
<tr>
<td>Sequence Id. No. 12</td>
<td>AGC CTA TGC TAC GAT GCA ATT GCA AGT ATT TAT TAC ACT CAT GCC ATA GCA CAC TAG TGC ATC TG</td>
</tr>
</tbody>
</table>

The PCR amplified deletion cassette was used for the transformation of the prototrophic diploid <i>Sacharomyces cerevisiae</i> strain CEN.PK122 developed by SRD.CEN.PK 122 (Mata, alpha, URA3, URA3, HIS3, HIS3, LEU2, LEU2, TRPI, TRP1, MAL2-8', MAL2-8', SUC2, SUC2)

For selection of transformants, geneticin (G-418 sulfate, Life Technologies) was added at a final concentration of 200 mg/l. After tetrad analysis, G418 resistant spores were subsequently analyzed by diagnostic PCR to confirm the correct deletion of the corresponding PDC gene and to determine the mating type of the haploid strain.

To obtain a strain deleted for the three PDC genes, PDC1, PDC5 and PDC6, the haploid deletion strains were subsequently crossed. To obtain the two double deletion strains, pdc1::Kan<sup>R</sup> pdc6::Kan<sup>R</sup> and pdc5::Kan<sup>R</sup> pdc6::Kan<sup>R</sup>, the corresponding haploid strains were crossed. After tetrad analysis, spores showing the non-parental ditype for the Kan<sup>R</sup> marker were subsequently analyzed by diagnostic PCR to confirm the correct deletion of both genes and to determine the mating type. The resulting double deletion strains were crossed to obtain the triple deletion strain. After tetrad analysis there was obtained by diagnostic PCR for spores which are deleted for the three PDC genes.

To eliminate the Kan<sup>R</sup> marker from the successfully disrupted gene the haploid deletion strains (single, double and triple mutants) were transformed with the cre recombinase plasmid, pPK-ILV2<sup>SMR</sup> (developed by SRD). Plasmid pPK-ILV2<sup>SMR</sup> contains the cre-recombinase under the control of the GAL1 promoter and as dominant selection marker the ILV2 resistance gene, which allows yeast cells transplasmid pPK-ILV2<sup>SMR</sup> to grow in the presence of sulfofomethuron methyl (30 mg/l). Correct excision of the Kan<sup>R</sup> marker was subsequently analyzed by diagnostic PCR with whole yeast cells. To remove plasmid pPK-ILV2<sup>SMR</sup> the yeasts cells were incubated for an appropriate time without sulfofomethuron methyl in the medium and subsequently searching for sulfofomethuron methyl sensitive cells.

The following table shows the resulting yeast strains. The numbers in parentheses indicate the deleted nucleotides (AA/G=1) of the corresponding genes. In the case of negative numbers means the first number the deleted nucleotides upstream of the ATG and the second number the deleted nucleotides downstream of the STOP codon.
| CEN.PK184 MA15 URA3 HIS3 LEU2 TRP1 MAL2-86 | Suc2 pdc1(-6,-2):loxP | CEN.PK186 MA15 URA3 HIS3 LEU2 TRP1 MAL2-86 | Suc2 pdc5(-6,-2):loxP | CEN.PK187 MA15 URA3 HIS3 LEU2 TRP1 MAL2-86 | Suc2 pdc6(-6,-2):loxP | CEN.PK188 MA15 URA3 HIS3 LEU2 TRP1 MAL2-86 | Suc2 pdc6(-6,-2):loxP | CEN.PK189 MA15 URA3 HIS3 LEU2 TRP1 MAL2-86 | Suc2 pdc6(-6,-2):loxP |
| CEN.PK190 MA15 URA3 HIS3 LEU2 TRP1 MAL2-86 | Suc2 pdc6(-6,-2):loxP | CEN.PK201 MA15 URA3 HIS3 LEU2 TRP1 MAL2-86 | Suc2 pdc2(101, 2730):loxP |

| TABLE |

Mainly we used the strains CEN.PK211 and CEN.PK182 which, in the tables summarizing the data obtained, are named CEN.PK113A PDC2 and CEN.PK113 A PDC2. Using a similar approach a S. cerevisiae GFR18U strain (Mat alpha, his3, leu2, ura3) bearing a deletion in the PDC2 gene was build (GFR18U A PDC2; Mat alpha, his3, leu2, ura3, pdc2::AP1T). We used the AP1T gene, conferring G418 resistance, as a marker of the integration, isolated from the plasmid pA6a-KanMX (Wach et al. supra). For the strain bearing deletions in the PDC1, PDC5 and PDC6 genes, the PDC activity is zero. For the strains bearing a deletion in the PDC2 gene, the PDC activity is about 20–40% of the level determined in the wild-type strains. Isolation of the K. lactis BM3-12D [pLAZ10]

A double deletant strain Klpld1a/Klpld1a was selected from the haploid segregant population of a diploid strain obtained by crossing strain MW341-5/Klpld1a (MA1A, lac4-8, leu2 lys1-A1, ura1-A1, Klpld1::URA3) obtained as previously described in Bianchi et al. 1996, Mol. Microbiol. 19(1), 27–36; Destrieul et al., submitted) with strain CBS2359/Klpld1a (MA1A, URA3-48, Klpld1:: Tn5BLS) Deletion of the PDA1 gene, encoding for the pyruvate dehydrogenase complex E1-alpha subunit (EC1.2.4.1) (the DNA sequence has been obtained by Stecchina H. Y., Faculty of Mathematics and Natural Sciences, Clusius Laboratory, Leiden, The Netherlands—the DNA sequence is also available at the accession n. AF023920 of the Genbank Sequence Database provided by the National Center of Biotechnology—NCBI—retrieved [online] [retrieved Jul. 18, 1997] from the Internet: <URL:http://www.ncbi.nlm.gov/entrez/query.fcgi?cmd=Retrieve&db= Nucleotide&list_uids=2558802&dopt=GenBank>, in the yeast strain CBS2359 has been obtained following the classical PCR approach and yeast transformation described throughout the text. We used the marder Tn5BLS (Gatignol et al., Gene, 91: 35, 1990) conferring phleomycin resistance, as a marker of the integration.

The double deletant strain, called BM1-3C (MA1A, leu2, Klpld1:: URA3; Klpld1:: Tn5BLS), was selected as a phleomycin resistant/antimycin sensitive segregant strain. The vector pLAZ10 was then genetically transferred to the double deletant strain as follows: A pLAZ10 transformant of the Klpld1:: URA3 strain PML8 (MA1A, ade1-600, ura1-A1, Klpld1:: URA3; Bianchi et al., Mol. Microbiol. 19: 27–36; 1996) was crossed with strain MW109-8C (MA1X, lys1-A1, trp1-A1). After sporulation of the resulting diploid strain, a genetin resistant/antimycin sensitive strain, called strain 7C (MA1X, ade1-600, lys1-A1, Klpld1:: URA3; plAZ10), was selected.

Strain BM1-3C and strain 7C were crossed and phleomycin resistant/antimycin resistant haploid segregant strains were selected after sporulation of the obtained diploid strain. All of these strains (MA1X, ade1-600, lys1-A1, Klpld1:: URA3; plAZ10) were crossed with the prototroph strain BM3-12D (Klpld1:: URA3; Klpld1:: Tn5BLS, plAZ10) was chosen for further experiments.

Transformation of Klyuyveromyces Yeast PM6-7A and PMI/ C1 with the Vectors pEPL2 and pEPL4.

PM6-7A and PMI/C1 cells were grown in YPD medium until a concentration of 0.5×10^6 cells/ml harvested, washed once in water, twice in 1M sorbitol, and resuspended in 1M sorbitol at a concentration of 2×10^6 cells/ml. Cells were electroporated (7.5 kV/cm, 25 μF, 200 Ω; GenePulser, Biorad, Hercules, Calif.) in the presence of 5–10 microg of pEPL2 or pEPL4. Selection of URA^+ transformants was carried out in synthetic solid medium without uracil (0.7% w v Yeast Nitrogen Base, 2 w/v % glucose, 200 mg/ml adenine, 2 w/v % agar).

Transformation of Torulaspora Yeast with the Vector pPAA-ADH.

CBS817 cells were grown in YPD medium until a concentration of 6×10^7 cells/ml harvested, washed once in water, twice in 1M sorbitol, and resuspended in 1M sorbitol at a concentration of 2×10^6 cells/ml. Cells were electroporated (1.5 kV, 7.5 kV/cm, 25 μF, 200 Ω; GenePulser, Biorad, Hercules, Calif.) in the presence of 1 μg of pPAA-ADH.

Cells were grown overnight in sterile microbiological tubes containing 5 ml of YEPD, Sorbitol 1M. Selection of G418^+ transformants was carried out in solid medium (2 w/v % glucose, 2 w/v % Peptone, 1 w/v % Yeast extract, 2 w/v % agar, 200 μg/ml G418 (Gibco BRL, cat. 11811-031). Transformation of Saccharomyces Yeast Cells with the Vectors pLCS, PLC7, pB1, pBST2, pBME2, pPAI-ADH, pLCS-kanMX and pJEN1.

GFR18U (described above), GFR18UPAD2 (described above), GFR18UPLC5 (Mat alpha, his3, leu2, ura3: TPI-LDH), CENPK113 (Mat a; CBS8340), CENPK1-1 (Mat a, ura3), CENPK113A PDC1 PDC5 A PDC6 (described above) and CENPK113A PDC2 (described above) yeast cells were grown in rich YPD complete medium (2% w/v yeast extract, 1% w/v peptone, 2% w/v glucose) until a concentration of 2×10^6 cells/ml, washed once in 0.1 M lithium acetate, 1 mM EDTA, 10 mM Tris-HCl, pH 8, harvested, and resuspended in 0.1 M lithium acetate, 1 mM EDTA, 10 mM Tris-HCl, pH 8, at a concentration of 2×10^6 cells/ml. 100 μl of the cellular
suspension were incubated 5 minutes with 5–10 µg of vector (i.e., previously linearized in the auxotrophic marker in the case of pLCS, pLC7, pBL1, pBST2, pBME2, pLC5-kanMX, pJEN1). After the addition of 280 µl of PEG 4000, the cells were incubated for at least 45° at 30° C. 43 µl of DMSO were added and the suspension was incubated at 42° C. The cells were washed twice with water and plated onto selective medium. For the isolation of CENPK-1 strain (ura3), CENPK113 cells were grown in media containing 5-fluorouracil acid (see also above).

Single transformed clones were scored with 0.7% w/v Yeast Nitrogen Base, 2 w/v % glucose, 2 w/v % agar plus the appropriate supplements or G418 as indicated. For selection of G418 transformants, cells were also scored on 2 w/v % glucose, 2 w/v % Peptone, 1 w/v % Yeast extract, 2 w/v % agar, 200 µg/ml G418 (Gibco BRL, cat. 11811–031).

Transformed Strain: Supplements.

GRF18U[pLAT-ADH]: 200 mg/l uracile, 200 mg/l leucine, 200 mg/l histidine, 200 mg/l G418.
GRF18U[pB1]: 200 mg/l leucine, 200 mg/l histidine.
GRF18U[pLC5]: 200 mg/l leucine, 200 mg/l histidine.
GRF18U[pLC5][pL7]: 200 mg/l leucine.
GRF18U[pBM2]: 200 mg/l leucine, 200 mg/l histidine.
GRF18U[pBST2]: 200 mg/l leucine, 200 mg/l histidine.
GRF18U[pLC5][pJEN1]: 200 mg/l leucine.
GRF18U[pDP2][pLC5]: 200 mg/l leucine, 200 mg/l histidine.
CENPK-[pLC5]: no supplements
CENPK113[pLC5-kanMX]: 200 mg/l G418
CENPK113[pDP2][pLC5-kanMX]: 200 mg/l G418
List of the Expression Vectors Used:

<table>
<thead>
<tr>
<th>Name</th>
<th>LDH source</th>
<th>promoter</th>
<th>Host, Selective marker</th>
</tr>
</thead>
<tbody>
<tr>
<td>pEPL2</td>
<td>Bovine</td>
<td>KLPDCA</td>
<td>K. lactis, URA3. (FIG. 4)</td>
</tr>
<tr>
<td>pEPL4</td>
<td>L. casei</td>
<td>KLPDCA</td>
<td>K. lactis, APT1.</td>
</tr>
<tr>
<td>PLAZ10</td>
<td>Bovine</td>
<td>KLPDCA</td>
<td>K. lactis, APT1.</td>
</tr>
<tr>
<td>pLC5</td>
<td>L. casei</td>
<td>SCP1</td>
<td>S. cerevisiae, URA3. (FIG. 5)</td>
</tr>
<tr>
<td>pLC5-kanMX</td>
<td>L. casei</td>
<td>SCP1</td>
<td>S. cerevisiae, URA3.</td>
</tr>
<tr>
<td>pBM2</td>
<td>B. megate-</td>
<td>SCP1</td>
<td>S. cerevisiae, URA3.</td>
</tr>
<tr>
<td>pBST2</td>
<td>B. stearo-</td>
<td>SCP1</td>
<td>S. cerevisiae, URA3.</td>
</tr>
<tr>
<td>pB1</td>
<td>Bovine</td>
<td>SCP1</td>
<td>S. cerevisiae, URA3.</td>
</tr>
<tr>
<td>pLC7</td>
<td>L. casei</td>
<td>SCP1</td>
<td>S. cerevisiae, HIS3.</td>
</tr>
<tr>
<td>pJEN1</td>
<td>//</td>
<td>SCP1</td>
<td>S. cerevisiae, HIS3.</td>
</tr>
<tr>
<td>pLAT-ADH</td>
<td>Bovine</td>
<td>SCADMB1</td>
<td>APT1-URA3. (FIG. 6)</td>
</tr>
</tbody>
</table>

KL = K. lactis’s promoter
SC = S. cerevisiae’s promoter
B. Ste = Bacillus stearothermophilus
pJEN1 has been used for the overexpression of the JEN1 gene.

**BATCH TESTS**


Clones obtained by the transformation procedure above described were tested in batch culture during growth on minimum synthetic medium (1.3% w/v Yeast Nitrogen Base—aa (Difco, Detroit, Mich.), 200 mg/l adenine, 50 g/l glucose). The media used were both buffered or not with 200 mM phosphate buffer to a pH of 5.6.

Cells were preinoculated in the same test’s medium. Exponentially growing cells were inoculated in flask (300 ml volume) containing 100 ml of fresh medium. The flasks were incubated at 30° C. in a shaking bath (Dubnoff, 150 rpm), and fermentation was monitored at regular time points. Cell number concentration was determined with an electronic Coulter counter (Coulter Counter ZBI Coulter Electronics Harpenden, G B, Porro et al., Res. Microbiol. (1991) 142, 535–539), after sonication of the samples to avoid cellular aggregates (Sonicator Fisher 300, medium point, Power 35%, 10 seconds) (FIGS 7 and 8 and Tab.3).


Clones obtained by the procedure above described were tested in batch culture during growth on minimum synthetic medium (1.3% w/v Yeast Nitrogen Base—aa (Difco, Detroit, Mich.), 50 g/l glucose, 20 g/l ethanol, 200 mg/l G418). The media used were buffered with 200 mM phosphate buffer to a pH of 5.6.

Cells were preinoculated in the same test’s medium. Exponentially growing cells were inoculated in flask (300 ml volume) containing 100 ml of fresh medium. The flasks were incubated at 30° C. in a shaking bath (Dubnoff, 150 rpm), and fermentation was monitored at regular intervals. Cell number concentration was determined with an electronic Coulter counter (Coulter Counter ZBI Coulter Electronics Harpenden, G B, Porro et al., Res. Microbiol. (1991) 142, 535–539), after sonication of the samples to avoid cellular aggregates (Sonicator Fisher 300, medium point, Power 35%, 10 seconds). At the beginning, cells used ethanol and then transformed glucose to lactic acid with very high yield (>0.75; g of lactic acid/g glucose consumed) (Tab.5).


Clones obtained by the transformation procedure above described were tested in batch culture during growth on minimum synthetic medium (1.3% w/v Yeast Nitrogen Base—aa (Difco, Detroit, Mich.), 20 g/l glucose, 200 mg/l G418). The media used were not buffered.

Cells were preinoculated in the same test’s medium. Exponentially growing cells were inoculated in flask (300 ml volume) containing 100 ml of fresh medium. The flasks were incubated at 30° C. in a shaking bath (Dubnoff, 150 rpm), and fermentation was monitored at regular intervals. Cell number concentration was determined with an electronic Coulter counter (Coulter Counter ZBI Coulter Electronics Harpenden, G B, Porro et al., Res. Microbiol. (1991) 142, 535–539), after sonication of the samples to avoid cellular aggregates (Sonicator Fisher 300, medium point, Power 35%, 10 seconds) (FIG. 10 and Tab.3).


Clones obtained by the transformation procedure above described were tested in batch culture during growth on minimum synthetic medium (1.3% w/v Yeast Nitrogen Base—aa (Difco, Detroit, Mich.), 50 g/l glucose, 200 mg/l G418). The media used were not buffered.

Cells were preinoculated in the same test’s medium. Exponentially growing cells were inoculated in flask (300 ml volume) containing 100 ml of fresh medium. The flasks were incubated at 30° C. in a shaking bath (Dubnoff, 150 rpm), and fermentation was monitored at regular intervals. Cell number concentration was determined with an electronic Coulter counter (Coulter Counter ZBI Coulter Electronics Harpenden, G B, Porro et al., Res. Microbiol. (1991) 142, 535–539), after sonication of the samples to avoid...
cellular aggregates (Sonicator Fisher 300, medium point, Power 35%, 10 seconds) (FIG. 11 and Tab. 3).

Batch Analysis of Saccharomyces GFR18U[pLAI-ADH], GFR18U[pB1], GFR18U[pLC5], GFR18U[pLC7], GFR18U[pBM2], GFR18U[pBST2], CENPK1-[pLC5] Transformed Cells.

Clones obtained by the transformation procedure above described were tested in batch culture during growth on minimum synthetic medium (1.3% w/w Yeast Nitrogen Base—aa (Difco, Detroit, Mich.), 50 g/l glucose and appropriate supplements (see above). The media used were not buffered. Cells were preincubated in the same medium. Exponentially growing cells were inoculated in flask (300 ml volume) containing 100 ml of fresh medium. The flasks were incubated at 30°C in a shaking bath (Dubnoff, 150 rpm), and fermentation was monitored at regular intervals. Cell number concentration was determined with an electronic Coulter counter (Coulter Counter ZBI Coulter Electronics Harpenden, GB, Porro et al., Res. Microbiol. (1991) 142, 535–539), after sonication of the samples to avoid cellular aggregates (Sonicator Fisher 300, medium point, Power 35%, 10 seconds) (Tab. 3).

Batch Analysis—spinner flask—of Saccharomyces, GFR18UAPDC2[pLC5], CENPK113 [pLC5-kanMX], CENPK113APDC2[pLC5-kanMX] and CENPK113APDC1ΔAPDC5APDC6[pLC5-kanMX] Transformed Cells.

Clones obtained by the procedures above described were tested in batch culture during growth on rich medium (1.0% w/w Yeast Extract, 2% w/v Peptone, 100 g/l glucose). The media were not buffered.

Cells were preincubated on Yeast Extract–Peptone–ethanol (5g/l) media. 100 ml were inoculated in spinner flasks (5 Cells working volume: 10ml). The spinner flasks were incubated at 30°C, agitation: 55 rpm. Fermentation was monitored at regular intervals (Tables A,B,C and Table 3).

The LDH specific activity from the different transformed strains, was higher than 5 U/mg of total cell proteins.

LDH Activity Assay

Bovine LDH. About 10^8 cells were harvested, washed in 50 mM phosphate buffer, pH 7.5, and resuspended in the same buffer. Cells were lysed with 5 cycles of vigorous vortexing in presence of glass microbeads (diameter 400 μm, SIGMA, G-8772) at 4°C. Cellular debris were removed by centrifugation (Eppendorf, Hamburg, D 5415 C, 13600 RCF, 10 min), and protein extracts’ concentration was determined by Micro Assay, Biorad, Hercules, Calif. (cat. 500-0006).

About 0.2 mg of extract were tested for LDH activity using SIGMA (St. Louis, Mo.) kit DG1340-UV, according to manufacturer’s instructions.

Bacterial LDHs. About 10^9 cells were harvested, washed in 50 mM phosphate buffer, pH 7.5, and resuspended in the same buffer. Cells were lysed with 5 cycles of vigorous vortexing in presence of glass microbeads (diameter 400 μm, SIGMA, G-8772) at 4°C. Cellular debris were removed by centrifugation (Eppendorf, Hamburg, D 5415 C, 13600 RCF, 10 min), and protein extracts’ concentration was determined by Micro Assay, Biorad, Hercules, Calif. (cat. 500-0006).

Cellular extract was tested for LDH activity using: 0.01 ml of 12.8 mM NADH, 0.1 ml of 2 mM fructose, 6-diphosphate, 0.74 ml of 50 mM acetyl buffer (pH=5.6), 0.05 ml of properly diluted cell extract and 0.1 ml sodium pyruvate 100 mM.

LDH activity was assayed as micromoles of NADH oxidised per min, per mg of total cell extract at 540 nm, 25°C.

Metabolites Dosage in the Growth Medium.

Samples from the growth medium, obtained after removing cells by centrifugation, were analysed for the presence of glucose, ethanol, I(-)- and D(-)-lactic acid using kits from Boehringer Mannheim, Mannheim Del., (F. 716251, 176290, and 1112821 respectively), according to manufacturer’s instructions.

Experimental batch tests related to the Kluuyveromyces PM6-7A[pEPL2] and PMCl[pEPL2] transformed yeasts are shown in FIGS. 7A, 7B and FIGS. 8A, 8B.

Experimental data related to the Zygossaccharomyces ATCC60483 [pLAI-ADH] transformed yeasts are shown in FIG. II.

Experimental data related to the Saccharomyces CENPK113[pLC5-KanMX]CENPK113 ΔAPDC2[pLC5- kanMX] and CENPK113Δ APDC3ΔAPDC5ΔPC6[pLC5- kanMX] transformed yeasts growing in spinner-flask are shown in Tables A,B,C.

TABLE A

<table>
<thead>
<tr>
<th>Time [h]</th>
<th>pH</th>
<th>OD_{560}</th>
<th>glucose [g·l^{-1}]</th>
<th>ethanol [g·l^{-1}]</th>
<th>lactate [g·l^{-1}]</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0</td>
<td>5.7</td>
<td>0.31</td>
<td>88.4 ± 0.3</td>
<td>0.2</td>
<td>0</td>
</tr>
<tr>
<td>19.0</td>
<td>3.01</td>
<td>8.7 ± 0.2</td>
<td>65. ± 0.1</td>
<td>25</td>
<td>27.4 ± 0.2</td>
</tr>
<tr>
<td>25.0</td>
<td>3.05</td>
<td>10.41 ±0.01</td>
<td>0.4 ± 0.2</td>
<td>37</td>
<td>30.1 ± 0.4</td>
</tr>
<tr>
<td>45.25</td>
<td>3.07</td>
<td>13.2 ± 0.2</td>
<td>0.06 ± 0.03</td>
<td>37</td>
<td>31.1 ± 0.2</td>
</tr>
<tr>
<td>70.75</td>
<td>3.08</td>
<td>10.6 ± 0.3</td>
<td>0</td>
<td>26</td>
<td>30.7 ± 0.1</td>
</tr>
<tr>
<td>92.0</td>
<td>3.08</td>
<td>12.2 ± 0.8</td>
<td>0</td>
<td>26</td>
<td>29.5 ± 0.1</td>
</tr>
</tbody>
</table>

TABLE B

<table>
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<tr>
<th>Time [h]</th>
<th>pH</th>
<th>OD_{560}</th>
<th>glucose [g·l^{-1}]</th>
<th>ethanol [g·l^{-1}]</th>
<th>lactate [g·l^{-1}]</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0</td>
<td>5.75</td>
<td>0.22</td>
<td>87 ± 0.4</td>
<td>0.2</td>
<td>0</td>
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<tr>
<td>19.0</td>
<td>3.20</td>
<td>2.2 ± 0.2</td>
<td>47.8 ± 0.1</td>
<td>8</td>
<td>17.3 ± 0.1</td>
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<tr>
<td>25.0</td>
<td>3.07</td>
<td>4.45 ± 0.1</td>
<td>36.3 ± 0.1</td>
<td>11</td>
<td>25.4 ± 0.1</td>
</tr>
<tr>
<td>45.25</td>
<td>2.96</td>
<td>5.31 ± 0.02</td>
<td>24.0 ± 0.1</td>
<td>15</td>
<td>38.0 ± 0.1</td>
</tr>
<tr>
<td>70.75</td>
<td>2.98</td>
<td>4.8 ± 0.1</td>
<td>12.9 ± 0.1</td>
<td>19</td>
<td>42.4 ± 0.1</td>
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<tr>
<td>92.0</td>
<td>2.95</td>
<td>5.3 ± 0.1</td>
<td>8.47 ± 0.01</td>
<td>24</td>
<td>43.1 ± 0.1</td>
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</table>

TABLE C

<table>
<thead>
<tr>
<th>Time [h]</th>
<th>pH</th>
<th>OD_{560}</th>
<th>glucose [g·l^{-1}]</th>
<th>ethanol [g·l^{-1}]</th>
<th>lactate [g·l^{-1}]</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0</td>
<td>5.74</td>
<td>0.82 ± 0.01</td>
<td>92 ± 1</td>
<td>0.171 ± 0.005</td>
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<tr>
<td>10.0</td>
<td>5.16</td>
<td>1.185 ± 0.02</td>
<td>93 ± 1</td>
<td>0.715 ± 0.00</td>
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<tr>
<td>23.5</td>
<td>4.61</td>
<td>1.28 ± 0.03</td>
<td>94 ± 0.2</td>
<td>1.76 ± 0.1</td>
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<tr>
<td>49.25</td>
<td>4.05</td>
<td>1.36 ± 0.03</td>
<td>92.8 ± 0.8</td>
<td>3.614 ± 0.005</td>
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</tr>
<tr>
<td>73.0</td>
<td>3.79</td>
<td>1.27 ± 0.03</td>
<td>89.0 ± 0.7</td>
<td>5.17 ± 0.01</td>
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<tr>
<td>106.0</td>
<td>3.60</td>
<td>1.25 ± 0.02</td>
<td>80 ± 2</td>
<td>6.84 ± 0.006</td>
<td></td>
</tr>
<tr>
<td>122.5</td>
<td>3.57</td>
<td>1.23 ± 0.05</td>
<td>81.24 ± 0.06</td>
<td>7.596 ± 0.006</td>
<td></td>
</tr>
<tr>
<td>167.0</td>
<td>3.43</td>
<td>1.17 ± 0.08</td>
<td>75 ± 1</td>
<td>8.5 ± 0.2</td>
<td></td>
</tr>
</tbody>
</table>

All the results obtained from transformed Kluuyveromyces, Torulaspora, Zygossaccharomyces and Sac-
charomyces yeasts are summarized and compared in Table 3. The yield is the amount of lactic acid produced (g/L) divided by the amount of glucose consumed (g/L). The percentage of free lactic acid is obtained from the Henderson-Hasselbalch equation:

\[ \text{pH} = \text{pK}_a + \log \left( \frac{[\text{Lactate}]}{[\text{Free Lactic Acid}]} \right) \]

where the pKₐ for lactic acid is 3.86.

Comparison of the data reported in Table 3A vs Table 3B clearly shows that in different yeasts genera, the production of lactic acid with higher yield on glucose can be obtained by changing the relative ratio—at cellular level—of the LDH and PDC activities. Such goal can be obtained following at least two studies approaches:

(1) by reducing the PDC activity (compare data from transformed K. lactis hosts: PM6-7A Vs PM6/CI; compare data from transformed S. cerevisiae hosts GRF18U Vs GRF18APDC2 and CENPK113 Vs CENPK113APDC1APDC5APDC6)

(2) by increasing the LDH gene copy number and therefore the LDH activity (compare data from S. cerevisiae host: GRF18U[pLC5] Vs GRF18U[pLC5][pLC7]; the LDH heterologous activity in the two strains is 5–6 and 7–6 U/mg of total cell proteins, respectively).

Further, higher yields can be obtained by manipulating the composition of the growth medium. Also in this case, a reduced ethanol production was observed (see also Table 4).

Table 3A

<table>
<thead>
<tr>
<th>Phosphate Buffer</th>
<th>Lactic Acid (g/L)</th>
<th>Yield (g/g)</th>
<th>Final pH</th>
<th>% Free Lactic Acid</th>
</tr>
</thead>
<tbody>
<tr>
<td>PM6-7A (negative control)</td>
<td>0.0</td>
<td>0.000</td>
<td>2.5</td>
<td>00</td>
</tr>
<tr>
<td>PM6-7A[pEPL2] (Fig. 7)</td>
<td>1.2</td>
<td>0.024</td>
<td>2.0</td>
<td>99</td>
</tr>
<tr>
<td>PM6-7A[pEPL2] (Fig. 7)</td>
<td>4.3</td>
<td>0.087</td>
<td>3.0</td>
<td>88</td>
</tr>
<tr>
<td>PM6-7A[pEPL4] (Fig. 7)</td>
<td>1.1</td>
<td>0.022</td>
<td>2.1</td>
<td>99</td>
</tr>
<tr>
<td>pM6-7A[pEPL4] (Fig. 7)</td>
<td>4.5</td>
<td>0.090</td>
<td>3.0</td>
<td>88</td>
</tr>
</tbody>
</table>

Table 3B

<table>
<thead>
<tr>
<th>Phosphate Buffer</th>
<th>Lactic Acid (g/L)</th>
<th>Yield (g/g)</th>
<th>Final pH</th>
<th>% Free Lactic Acid</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kloeveryces yeasts</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GRF18U[pLC5]</td>
<td>8.297</td>
<td>0.165</td>
<td>2.0</td>
<td>88</td>
</tr>
<tr>
<td>GRF18U[pME2]</td>
<td>5.927</td>
<td>0.118</td>
<td>3.0</td>
<td>88</td>
</tr>
<tr>
<td>GRF18U[pB11]</td>
<td>0.320</td>
<td>0.06</td>
<td>3.1</td>
<td>87</td>
</tr>
<tr>
<td>GRF18U[pB11]</td>
<td>1.5</td>
<td>0.020</td>
<td>3.0</td>
<td>88</td>
</tr>
<tr>
<td>GRF18U[pB11]</td>
<td>1.8</td>
<td>0.030</td>
<td>3.0</td>
<td>88</td>
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<tr>
<td><strong>CENPK113[pLC5]</strong></td>
<td>29.5</td>
<td>0.338</td>
<td>3.0</td>
<td>88</td>
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<tr>
<td><strong>CENPK113[pLC5]</strong></td>
<td>29.5</td>
<td>0.338</td>
<td>3.0</td>
<td>88</td>
</tr>
<tr>
<td><strong>CENPK113[pLC5]</strong></td>
<td>29.5</td>
<td>0.338</td>
<td>3.0</td>
<td>88</td>
</tr>
</tbody>
</table>

Table 3A-continued

<table>
<thead>
<tr>
<th>Phosphate Buffer</th>
<th>Lactic Acid (g/L)</th>
<th>Yield (g/g)</th>
<th>Final pH</th>
<th>% Free Lactic Acid</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kloeveryces yeasts</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PM6-7A (negative control)</td>
<td>0.0</td>
<td>0.000</td>
<td>2.5</td>
<td>00</td>
</tr>
<tr>
<td>PM6-7A[pEPL2] (Fig. 7)</td>
<td>1.2</td>
<td>0.024</td>
<td>2.0</td>
<td>99</td>
</tr>
<tr>
<td>PM6-7A[pEPL2] (Fig. 7)</td>
<td>4.3</td>
<td>0.087</td>
<td>3.0</td>
<td>88</td>
</tr>
<tr>
<td>PM6-7A[pEPL4] (Fig. 7)</td>
<td>1.1</td>
<td>0.022</td>
<td>2.1</td>
<td>99</td>
</tr>
<tr>
<td>pM6-7A[pEPL4] (Fig. 7)</td>
<td>4.5</td>
<td>0.090</td>
<td>3.0</td>
<td>88</td>
</tr>
<tr>
<td>Torulaspora yeasts</td>
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<td></td>
</tr>
<tr>
<td>CBS817 (negative control)</td>
<td>0.0</td>
<td>0.000</td>
<td>2.9</td>
<td>00</td>
</tr>
<tr>
<td>CBS817[pLAT-ADH] (Fig. 10)</td>
<td>1.0</td>
<td>0.058</td>
<td>2.8</td>
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<td>ATCC60483 (negative control)</td>
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<td>0.000</td>
<td>2.5</td>
<td>00</td>
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<td>ATCC60483[pLAT-ADH] (Fig. 11)</td>
<td>1.2</td>
<td>0.029</td>
<td>2.4</td>
<td>96</td>
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<tr>
<td>ATCC60497 (negative control)</td>
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<td>0.000</td>
<td>2.5</td>
<td>00</td>
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<tr>
<td>ATCC60497[pLAT-ADH]</td>
<td>0.9</td>
<td>0.018</td>
<td>2.4</td>
<td>95</td>
</tr>
<tr>
<td>Saccharomyces yeasts</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GRF18U (negative control)</td>
<td>0.0</td>
<td>0.000</td>
<td>3.1</td>
<td>00</td>
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<tr>
<td>GRF18U[pLAT-ADH]</td>
<td>2.1</td>
<td>0.040</td>
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**Experimental Data Related to the Saccharomyces GRF18U[pLC5][pLC7] Growing in a Manipulated Mineral Medium (Table 4).**

Lactic acid production by Saccharomyces GRF18U[pLC5][pLC7] transformed cells was also carried out growing the cells in a synthetic medium (D. Porro et al., Development of a pH controlled fed-batch system for budding yeast. Res. in Microbiol., 142, 535–539, 1991). In the synthetic medium used, the source of Mg and Zn salts are MgSO₄ (5 mM) and ZnSO₄·7H₂O (0.02 mM), respectively. Production was tested in aerobic batch culture (glucose concentration 50 g/l) as above described for the other transformed Saccharomyces cells. It has been found that depletion of both MgSO₄ and ZnSO₄·7H₂O yielded higher yield and higher lactic acid productivities. In fact, these minerals could be required as cofactor for the enzymatic activities leading to ethanol production. Data are shown in Table 4.
Table 4. (+) L-lactic acid production by *Saccharomyces GRF18[pLC5][pLC7] transformed cells during batch growth in manipulated mineral media.

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>-Mg</th>
<th>-Zn</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lactic acid production, g/l</td>
<td>9.23</td>
<td>13.74</td>
<td>13.74</td>
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<tr>
<td>Yield, g/g</td>
<td>0.20</td>
<td>0.29</td>
<td>0.29</td>
</tr>
<tr>
<td>Productivity, g/l/hr</td>
<td>0.38</td>
<td>0.42</td>
<td>0.61</td>
</tr>
<tr>
<td>Ratio ethanol/lactic acid, mM/M</td>
<td>2.78</td>
<td>2.31</td>
<td>3.99</td>
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</tbody>
</table>

Legend:
Control: complete synthetic medium (Ros. in Microbiol., 142, 535–539, 1991; enclosed).
-Mg: identical to the control but with out MgSO4.
-Zn: identical to the control but with out ZnSO4, w 71.0.

For all the tests, the final pH value was lower than 3.0 and therefore the % of free lactic acid was higher than 88%. Lactic Acid Production by Yeast Cells Overexpressing the JEN1 Gene.

Better lactic acid productions and lower ethanol productions have been obtained by overexpression of the JEN1 gene, encoding for the lactate transporter.

GRF18U[pLC5] (i.e., negative control) and GRF18U[pLC5][pJEN1] have been grown in media containing 2% glucose, 0.67% YNB w/v and supplements (i.e., 100 mg/l leucine-histidine, and 100 mg/l leucine, respectively).

Cells were preincubated in the same test’s medium. Exponentially growing cells were inoculated in flask (300 ml volume) containing 100 ml of fresh medium. The flasks were incubated at 30°C in a shaking bath (Dubnoff, 150 rpm), and fermentation was monitored at regular intervals. Cell number concentration was determined with an electronic Coulter counter (Coulter Counter ZBI Coulter Electronics Harpenden, G B, Porro et al., Res. Microbiol. (1991) 142, 535–539), after sonication of the samples to avoid cellular aggregates (Sonicator Fisher 300, medium point, Power 35%, 10 seconds).

TABLE 5

<table>
<thead>
<tr>
<th></th>
<th>Lactate</th>
<th>Ethanol</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>g/l</td>
<td>g/l</td>
</tr>
<tr>
<td>GRF18U[pLC5]</td>
<td>3.33</td>
<td>4.39</td>
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<tr>
<td>GRF18U[pLC5][pJEN1]</td>
<td>6.06</td>
<td>4.23</td>
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</tbody>
</table>

Continuous Lactic Acid Production.
Continuous and stable productions of lactic acid have been obtained for more than 2 weeks by means of classical chemostat cultures (the continuous flow of fresh medium to the bioreactor supported specific growth rate ranging between 0.01 and 0.3 hr⁻¹) using both the transformed *K. lactis* PM6-7A [pEPL2], PM1/C[pEPL2] and the transformed *S. cerevisiae* GRF18U[pLC5][pLC7] strains.

FED-BATCH TESTS
Lactic Acid Production by PM1/C1[pEPL2] in a Stirred-tank Fermentor.
Lactic acid production by PM1/C1[pEPL2] was further tested by cultivation in a 1-liter stirred-tank fermentor containing 8 liters of nutrient medium (30 g dry solids/L, light cone steep water, A. E. Staley Manufacturing Co., Decatur, IL; 10 g/L Difco yeast extract, Difco, Detroit, Mich.; 200 mg/L adenine, 50 g/L glucose). The fermentor was kept at 30°C, agitated at 400 rpm, and aerated at 2 liters/min throughout. Antifoam (Antifoam 1520, Dow Corning Corp., Midland, Mich.) was added as needed to control foaming. Glucose was fed as needed to maintain a residual concentration in the fermentation medium of about 25–50 g/L. When controlled, the pH was maintained by automatic addition of 14.8 M ammonium hydroxide in water. Lactic acid production at acidic pH was tested as follows: (1) The fermentation pH was controlled at 4.5 throughout the fermentation. (2) The initial fermentation pH was controlled at 4.0 until 80 ml of 14.8 M ammonium hydroxide were added. Then pH control was discontinued. (3) The initial fermentation pH was 5.0 and no neutralizing agent was added during the fermentation. The results are shown in Table 6. The elapsed time was measured from the time of inoculation. Samples from the fermentation, obtained after removing cells by filtration, were analyzed for the presence of glucose and L(+)-lactic acid using a YSI Model 2700 Select Biochemistry Analyzer (Yellow Springs Instrument Co., Inc., Yellow Springs, Ohio). Ethanol, measured by gas chromatography, was not detected in any of the fermentations. Yield and % free lactic acid were calculated as previously described. Inocula for the fermentations were prepared by pre-culturing PM1/C[pEPL2] in 50 ml of minimum synthetic medium (1.3% w/v Yeast Nitrogen Base-aa (Difco, Detroit, Mich.), 200 mg/L adenine, 5 g/L ammonium sulfate, 50 g/L glucose) in a 250 ml baffled Erlenmeyer flasks for 30 hr at 30°C and 300 rpm in an incubator shaker (Model G-24, New Brunswick Scientific Co., Inc., Edison, N.J.).

Similar results were obtained using the bacterial LDH gene (plasmid pEPL4; data not shown).

TABLE 6

| Lactic Acid Production by *Kluyveromyces* PM1/C1[pEPL2] cells in a Fermentor. |
|-------------------------------------|--------|--------|--------|--------|--------|
| Elapsed Time (hr)                  | NH₄OH Added (M) | Acid (g/l) | Yield (g/g) | Final pH | Lactic Acid % Free |
| Case 1                             | 137    | 1.31   | 109     | 0.59   | 4.5    | 19     |
| Case 2                             | 97     | 0.14   | 35      | 0.44   | 3.0    | 88     |
| Case 3                             | 72     | 0      | 29      | 0.38   | 2.8    | 92     |

Lactic Acid Production by BM3-12D[pLAZ10] in a Stirred-tank Fermentor.
Lactic acid production by BM3-12D[pLAZ10] was further tested by cultivation in a 1 liter stirred-tank fermentor containing 0.8 liters of nutrient medium (6.7 g/NYB/Yeast Nitrogen Base — Difco, Detroit, Mich., 45 g/L glucose, 2% v/v ethanol, G418 200 mg/l). The fermentor was kept at 30°C, agitated at 400 rpm, and aerated at 0.8 liters/min throughout. Antifoam (Antifoam 1520, Dow Corning Corp., Midland, Mich.) was added as needed to control foaming. Transformed cells first used ethanol for the production of biomass (first 50 hrs of growth) and then transformed glucose to L(+)-lactic acid. The pH was maintained at 4.5 by automatic addition of 2 M KOH. Glucose was fed as needed to maintain a residual concentration in the fermentation medium of about 35–45 g/L.

The results are shown in FIG. 9 and Table 7 (Case 1). The elapsed time was measured from the time of inoculation.
Samples from the fermentation, obtained after removing cells by filtration, were analyzed for the presence of glucose, ethanol and L(+)-lactic acid using a standard enzymatic analysis as described in Porro et al. 1995 (suDra). After T=50 hr, ethanol was not detected in any of sample-test. Yield and % free lactic acid were calculated as previously described.

Inocula for the fermentations were prepared by preculturing BM3-12D[pLAZ10] in 50 mL of minimum synthetic medium (1.3% w/v Yeast Nitrogen Base—aa (Difco, Detroit, Mich.), 2% w/v ethanol, G418 200 mg/L) in a 250 mL baffled Erlenmeyer flasks for 40 hr at 30°C and 300 rpm in an incubator shaker (Model G-24, New Brunswick Scientific Co., Inc., Edison, N.J.).

In a different experiment (Table 7, Case 2), the initial fermentation pH was 5.4 and no neutralizing agent was added during the fermentation.

### Table 7

<table>
<thead>
<tr>
<th>Elapsed Time (hr)</th>
<th>Lactic Acid (g/L)</th>
<th>Yield (g/g)</th>
<th>Final pH</th>
<th>Lactic Acid</th>
</tr>
</thead>
<tbody>
<tr>
<td>Case 1 474</td>
<td>60.3</td>
<td>0.854</td>
<td>4.5</td>
<td>19</td>
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<tr>
<td>Case 2 498</td>
<td>32.3</td>
<td>0.881</td>
<td>3.6</td>
<td>65</td>
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**Sequence Listing**

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<220> FEATURE:
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<210> SEQ ID NO 2
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<400> SEQUENCE: 2

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SEQ ID NO 5
LENGTH: 59
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: Oligomer complementary to S. cerevisiae PDC1 and anneals to loxP-KanSRD-loxP cassette
SEQUENCE: 5

ttctacttcat aaacctacgg cacaataac cagccagtgaagc ttcgtaacgc

SEQ ID NO 6
LENGTH: 62
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: Antisense oligomer complementary to S. cerevisiae PDC1 and anneals to loxP-KanSRD-loxP cassette
SEQUENCE: 6

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tg

SEQ ID NO 7
LENGTH: 59
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: Oligomer complementary to S. cerevisiae PDC5 and anneals to loxP-KanSRD-loxP cassette
SEQUENCE: 7

atcaatctca aagaagaaa cacaaataaa taacaagaag caggtgaagc ttcgtaacgc

SEQ ID NO 8
LENGTH: 62
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: Antisense oligomer complementary to S. cerevisiae PDC5 and anneals to loxP-KanSRD-loxP cassette
SEQUENCE: 8

aataataaca aacggtgtaat ctgagtttt tataatata caggtggcca ctagggatct
tg

SEQ ID NO 9
LENGTH: 59
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: Oligomer complementary to S. cerevisiae PDC6 and anneals to loxP-KanSRD-loxP cassette
SEQUENCE: 9
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SEQ ID NO 10
LENGTH: 62
TYPE: DNA
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: Antisense oligomer complementary to S. cerevisiae PDC6 and anneals to loxP-KanSRD-loxP cassette
What is claimed is:
1. A yeast strain transformed with at least one copy of a gene coding for lactate dehydrogenase functionally linked to a promoter sequence allowing the expression of the gene in the yeast strain, wherein the yeast strain is of Torulaspora or Zygosaccharomyces.

2. The yeast strain of claim 1, wherein the transformed yeast strain lacks ethanol production ability or has a reduced ethanol production ability with respect to a wild-type yeast of the same strain which has been similarly transformed with the gene encoding lactate dehydrogenase.

* * * * *
UNITED STATES PATENT AND TRADEMARK OFFICE
CERTIFICATE OF CORRECTION

PATENT NO. : 6,429,006 B1
DATED : August 6, 2002
INVENTOR(S) : Danilo Porro, Michele Bianchi, Bianca Maria Ranzi, Laura Frontali, Marina Vai, Aaron Adrian Winkler and Lilia Alberghina

It is certified that error appears in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

Title page,
Item [73], correct the spelling of the assignee to the following:
-- A.E. Staley Manufacturing Co. --

Signed and Sealed this
Fifteenth Day of October, 2002

Austen:

[Signature]

JAMES E. ROGAN
Assesting Officer
Director of the United States Patent and Trademark Office