McLeod *et al. BMC Microbiology* 2011, **11**:145 http://www.biomedcentral.com/1471-2180/11/145



RESEARCH ARTICLE

Open Access

Global transcriptome response in *Lactobacillus* sakei during growth on ribose

Anette McLeod^{1,2*}, Lars Snipen², Kristine Naterstad¹ and Lars Axelsson¹

Abstract

Background: Lactobacillus sakei is valuable in the fermentation of meat products and exhibits properties that allow for better preservation of meat and fish. On these substrates, glucose and ribose are the main carbon sources available for growth. We used a whole-genome microarray based on the genome sequence of *L. sakei* strain 23K to investigate the global transcriptome response of three *L. sakei* strains when grown on ribose compared with glucose.

Results: The function of the common regulated genes was mostly related to carbohydrate metabolism and transport. Decreased transcription of genes encoding enzymes involved in glucose metabolism and the L-lactate dehydrogenase was observed, but most of the genes showing differential expression were up-regulated. Especially transcription of genes directly involved in ribose catabolism, the phosphoketolase pathway, and in alternative fates of pyruvate increased. Interestingly, the methylglyoxal synthase gene, which encodes an enzyme unique for *L. sakei* among lactobacilli, was up-regulated. Ribose catabolism seems closely linked with catabolism of nucleosides. The deoxyribonucleoside synthesis operon transcriptional regulator gene was strongly up-regulated, as well as two gene clusters involved in nucleoside catabolism. One of the clusters included a ribokinase gene. Moreover, *hprK* encoding the HPr kinase/phosphatase, which plays a major role in the regulation of carbon metabolism and sugar transport, was up-regulated, as were genes encoding the general PTS enzyme I and the mannose-specific enzyme II complex (EII^{man}). Putative catabolite-responsive element (*cre*) sites were found in proximity to the promoter of several genes and operons affected by the change of carbon source. This could indicate regulation by a catabolite control protein A (CcpA)-mediated carbon catabolite repression (CCR) mechanism, possibly with the EII^{man} being indirectly involved.

Conclusions: Our data shows that the ribose uptake and catabolic machinery in *L. sakei* is highly regulated at the transcription level. A global regulation mechanism seems to permit a fine tuning of the expression of enzymes that control efficient exploitation of available carbon sources.

Background

The Lactobacillus sakei species belongs to the lactic acid bacteria (LAB), a group of Gram-positive organisms with a low G+C content which produce lactic acid as the main end product of carbohydrate fermentation. This trait has, throughout history, made LAB suitable for production of food. Acidification suppresses the growth and survival of undesirable spoilage bacteria and human pathogens. L. sakei is naturally associated with the meat and fish environment, and is important in the meat industry where it is used as starter culture for sausage fermentation [1,2]. The bacterium shows great potential as a protective

culture and biopreservative to extend storage life and ensure microbial safety of meat and fish products [3-6]. The genome sequence of *L. sakei* strain 23K has revealed a metabolic repertoire which reflects the bacterium's adaption to meat products and the ability to flexibly use meat components [7]. Only a few carbohydrates are available in meat and fish, and *L. sakei* can utilize mainly glucose and ribose for growth, a utilization biased in favour of glucose [7-9]. The species has been observed as a transient member of the human gastrointestinal tract (GIT) [10,11], and ribose may be described as a commonly accessible carbon source in the gut environment [12]. Transit through the GIT of axenic mice gave mutant strains which grow faster on ribose compared with glucose [13].

Glucose is primarily transported and phosphorylated by the phosphoenolpyruvate (PEP)-dependent carbohydrate

Full list of author information is available at the end of the article



^{*} Correspondence: anette.mcleod@nofima.no

¹Nofima Mat AS, Norwegian Institute of Food, Fisheries and Aquaculture Research, Osloveien 1, Ås, NO-1430, Norway

phosphotransferase system (PTS). A phosphorylation cascade is driven from PEP through the general components enzyme I (EI) and the histidine protein (HPr), then via the mannose-specific enzyme II complex (EII^{man}) to the incoming sugar. Moreover, glucose is fermented through glycolysis leading to lactate [7,8,14]. Ribose transport and subsequent phosphorylation are induced by the ribose itself and mediated by a ribose transporter (RbsU), a Dribose pyranase (RbsD), and a ribokinase (RbsK) encoded by rbsUDK, respectively. These genes form an operon with rbsR which encodes the local repressor RbsR [15,16]. The phosphoketolase pathway (PKP) is used for pentose fermentation ending with lactate and other end products [8,17]. L. sakei also has the ability to catabolize arginine, which is abundant in meat, and to catabolize the nucleosides inosine and adenine, a property which is uncommon among lactobacilli [7,18].

By proteomics, we recently identified proteins involved in ribose catabolism and the PKP to be over-expressed during growth on ribose compared with glucose, while several glycolytic enzymes were less expressed. Moreover, also enzymes involved in pyruvate- and glycerol/glycerolipid metabolism were over-expressed on ribose [19]. Bacteria often use carbon catabolite repression (CCR) in order to control hierarchical utilization of different carbon sources. In low G+C content Gram-positive bacteria, the dominant CCR pathway is mediated by the three main components: (1) catabolite control protein A (CcpA) transcriptional regulator; (2) the histidine protein (HPr); and (3) catabolite-responsive element (cre) DNA sites located in proximity to catabolic genes and operons, which are bound by CcpA [20-23]. The HPr protein has diverse regulatory functions in carbon metabolism depending on its phosphorylation state. In response to high throughput through glycolysis, the enzyme is phosphorylated at Ser46 by HPr kinase/phosphorylase (HPrK/P). This gives P-Ser-HPr which can bind to CcpA and convert it into its DNAbinding-competent conformation. However, when the concentration of glycolytic intermediates drop, the HPrK/ P dephosphorylates P-Ser-HPr [20,22-24]. Under low glucose concentrations, HPr is phosphorylated by E1 of the PTS at His15 to give P-His-HPr, which has a catalytic function in the PTS and regulatory functions by phosphorylation of catabolic enzymes and transcriptional regulators with a PTS regulation domain (PRD). Several P-EIIBs also phosphorylate different types of non-PTS proteins and regulate their activities [20-22]. Evidence for regulatory processes resembling glucose repression was shown both during lactose utilization [25] and catabolism of arginine [26,27] in *L. sakei*. A *cre* site has been reported upstream of the rbs operon [28], thus CcpA could likely be acting on the rbs operon as well as other catabolic genes and operons in this bacterium.

In the present study, we use a microarray representing the *L. sakei* 23K genome and an additional set of sequenced *L. sakei* genes, to investigate the global transcriptome response of three *L. sakei* strains when grown on ribose compared with glucose. Moreover, we predict the frequency of *cre* sites presumed to be involved in CCR in the *L. sakei* 23K genome sequence. Our objective was to identify differentially expressed genes between growth on the two sugars, and to increase the understanding of how the primary metabolism is regulated.

Methods

Bacterial strains, media and growth conditions

L. sakei 23K is a plasmid-cured sausage isolate [29], and its complete genome sequence has been published [7]. L. sakei LS 25 is a commercial starter culture strain for salami sausage [30]. L. sakei MF1053 originates from fermented fish (Norwegian "rakfisk") [9]. The strains were maintained at -80°C in MRS broth (Oxoid) supplemented with 20% glycerol. Growth experiments were performed in a defined medium for lactobacilli [31] supplemented with 0.5% glucose (DMLG) or 0.5% ribose + 0.02% glucose (DMLRg) as described previously [19]. Samples were extracted at three different days from independent DMLG and DMLRg cultures from each strain grown at 30°C to mid-exponential phase (OD₆₀₀ = 0.5-0.6) for a total of three sample sets (parallels).

Microarrays

The microarrays used have been described by Nyquist et al. [32], and a description is available at http://migale.jouy. inra.fr/sakei/Supplement.html/. 70-mer oligonucleotide probes representing the *L. sakei* strain 23K genome and an additional set of sequenced *L. sakei* genes were printed in three copies onto epoxy glass slides (Corning).

RNA extraction

Total RNA extraction was performed using the RNeasy Protect Mini Prep Kit (Qiagen) as described by Rud et al. [33]. The concentration and purity of the total RNA was analysed using NanoDrop ND-1000 (NanoDrop Technologies), and the quality using Agilent 2100 Bioanalyzer (Agilent Technologies). Sample criteria for further use in the transcriptome analysis were A_{260}/A_{280} ratio superior to 1.9 and 23S/16S RNA ratio superior to 1.6.

cDNA synthesis, labeling, and hybridization

cDNA was synthesized and labeled with the Fairplay III Microarray Labeling Kit (Stratagene, Agilent Technologies) as described previously [34]. After labeling, unincorporated dyes were removed from the samples using the QIAQuick PCR purification kit (Qiagen). The following prehybridization, hybridization, washing, and

drying of the arrays were performed in a Tecan HS 400 Pro hybridization station (Tecan) as described by Nyquist et al. [32]. For studying the carbon effects, samples from DMLG and DMLRg were co-hybridized for each of the three strains. Separate hybridizations were performed for each strain on all three biological parallels. In order to remove potential biases associated with labelling and subsequent scanning, a replicate hybridization was performed for each strain for one of the three parallels, where the Cy3 and Cy5 dyes (GE Healthcare) used during cDNA synthesis were swapped. The hybridized arrays were scanned at wavelengths 532 nm (Cy3) and 635 nm (Cy5) with a Tecan scanner LS (Tecan). GenePix Pro 6.0 (Molecular Devices) was used for image analysis, and spots were excluded based on slide or morphology abnormalities.

Microarray data analysis

Downstream analysis was done by the Limma package http://www.bioconductor.org in the R computing environment http://www.r-project.org. Pre-processing and normalization followed a standard procedure using methods described by Smyth & Speed [35], and testing for differential expressed genes were done by using a linear mixed model as described by Smyth [36]. A mixed-model approach was chosen to adequately describe between-array variation and still utilize probe-replicates (three replicates of each probe in each array). An empirical Bayes smoothing of gene-wise variances was conducted according to Smyth et al. [37], and for each gene the p-value was adjusted to control the false discovery rate (FDR), hence all p-values displayed are FDR-adjusted (often referred to as q-values in the literature).

Validation of microarray data by qRT-PCR analysis

The microarray results were validated on selected regulated genes for the LS 25 strain by quantitative real-time reverse transcriptase PCR (qRT-PCR) performed as described previously [38]. Primers and probes (Additional file 1, Table S3) were designed using Primer Express 3.0 (Applied Biosystems). Relative gene expression was calculated by the ΔC_T method, using the DNA gyrase subunit alpha gene (*gyrA*) as the endogenous reference gene.

Microarray accession numbers

The microarray data have been deposited in the Array Express database http://www.ebi.ac.uk/arrayexpress/under the accession numbers A-MEXP-1166 (array design) and E-MEXP-2892 (experiment).

Sequence analysis

A prediction of *cre* sites in the *L. sakei* 23K genome sequence (GeneBank acc. no. CR936503.1), both strands, was performed based on the consensus sequence

TGWNANCGNTNWCA (W = A/T, N = A/T/G/C), confirmed in Gram-positive bacteria [39]. We made a search with the consensus sequence described by the regular expression T-G-[AT]-X-A-X-C-G-X-T-X-[AT]-C-A, allowing up to two mismatches in the conserved positions except for the two center position, highlighted in boldface. All computations were done in R http://www.r-project.org.

Results and Discussion

Selection of L. sakei strains and growth conditions

We have previously investigated *L. sakei* strain variation [9], and used proteomics to study the bacterium's primary metabolism [19], providing us with a basis for choosing strains with interesting differences for further studies. The starter culture strain LS 25 showed the fastest growth rates in a variety of media, and together with strain MF1053 from fish, it fermented the highest number of carbohydrates [9]. The LS 25 strain belongs to the L. sakei subsp. sakei, whereas the 23K and MF1053 strains belong to L. sakei subsp. carnosus [9,19]. By identification of differentially expressed proteins caused by the change of carbon source from glucose to ribose, LS 25 seemed to down-regulate the glycolytic pathway more efficiently than other strains during growth on ribose [19]. For these reasons, LS 25 and MF1053 were chosen in addition to 23K for which the microarray is based on. Nyquist et al. [32] recently investigated the genomes of various L. sakei strains compared to the sequenced strain 23K by comparative genome hybridization (CGH) using the same microarray as in the present study. A large part of the 23K genes belongs to a common gene pool invariant in the species, and the status for each gene on the array is known for all the three strains [32].

As glucose is the preferred sugar, *L. sakei* grows faster when glucose is utilized as the sole carbon source compared with ribose [8,9,15]. However, glucose stimulates ribose uptake and a possible co-metabolism of these two sugars present in meat and fish has been suggested, a possibility that give the organism an advantage in competition with other microbiota [15,16,40]. To obtain comparable 2-DE gels between samples issued from bacteria grown on the two carbohydrates in our recent proteomic analysis, growth on ribose was enhanced by adding small amounts of glucose [19]. For the present transcriptome analysis we therefore chose the same growth conditions.

Global gene expression patterns

A microarray representing the *L. sakei* 23K genome and an additional set of sequenced *L. sakei* genes was used for studying the effect of carbon source on the transcriptome of *L. sakei* strains 23K, MF1053 and LS 25. Genes displaying a significant differential expression with a \log_2 ratio > 0.5 or < -0.5 were classified into functional categories according to the *L. sakei* 23K genome database

http://migale.jouy.inra.fr/sakei/genome-server and are listed in Table 1. The 23K strain showed differential expression for 364 genes within these limits, MF1053 and LS 25 for 223 and 316 genes, respectively. Among these, 88, 47 and 82, respectively, were genes belonging to the category of genes of 'unknown' function. Eighty three genes, the expression of which varied depending on the carbon source, were common to the three strains, among which 52 were up-regulated and 31 down-regulated during growth on ribose (Figure 1). The function of these common regulated genes was mostly related to carbohydrate transport and metabolism (34 genes, Table 1). The reliability of the microarray results was assessed by qRT-PCR analysis using selected regulated genes in the LS 25 strain. As shown in Table S4 in the additional material (Additional file 1), the qRT-PCR results were in agreement with the data obtained by the microarrays.

Several of the up-regulated genes are located in operons, an organisation believed to provide the advantage of coordinated regulation. In addition, in order to discriminate genes induced by growth on ribose from those repressed by glucose (submitted to CCR mediated by CcpA), a search of the complete genome sequence of *L. sakei* 23K [7] was undertaken, with the aim to identify putative cre sites. The search revealed 1962 hits, most of which did not have any biological significance considering their unsuitable location in relation to promoters. Relief of CcpAmediated CCR likely occur for many of the up-regulated genes in the category of carbohydrate transport and metabolism. Putative cre sites were identified in their promoter region, as well as for some genes involved in nucleoside and amino acid transport and metabolism (Table 2). In the other gene categories, the presences of putative *cre* sites were rare. With regard to gene product, the L. sakei genome shares high level of conservation with Lactobacillus plantarum [7], and high similarity of catabolic operon organization. The role of CcpA in CCR in L. plantarum has been established, and was shown to mediate regulation of the pox genes encoding pyruvate oxidases [41,42]. During growth on ribose, *L. plantarum* induces a similar set of genes as observed in the present study, and putative *cre* sites were identified in the upstream region of several genes involved [33].

Ribose catabolism and PKP

Confirming its major role in ribose transport and utilization in $L.\ sakei$, and in agreement with previous findings [16], our microarray data revealed a strong up-regulation (Table 1; $\log_2 = 2.8$ -4.3) of rbsUDK. The genes encoding an additional putative carbohydrate kinase belonging to the ribokinase family and a putative phosphoribosyl isomerase, lsa0254 and lsa0255, respectively, previously suggested to be involved in catabolism of ribose in $L.\ sakei$ [7], were induced in all the strains (Table 1). Recent

CGH studies revealed that some L. sakei strains which were able to grow on ribose did not harbour the rbsK gene, whereas lsa0254 was present in all strains investigated [32]. This second ribokinase could therefore function as the main ribokinase in some *L. sakei* strains. The rbsK sequence could also differ considerably from that of 23K in these strains. The PKP showed an obvious induction with an up-regulation (2.2-3.2) of the xpk gene encoding the key enzyme xylulose-5-phosphate phosphoketolase (Xpk). This enzyme connects the upper part of the PKP to the lower part of glycolysis by converting xylulose-5-phosphate into glyceraldehyde-3-phosphate and acetyl-phosphate. Acetyl-phosphate is then converted to acetate and ATP by acetate kinase (Ack). Supporting our results, previous proteomic analysis showed an over-expression of RbsK, RbsD and Xpk during growth on ribose [15,16,19]. The induction of ribose transport and phosphorylation, and increased phosphoketolase and acetate kinase activities were previously observed during growth on ribose [15]. Three genes encoding Ack are present in the 23K genome [7], as well as in MF1053 and LS 25 [32]. A preferential expression of different ack genes for the acetate kinase activity seem to exist. The ack2 gene was up-regulated in all the strains, while ack1 was up-regulated and ack3 downregulated in 23K and LS 25 (Table 1). An illustration of the metabolic pathways with genes affected by the change of carbon source from glucose to ribose in L. sakei is shown in Figure 2.

As a consequence of the pentose-induced PKP, genes involved in PKP-metabolism of glucose, such as gntZ, gntK and zwf, were down-regulated (Table 1, Figure 2). The glycolytic pathway was clearly repressed, supporting previous findings [15,19]. Among these genes were pfk (0.5-1.1) encoding 6-phosphofructokinase (Pfk), and fba (0.7-1.1) coding for fructose-bisphosphate aldolase, both acting at the initial steps of glycolysis. In addition, gpm3 encoding one of the five phosphoglycerate mutases present in the 23K genome, acting in the lower part of glycolysis, was also down-regulated (0.7-0.9). MF1053 down-regulated pyk (0.7) encoding pyruvate kinase (Pyk) that competes for PEP with the PTS (Figure 2). Its activity results in the production of pyruvate and ATP, and it is of major importance in glycolysis and energy production in the cell. MF1053 also showed a stronger downregulation of pfk than the other strains (Table 1). Similar to several other lactobacilli, pfk is transcribed together with pyk [43,44], and in many microorganisms the glycolytic flux depends on the activity of the two enzymes encoded from this operon [43,45]. At the protein level, we previously observed both Pfk and Pyk expressed at a lower level for all the three strains [19], however this was not confirmed at the level of gene expression for 23K and LS 25. We could also not confirm the lower

Table 1 Genes with significant differential expression in three *L. sakei* strains grown on ribose compared with glucose, FDR adjusted p-value less than 0.01 and \log_2 of > 0.5 or < -0.5 (\log_2 values > 1.0 or < -1.0 are shown in bold)

Gene locus	Gene	Description	23K	MF1053	LS 25
Carbohydı	ate transpo	ort and metabolism			
Transport/	binding of	carbohydrates			
LSA0185*	galP	Galactose:cation symporter	1.2		1.7
LSA0200*	rbsU	Ribose transport protein	2.8	3.5	4.3
LSA0353*	lsa0353	Putative cellobiose-specific PTS, enzyme IIB	3.6	1.3	2.5
LSA0449*	manL	Mannose-specific PTS, enzyme IIAB	2.1	2.5	1.5
LSA0450*	manN	Mannose-specific PTS, enzyme IIC	1.9	2.0	1.4
LSA0451*	manM	Mannose-specific PTS, enzyme IID	2.4	1.0	2.1
LSA0651*	glpF	Glycerol uptake facilitator protein, MIP family	3.4	4.7	3.4
LSA1050*	fruA	Fructose-specific PTS, enzyme IIABC			0.9
LSA1204*	lsa1204	Putative sugar transporter		1.1	
LSA1457*	lsa1457	Putative cellobiose-specific PTS, enzyme IIC		2.3	
LSA1462*	ptsl	PTS, enzyme I	0.8	1.7	0.9
LSA1463*	ptsH	Phosphocarrier protein HPr (histidine protein)		1.2	0.9
LSA1533	lsa1533	Putative cellobiose-specific PTS, enzyme IIA		2.5	2.1
LSA1690	lsa1690	Putative cellobiose-specific PTS, enzyme IIC	0.9		
LSA1792*	scrA	Sucrose-specific PTS, enzyme IIBCA	0.8		1.1
Metabolisı	m of carbo	hydrates and related molecules			
LSA0123*	lsa0123	Putative sugar kinase, ROK family	1.2		
LSA0198	ack1	Acetate kinase (acetokinase)	1.7		1.3
LSA0254*	lsa0254	Putative carbohydrate kinase	2.4	0.8	1.8
LSA0292*	budC	Acetoin reductase (acetoin dehydrogenase) (meso-2,3-butanediol dehydrogenase)	3.4	2.3	3.4
LSA0444	lsa0444	Putative malate dehydrogenase	3.4	D	2.1
LSA0516	hprK	Hpr kinase/phosphorylase	2.0	1.6	1.2
LSA0664*	loxL1N	L-lactate oxidase (N-terminal fragment), degenerate	1.2		0.7
LSA0665*	loxLl	L-lactate oxidase (central fragment), degenerate	1.0		
LSA0666*	loxL1C	L-lactate oxidase (C-terminal fragment), degenerate	1.0		
LSA0974*	pflB	Formate C-acetyltransferase (pyruvate formate-lyase) (formate acetyltransferase)	4.0		
LSA0981	aldB	Acetolactate decarboxylase (alpha-acetolactate decarboxylase)		0.6	1.9
LSA0982	als	Acetolactate synthase (alpha-acetolactate synthase)			1.9
LSA0983	lsa0983	Putative aldose-1 epimerase	0.6		
LSA1032	pyk	Pyruvate kinase		-0.7	
LSA1080	lsa1080	Myo-inositol monophosphatase	0.6		0.8
LSA1082	pdhD	Pyruvate dehydrogenase complex, E3 component, dihydrolipoamide dehydrogenase	2.8	2.5	2.1
LSA1083	pdhC	Puruvate dehydrogenase complex, E2 component, dihydrolipoamide acetyltransferase	3.4	3.7	2.7
LSA1084	pdhB	Pyruvate dehydrogenase complex, E1 component, beta subunit	3.2	3.3	2.2
LSA1085	pdhA	Pyruvate dehydrogenase complex, E1 component, alpha subunit	2.9	3.5	2.4
LSA1141*	ррдК	Pyruvate phosphate dikinase	1.0		0.9
LSA1188*	pox1	Pyruvate oxidase	2.3	3.1	2.1
LSA1298	ack2	Acetate kinase (acetokinase)	1.1	0.9	0.9
LSA1343*	eutD	Phosphate acetyltransferase (phosphotransacetylase)	2.0	1.0	1.6
LSA1381	Isa1381	Putative acylphosphatase	-0.6	-0.5	110
LSA1399*	loxL2	L-lactate oxidase	3.4	U	
LSA1630	Isa1630	Putative sugar kinase, ROK family	-0.6	J	-0.6
LSA1630*	nanA	N-acetylneuraminate lyase	2.0		-0.0 D
LSA1641*	nanE	N-acylglucosamine/mannosamine-6-phosphate 2-epimerase	0.9		D
			0.9 1.8		D
LSA1643*	lsa1643	Putative sugar kinase, ROK family			1 1
LSA1668	ack3	Acetate kinase (acetokinase) Pyruvate oxidase	-0.7 0.7		-1.1

Table 1 Genes with significant differential expression in three *L. sakei* strains grown on ribose compared with glucose, FDR adjusted p-value less than 0.01 and \log_2 of > 0.5 or < -0.5 (\log_2 values > 1.0 or < -1.0 are shown in bold) (Continued)

Intermedia		liem			
Intermedia LSA0255*	lsa0255	Putative phosphoribosyl isomerase	2.0	1.0	1.6
		e metabolic pathway	2.0	1.0	1.0
LSA0201*	rbsD	D-ribose pyranase	2.5	2.5	3.4
LSA0201*	rbsK	Ribokinase	3.0	3.9	4.3
LSA0202 LSA0289*	xpk	Xylulose-5-phosphate phosphoketolase	3.2	2.3	2.6
LSA0289	gntZ	6-phosphogluconate dehydrogenase	-1.2	-0.9	-1.7
LSA0297	gntK	Gluconokinase	-0.8	-0.9	-1.7
LSA0381	zwf	Glucose-6-phosphate 1-dehydrogenase	-0.6	-0.6	-0.6
LSA0649*	glpK	Glycerol kinase	3.4	4.8	2.1
LSA0650*	glpD glpD	Glycerol-3-phosphate dehydrogenase	2.3	2.2	2.0
LSA0764*	galK	Galactokinase	1.1	0.7	1.8
LSA0765*	galE1	UDP-glucose 4-epimerase	1.1	0.7	1.2
LSA0765*	galT	Galactose-1-phosphate uridylyltransferase	1.2	0.8	2.0
LSA0760*	galM	Aldose 1-epimerase (mutarotase)	1.3	0.0	2.0
LSA0707 LSA1146*	manA	Mannose-6-phosphate isomerase	1.4	1.3	1.5
LSA1140	Isa1531	Putative beta-glucosidase	1.4	0.7	0.9
LSA1581	nagA	N-acetylglucosamine-6-phosphate deacetylase	0.6	0.7	0.5
LSA1685	rpiA	Ribose 5-phosphate epimerase (ribose 5-phosphate isomerase)	0.0	1.1	0.8
LSA1710*	lacM	Beta-galactosidase, small subunit (lactase, small subunit)	3.3		1.2
LSA1711*	lacL	Beta-galactosidase, large subunit (lactase, large subunit)	3.0	1.5	1.7
LSA1790*	scrK	Fructokinase	5.0	1.0	1.1
LSA1791*	dexB	Glucan 1,6-alpha-glucosidase (dextran glucosidase)		1.0	1.1
LSA1795	melA	Alpha-galactosidase (melibiase)			-0.6
Glycolytic		April guidetosidase (meilolase)			0.0
LSA0131	gpm2	Phosphoglycerate mutase		0.7	
LSA0206	gpm3	Phosphoglycerate mutase	-0.7	-0.8	-0.9
LSA0609*	gloAC	Lactoylglutathione lyase (C-terminal fragment), authentic frameshift	1.1	0.0	0.7
LSA0803	gpm4	Phosphoglycerate mutase	0.5		0.5
LSA1033	pfk	6-phosphofructokinase	-0.6	-1.1	-0.5
LSA1157	mgsA	Methylglyoxal synthase	2.3	1.4	1.7
LSA1179	pgi	Glucose-6-phosphate isomerase	0.5		•••
LSA1527	fba	Fructose-bisphosphate aldolase	-1.0	-0.7	-1.1
LSA1606	ldhL	L-lactate dehydrogenase	-1.0	-0.9	-1.5
25/11000	IGITE	E loctate dell'yarogenide		0.5	1.5
Nucleotide	transport	and metabolism			
Transport/	binding of	nucleosides, nucleotides, purines and pyrimidines			
LSA0013	Isa0013	Putative nucleobase:cation symporter	-0.9		-1.5
LSA0055	Isa0055	Putative thiamine/thiamine precursor:cation symporter			1.6
LSA0064	Isa0064	Putative nucleobase:cation symporter		-0.8	
LSA0259	lsa0259	Pyrimidine-specific nucleoside symporter	1.5		1.3
LSA0798*	Isa0798	Pyrimidine-specific nucleoside symporter	3.5	2.2	1.7
LSA0799*	lsa0799	Putative purine transport protein	4.4	2.7	2.9
LSA1210	lsa1210	Putative cytosine:cation symporter (C-terminal fragment), authentic frameshift	-0.8		-0.6
LSA1211	lsa1211	Putative cytosine:cation symporter (N-terminal fragment), authentic frameshit	-1.1		-0.9
Metabolisr	n of nucle	otides and nucleic acids			
LSA0010	Isa0010	Putative nucleotide-binding phosphoesterase			-0.6
LSA0023	Isa0023	Putative ribonucleotide reductase (Nrdl-like)	-0.5	D	D
LSA0063	purA	Adenylosuccinate synthetase (IMP-aspartate ligase)		-0.8	
LSA0139	guaA	Guanosine monophosphate synthase (glutamine amidotransferase)		-0.5	-0.8

Table 1 Genes with significant differential expression in three *L. sakei* strains grown on ribose compared with glucose, FDR adjusted p-value less than 0.01 and \log_2 of > 0.5 or < -0.5 (\log_2 values > 1.0 or < -1.0 are shown in bold) (Continued)

(Continue	a)				
LSA0252	iunH1	Inosine-uridine preferring nucleoside hydrolase	2.6	2.6	1.8
LSA0446	pyrDB	Putative dihydroorotate oxidase, catalytic subunit			0.9
LSA0489	Isa0489	Putative metal-dependent phosphohydrolase precursor	0.5		
_SA0533*	iunH2	Inosine-uridine preferring nucleoside hydrolase	1.2		
_SA0785	Isa0785	Putative NCAIR mutase, PurE-related protein	-2.3		-1.3
_SA0795*	deoC	2 Deoxyribose-5 phosphate aldolase	4.0	2.1	2.2
_SA0796*	deoB	Phosphopentomutase (phosphodeoxyribomutase)	5.5	4.1	3.2
_SA0797*	deoD	Purine-nucleoside phosphorylase	4.5	2.6	1.9
_SA0801*	pdp	Pyrimidine-nucleoside phosphorylase	1.8		
SA0940	nrdF	Ribonucleoside-diphosphate reductase, beta chain		1.0	0.6
SA0941	nrdE	Ribonucleoside-diphosphate reductase, alpha chain		1.0	0.6
SA0942	nrdH	Ribonucleotide reductase, NrdH-redoxin		1.1	
.SA0950	pyrR	Bifunctional protein: uracil phosphoribosyltransferase and pyrimidine operon transcriptional regulator	-0.6		
SA0993	rnhB	Ribonuclease HII (RNase HII)			0.6
SA1018	cmk	Cytidylate kinase			0.6
SA1097	lsa1097	Putative ADP-ribose phosphorylase, NUDIX family	0.5		
SA1352	lsa1352	Putative phosphomethylpyrimidine kinase	-0.8		
SA1651	lsa1651	Putative purine phosphoribosyltransferase, PRT family		0.8	
SA1661	lsa1661	Putative nucleotide hydrolase, NUDIX family		-0.5	
SA1805	dgk	Deoxyguanosine kinase	-1.0		-0.8
「ranscripti					
ranscripti	ion regulat				
.SA0130	Isa0130	Putative transcriptional regulator, LacI family	-0.6		
SA0132	lsa0132	Putative transcriptional regulator, MarR family	-0.6		
SA0161	lsa0161	Putative transcriptional regulator, ArsR family	-0.6		
SA0186	lsa0186	Putative transcriptional regulator, LytR family		0.8	0.6
SA0203	rbsR	Ribose operon transcriptional regulator, Lacl family	1.7		
SA0217	lsa0217	Putative thiosulfate sulfurtransferase with a ArsR-HTH domain, rhodanese family		-1.0	-0.7
SA0229	lsa0229	Putative transcriptional regulator, MerR family (N-terminal fragment), authentic frameshift	-0.5		
SA0269	Isa0269	Putative transcriptional regulator, TetR family			-0.6
SA0293	Isa0293	Putative DNA-binding protein, XRE family			-0.6
SA0356	rex1	Redox-sensing transcriptional repressor, Rex	-0.8	-0.5	-0.9
SA0603	cggR	Glycolytic genes regulator		-0.6	-0.6
SA0669	Isa0669	Putative transcription regulator, TetR family		-0.6	
SA0783	Isa0783	Putative transcriptional regulator, Fnr/Crp Family	-0.6		
_SA0800	deoR	Deoxyribonucleoside synthesis operon transcriptional regulator, GntR family	3.8	2.1	1.9
SA0835	Isa0835	Putative DNA-binding protein, XRE family	-0.6		
SA0848	rex	Redox-sensing transcriptional repressor, Rex	1.6	0.7	
SA0972	lsa0972	Putative transcriptional regulator, LysR family	0.9		
SA1201	lsa1201	Putative transcriptional regulator, GntR family	1.4	D	D
SA1322	glnR	Glutamine synthetase transcriptional regulator, MerR family	-1.4	-1.3	
SA1351	lsa1351	Putative transcritional regulator with aminotransferase domain, GntR family		-0.5	-0.6
SA1434	lsa1434	Putative transcriptional regulator, DUF24 family (related to MarR/PadR families)	-0.8		
SA1449	spxA	Transcriptional regulator Spx	1.0		0.6
SA1521	lsa1521	Putative transcriptional regulator, TetR family	0.6		
SA1554	lsa1554	Putative transcriptional regulator, LacI family	-0.7	-0.9	-0.5
SA1587	lsa1587	Putative transcriptional regulator, GntR family	0.6		
		· · · · · · · · · · · · · · · · · · ·			
SA1611	lsa1611	Putative DNA-binding protein, PemK family		-0.5	-0.7

Table 1 Genes with significant differential expression in three *L. sakei* strains grown on ribose compared with glucose, FDR adjusted p-value less than 0.01 and \log_2 of > 0.5 or < -0.5 (\log_2 values > 1.0 or < -1.0 are shown in bold) (Continued)

of coenzy panE thiE	and metabolism ormes and prostethic groups			
panE thiE				
thiE	2 dabydrapantosta 2 radystaca			
	2-dehydropantoate 2-reductase		0.8	
41-10	Thiamine-phosphate pyrophosphorylase (thiamine-phosphate synthase)			1.9
thiD	Phosphomethylpyrimidine kinase (HMP-phosphate kinase)			1.4
thiM	Hydroxyethylthiazole kinase (4-methyl-5-beta-hydroxyethylthiazole kinase)	1.0		1.8
lsa0183	Putative hydrolase, isochorismatase/nicotamidase family	-0.7		
Isa0840	Putative glutamate-cysteine ligase	0.6		
fhs	Formate-tetrahydrofolate ligase (formyltetrahydrofolate synthetase)	0.6		
lsa0980	Putative hydroxymethylpyrimidine/phosphomethylpyrimidine kinase, PfkB family	0.6		
folK	2-amino-4-hydroxy-6-hydroxymethyldihydropteridine pyrophosphokinase	0.6	U	
acpS	Holo-[acyl-carrier protein] synthase (holo-ACP synthase) (4'-phosphopantetheine transferase AcpS)	-1.0	-0.9	-0.9
lsa1664	Putative dihydrofolate reductase	1.6	1.1	1.5
atpC		0.6		
atpD				0.6
atpG				0.8
atpA				0.6
atpH				0.6
atpF				0.5
atpE	H(+)-transporting two-sector ATPase (ATP synthase), C subunit			0.7
-	-			
				-0.7
	•			-0.6
				-1.0
				-0.6
	·	-0.9		-1.3
	· · · · · · · · · · · · · · · · · · ·			-0.5
				-0.6
		1.1		
			0.7	0.5
		0.5		0.5
				0.6
	· · · · · · · · · · · · · · · · · · ·			
'				
	· · · · · · · · · · · · · · · · · · ·			
	· · · · · · · · · · · · · · · · · · ·			
	·			
		0.6	1.0	0.0
	-	4.4	-1.0	-0.8
lsa1645	Putative Na(+)/(+) antiporter	1.4		D
	folk acpS sa1664 uction an bioenerge atpC atpD atpB atpH atpH atpF atpE	Colic 2-amino-4-hydroxy-6-hydroxymethyldihydropteridine pyrophosphokinase Holo-[acyl-carrier protein] synthase (holo-ACP synthase) (4'-phosphopantetheine transferase AcpS) Putative dihydrofolate reductase uction and conversion Dioenergetics (ATP synthase) ATPC H(+)-transporting two-sector ATPase (ATP synthase), epsilon subunit ATPC H(+)-transporting two-sector ATPase (ATP synthase), beta subunit ATPC H(+)-transporting two-sector ATPase (ATP synthase), alpha subunit ATPC H(+)-transporting two-sector ATPase (ATP synthase), alpha subunit ATPC H(+)-transporting two-sector ATPase (ATP synthase), delta subunit ATPC H(+)-transporting two-sector ATPase (ATP synthase), B subunit ATPC H(+)-transporting two-sector ATPase (ATP synthase), C subunit H(+)-transport and metabolism Holing of inorganic ions H(+)-transporter and H(+)-transporter, HP-binding subunit HIST HIST H(+)-transporter, ATP-binding subunit HIST H(+)-transporter, HP-binding subunit HIST H(+)-transporter, H	Column C	Column C

Table 1 Genes with significant differential expression in three *L. sakei* strains grown on ribose compared with glucose, FDR adjusted p-value less than 0.01 and \log_2 of > 0.5 or < -0.5 (\log_2 values > 1.0 or < -1.0 are shown in bold) (Continued)

Continue	:u)				
LSA1703	lsa1703	Putative Na(+)/H(+) antiporter	-1.2		
LSA1704	Isa1704	Putative calcium-transporting P-type ATPase			-0.8
LSA1735	Isa1735	Putative cobalt ABC transporter, membrane-spanning subunit			-0.6
LSA1736	Isa1736	Putative cobalt ABC transporter, ATP-binding subunit	-0.6		
LSA1737	Isa1737	Putative cobalt ABC transporter, ATP-binding subunit	-0.7		
LSA1838	Isa1838	Putative metal ion ABC transporter, membrane-spanning subunit			-0.5
LSA1839	lsa1839	Putative metal ion ABC transporter, substrate-binding lipoprotein precursor			-0.6
	-	t and metabolism			
	_	amino acids			
LSA0125	Isa0125	Putative amino acid/polyamine transport protein	0.6		
LSA0189	Isa0189	Putative amino acid/polyamine transport protein			-0.7
LSA0311	Isa0311	Putative glutamate/aspartate:cation symporter	-1.1		-1.0
LSA1037	Isa1037	Putative amino acid/polyamine transport protein	1.0	0.8	0.5
LSA1219	lsa1219	Putative cationic amino acid transport protein	0.7		
LSA1415	Isa1415	Putative amino acid/polyamine transport protein	1.1		0.7
LSA1424	lsa1424	Putative L-aspartate transport protein	-1.4	-0.9	-1.2
LSA1435	lsa1435	Putative amino acid:H(+) symporter	1.0		0.8
LSA1496	lsa1496	Putative glutamine/glutamate ABC transporter, ATP-binding subunit		1.2	
LSA1497	lsa1497	Putative glutamine/glutamate ABC transporter, membrane-spanning/substrate-binding subunit precursor		0.7	
Transport	binding of	proteins/peptides			
LSA0702	оррА	Oligopeptide ABC transporter, substrate-binding lipoprotein precursor		1.3	1.0
LSA0703	оррВ	Oligopeptide ABC transporter, membrane-spanning subunit		0.8	0.8
LSA0704	оррС	Oligopeptide ABC transporter, membrane-spanning subunit		1.8	1.0
LSA0705	oppD	Oligopeptide ABC transporter, ATP-binding subunit		1.2	1.1
LSA0706	оррҒ	Oligopeptide ABC transporter, ATP-binding subunit		1.2	1.2
Protein fa	te				
LSA0053	рерО	Endopeptidase O	0.6		
LSA0133	pepR	Prolyl aminopeptidase	1.5		
LSA0226	pepN	Aminopeptidase N (lysyl-aminopeptidase-alanyl aminopeptidase)			-0.7
LSA0285	pepF1	Oligoendopeptidase F1			-0.7
LSA0320	pepD3	Dipeptidase D-type (U34 family)		-0.8	-0.5
LSA0424	pepV	Xaa-His dipeptidase V (carnosinase)	1.6		
LSA0643	рерХ	X-Prolyl dipeptidyl-aminopeptidase	0.6		
LSA0888	рерТ	Tripeptide aminopeptidase T	0.6		
LSA1522	pepS	Aminopeptidase S	0.5		
LSA1686	pepC1N	Cysteine aminopeptidase C1 (bleomycin hydrolase) (N-terminal fragment), authentic frameshift		1.6	
LSA1688	pepC2	Cysteine aminopeptidase C2 (bleomycin hydrolase)		0.7	
LSA1689	lsa1689	Putative peptidase M20 family	1.0		1.1
Metabolis	m of amino	acids and related molecules			
LSA0220_c	: dapE	Succinyl-diaminopimelate desuccinylase	-1.4		-1.5
LSA0316	sdhB	L-serine dehydratase, beta subunit (L-serine deaminase)	-0.7		
LSA0370*	arcA	Arginine deiminase (arginine dihydrolase)	1.9		
LSA0372*	arcC	Carbamate kinase	0.5		
LSA0463	Isa0463	Putative 2-hydroxyacid dehydrogenase	-0.7		
LSA0509	kbl	2-amino-3-ketobutyrate coenzyme A ligase (glycine acetyltransferase)	1.5		
LSA0510	Isa0510	L-threonine dehydrogenase (N-terminal fragment), authentic frameshift	2.0	0.5	
LSA0572*	tdcB	Threonine deaminase (threonine ammonia-lyase, threonine dehydratase, IlvA homolog)	2.2		1.7
LSA0922	serA	D-3-phosphoglycerate dehydrogenase	0.9		
LSA1134	glyA	Glycine/Serine hydroxymethyltransferase		0.7	

Table 1 Genes with significant differential expression in three *L. sakei* strains grown on ribose compared with glucose, FDR adjusted p-value less than 0.01 and \log_2 of > 0.5 or < -0.5 (\log_2 values > 1.0 or < -1.0 are shown in bold) (Continued)

LSA1321	glnA	Glutamate-ammonia ligase (glutamine synthetase)	-1.3	-1.0	
LSA1484	mvaS	Hydroxymethylglutaryl-CoA synthase	-0.7	-0.6	-0.7
LSA1693	asnA2	L-asparaginase	0.8		
Lipid trans	sport and m	netabolism			
	m of lipids				
LSA0045	cfa	Cyclopropane-fatty-acyl-phospholipid synthase	-1.3	-1.4	-1.4
LSA0644	Isa0644	Putative acyl-CoA thioester hydrolase	0.6		
LSA0812	fabZ1	(3R)-hydroxymyristoyl-[acyl-carrier protein] dehydratase		-0.7	0.5
LSA0813	fabH	3-oxoacyl-[acyl carrier protein] synthetase III			0.6
LSA0814	асрР	Acyl carrier protein			0.6
LSA0815	fabD	Malonyl-CoA:ACP transacylase		-0.7	0.7
LSA0816	fabG	3-oxoacyl-acyl carrier protein reductase		-0.7	
LSA0817	fabF	3-oxoacyl-[acyl carrier protein] synthetase II		-0.7	
LSA0819	fabZ	(3R)-hydroxymyristoyl-[acyl carrier proetin] dehydratase			0.7
LSA0820	accC	Acetyl-CoA carboxylase (biotin carbooxylase subunit)		-0.7	
LSA0821	accD	Acetyl-CoA carboxylase (carboxyl transferase beta subunit)			0.8
LSA0822	ассА	Acetyl-CoA carboxylase (carboxyl transferase alpha subunit)			0.6
LSA0823	fabl	Enoyl [acyl carrier protein] reductase			0.9
LSA0891	lsa0891	Putative lipase/esterase	1.2		
LSA1485	mvaA	Hydroxymethylglutaryl-CoA reductase	-0.5		
LSA1493	lsa1493	Putative diacylglycerol kinase	-0.6	-0.9	-0.7
LSA1652	ipk	4-diphosphocytidyl-2-C-methyl-D-erythritol kinase	-0.6		-0.7
Secondary	metabolite	s transport and metabolism			
Transport/	binding pro	oteins and lipoproteins			
LSA0046	Isa0046	Putative transport protein	-1.0	-0.6	-1.3
LSA0089	Isa0089	Putative drug transport protein	-2.1	-0.9	-0.8
LSA0094	Isa0094	Putative transport protein, Major Facilitator Super (MFS) family transporter	-0.7		-0.7
LSA0095	Isa0095	Putative transport protein	1.3	0.5	
LSA0128	lsa0128	Putative antimicrobial peptide ABC exporter, membrane-spanning/permease subunit			-0.5
LSA0187	lsa0187	Putative drug-resistance ABC transporter, two ATP-binding subunits		0.7	
LSA0219_b	lsa0219_b	Putative cyanate transport protein	-0.6		
LSA0232	lmrA	Multidrug ABC exporter, ATP-binding and membrane-spanning/permease subunits	-0.7		-0.7
LSA0270	Isa0270	Putative multidrug ABC exporter, membrane-spanning/permease subunit	-0.7		
LSA0271	Isa0271	Putative multidrug ABC exporter, ATP-binding subunit	-0.7		-0.6
LSA0272	Isa0272	Putative multidrug ABC exporter, ATP-binding and membrane-spanning/permease subunits	-0.6		-0.6
LSA0308	Isa0308	Putative drug:H(+) antiporter			-0.7
LSA0376	Isa0376	Putative transport protein	0.7		
LSA0420	Isa0420	Putative drug:H(+) antiporter (N-terminal fragment), authentic frameshift	-0.8		-1.1
LSA0469	Isa0469	Putative drug:H(+) antiporter	-0.6		-0.5
LSA0788	Isa0788	Putative facilitator protein, MIP family	-2.6		
LSA0936	Isa0936	Putative drug ABC exporter, membrane-spanning/permease subunit	1.1		
LSA0937	Isa0937	Putative drug ABC exporter, membrane-spanning/permease subunit	1.3		
LSA0938	Isa0938	Putative drug ABC exporter, ATP-binding subunit	1.2		
LSA0963	Isa0963	Integral membrane protein, hemolysin III related			
LSA1088	lsa1088	Putative multidrug ABC exporter, ATP-binding and membrane-spanning/permease subunits	0.5		
LSA1261	lsa1261	Putative autotransport protein	0.5		
LSA1340	lsa1340	Putative transport protein		-0.7	
LSA1366	lsa1366	Putative ABC exporter, ATP-binding subunit	-0.8		-1.0

Table 1 Genes with significant differential expression in three *L. sakei* strains grown on ribose compared with glucose, FDR adjusted p-value less than 0.01 and \log_2 of > 0.5 or < -0.5 (\log_2 values > 1.0 or < -1.0 are shown in bold) (Continued)

(Continue	ea)				
LSA1367	lsa1367	Putative ABC exporter, membrane-spanning/permease subunit	-0.8	-0.5	-0.8
LSA1420	Isa1417	Putative lipase/esterase		-1.1	
LSA1621	lsa1621	Putative drug:H(+) antiporter		-1.1	
LSA1642	Isa1642	Putative Solute:Na(+) symporter	3.4	1.8	D
LSA1872	lsa1872	Putative drug:H(+) antiporter		0.7	
LSA1878	lsa1878	Putative drug resistance ABC transporter, two ATP-binding subunits	-0.6		
Detoxifica	ation				
LSA0772	Isa0772	Hypothetical protein (TelA, telluric resistance family)	1.0		0.7
LSA1317	lsa1317	Putative chromate reductase	0.6	-0.7	
LSA1450	lsa1450	Putative metal-dependent hydrolase (beta-lactamase family III)			0.6
LSA1776	lsa1776	Putative 4-carboxymuconolactone decarboxylase	0.6		D
Translatio	on, ribosom	al structure and biogenesis			
Translatio	on initiation				
LSA1135	lsa1135	Putative translation factor, Sua5 family		0.7	0.6
Translatio	on elongatio	on			
LSA0251	efp1	Elongation factor P (EF-P)	0.5		
LSA1063	tuf	Elongation factor Tu (EF-Tu)	0.6		
Ribosoma	al proteins				
LSA0011	rpll	50S Ribosomal protein L9			-0.8
LSA0266	rpsN	30S ribosomal protein S14		0.7	-0.5
LSA0494	lsa0494	30S ribosomal interface protein S30EA	1.7		
LSA0696	rpmB	50S ribosomal protein L28			0.8
LSA1017	rpsA	30S Ribosomal protein S1	0.9		0.6
LSA1333	rpmG	50S ribosomal protein L33			0.6
LSA1666	rplL	50S ribosomal protein L7/L12	-0.6		
LSA1676	rpmG2	50S ribosomal protein L33			-0.6
LSA1750	rpIF	50S ribosomal protein L6		0.6	
LSA1755	rpsQ	30S ribosomal protein S17		0.5	
LSA1761	rpIB	50S ribosomal protein L2		0.6	
LSA1765	rpsJ	30S ribosomal protein S10	-0.7		
Protein sy	ynthesis				
LSA0377	tgt	Queuine tRNA-ribosyltransferase	-0.6		
LSA1546	gatB	Glutamyl-tRNA amidotransferase, subunit B		-0.5	
LSA1547	gatA	Glutamyl-tRNA amidotransferase, subunit A	-0.5		-0.5
RNA restr	riction and i	modification			
LSA0437	lsa0437	Hypothetical protein with an RNA-binding domain	-0.7		
LSA0443	Isa0443	Putative single-stranded mRNA endoribonuclease	2.7		1.9
LSA0738	dtd	D-tyrosyl-tRNA(tyr) deacylase	0.5		
LSA0794	trmU	tRNA (5-methylaminomethyl-2-thiouridylate)-methyltransferase		-0.9	
LSA1534	lsa1534	Putative ATP-dependent RNA helicase		0.9	
LSA1615	lsa1615	Putative ATP-dependent RNA helicase	-0.7	-0.8	-1.0
LSA1723	truA	tRNA pseudouridylate synthase A (pseudouridylate synthase I)	-0.7		-0.6
LSA1880	trmE	tRNA modification GTPase trmE	-0.7		
	/I-tRNA synt	hetases			
LSA0880	glyQ	Glycyl-tRNA synthetase, alpha subunit		0.7	
LSA0881	glyS	Glycyl-tRNA synthetase, beta subunit		0.7	
LSA1400	thrS	Threonyl-tRNA synthetase	0.6		
LSA1681	cysS	Cysteinyl-tRNA synthetase	-0.6		

Table 1 Genes with significant differential expression in three *L. sakei* strains grown on ribose compared with glucose, FDR adjusted p-value less than 0.01 and \log_2 of > 0.5 or < -0.5 (\log_2 values > 1.0 or < -1.0 are shown in bold) (Continued)

(Continue	ea) 				
DNA repli	cation, reco	ombination and repair	· · · · · · · · · · · · · · · · · · ·		
DNA repli	ication				
LSA0221	Isa0221	Putative transcriptional regulator, LysR family (C-terminal fragment), degenerate	-0.8	-0.9	-1.1
LSA0976	parE	Topoisomerase IV, subunit B		0.5	
Transpose	on and IS				
LSA1152_a	tnpA3- ISLsa1	Transposase of ISLsa1 (IS30 family)	-0.6		
Phage-rel	ated function	on			
LSA1292	lsa1292	Putative prophage protein	0.6		
LSA1788	lsa1788	Putative phage-related 1,4-beta-N-acetyl muramidase (cell wall hydrolase)	-1.0	D	D
DNA reco	mbination	and repair			
LSA0076	Isa0076	Putative DNA invertase (plasmidic resolvase)	-1.1	-1.5	-1.4
LSA0366	ruvA	Holliday junction DNA helicase RuvA			-0.5
LSA0382	dinP	DNA-damage-inducible protein P	-0.5		
LSA0487	recA	DNA recombinase A	-0.8		-1.1
LSA0523	uvrB	Excinuclease ABC, subunit B	-0.7		-0.5
LSA0524	uvrA1	Excinuclease ABC, subunit A	-1.2		-0.7
LSA0910	rexAN	ATP-dependent exonuclease, subunit A (N-terminal fragment), authentic frameshift	0.6		
LSA0911	rexAC	ATP-dependent exonuclease, subunit A (C-terminal fragment), authentic frameshift	0.7		
LSA0912	Isa0912	Putative ATP-dependent helicase, DinG family	0.6		0.8
LSA1162	lsa1162	DNA-repair protein (SOS response UmuC-like protein)		0.8	-0.6
LSA1405	fpg	Formamidopyrimidine-DNA glycosylase	-0.5	-0.6	-0.6
LSA1477	recX	Putative regulatory protein, RecX family	-0.6		
LSA1843	ogt	Methylated-DNA-protein-cysteine S-methyltransferase	-0.6		
DNA restr	riction and	modification			
LSA0143	Isa0143	Putative adenine-specific DNA methyltransferase	-0.7	D	D
LSA0921	Isa0921	Putative adenine-specific DNA methyltransferase	0.8		
LSA1299	lsa1299	Putative adenine-specific DNA methyltransferase	0.9	0.7	1.2
Information	on pathway	rs			
LSA0326	Isa0326	Putative DNA helicase		-0.6	U
DNA pack	aging and	segregation			
LSA0135	Isa0135	Hypothetical integral membrane protein, similar to CcrB			-0.6
LSA1015	hbsU	Histone-like DNA-binding protein HU	1.0		0.9
		omosome partitioning			
Cell divisi LSA0755	on divIVA	Call-division initiation protain (contum placement)			0.5
LSA0755 LSA0845	lsa0845	Cell-division initiation protein (septum placement) Putative negative regulator of septum ring formation	0.7		0.5
LSA1118	Isa1118	Rod-shape determining protein	0.7	0.6	0.5
LSA1118	ftsH	ATP-dependent zinc metalloendopeptidase FtsH (cell division protein FtsH)		0.0	-0.6
LSA1879	gidA	Cell division protein GidA	-0.6		-0.0
L3A10/9	giuA	Cell division protein dida	-0.0		
Cell envel	ope bioger	nesis, outer membrane			
LSA0280	murE	UDP-N-acetylmuramoylalanyl-D-glutamate-2,6-diaminopimelate ligase	-0.6	-0.6	-0.7
LSA0621	pbp2A	Bifunctional glycolsyltransferase/transpeptidase penicillin binding protein 2A	0.0	0.0	0.7
LSA0648	Isa0648	Putative penicillin-binding protein precursor (beta-lactamase class C)			1.0
LSA0862	Isa0862	N-acetylmuramoyl-L-alanine amidase precursor (cell wall hydrolase) (autolysin)	0.6		0.8
LSA0917	pbp1A	Bifunctional glycosyltransferase/transpeptidase penicillin-binding protein 1A	0.0		0.5
LSA1123	murA1	UDP-N-acetylglucosamine 1-carboxyvinyltransferase I		-0.5	0.5
LSA1123	pbp2B2	Bifuntional dimerisation/transpeptidase penicillin-binding protein 2B		0.7	0.7
LJ/11JJ4	μυμευε	promotion difficultation, transpeptidase penicinin sulfiding protein 20		0.7	0./

Table 1 Genes with significant differential expression in three *L. sakei* strains grown on ribose compared with glucose, FDR adjusted p-value less than 0.01 and \log_2 of > 0.5 or < -0.5 (\log_2 values > 1.0 or < -1.0 are shown in bold) (Continued)

Continue	:u)				
LSA1437	lsa1437	N-acetylmuramoyl-L-alanine amidase precursor (cell wall hydrolase) (autolysin)		-0.7	
LSA1441	bacA	Putative undecaprenol kinase (bacitracine resistance protein A)		0.6	
LSA1613	alr	Alanine racemase	-0.8	-0.9	-0.7
LSA1616	murF	UDP-N-acetylmuramoyl-tripeptide–D-alanyl-D-alanine ligase			-0.5
	-	Ilular processes			
LSA0162	Isa0162	Putative Bifunctional glycosyl transferase, family 8		-1.2	-1.5
LSA1246	lsa1246	Putative glycosyl transferase, family 2		-0.9	
LSA1558	lsa1558	Putative extracellular N-acetylmuramoyl-L-alanine amidase precursor (cell wall hydrolase/Lysosyme subfamily 2)			-0.6
	ity and seci	retion			
Protein se					
LSA0948	IspA 	Signal peptidase II (lipoprotein signal peptidase) (prolipoprotein signal peptidase)			0.5
LSA1884	oxaA2	Membrane protein chaperone oxaA			-0.6
Signal tra					
	nsduction	Two component system concer histiding kingso (Cnal/ fragment) deconcerts		O F	
LSA0561 LSA0692	sppKN Isa0692	Two-component system, sensor histidine kinase, (SppK fragment), degenerate Putative serine/threonine protein kinase		0.5 0.5	0.6
LSA1384	Isa1384	Two-component system, response regulator		0.5	0.0
L3A1304	1501304	Two-component system, response regulator		0.5	
		difications, protein turnover, chaperones			
Protein fo	•				
LSA0050	Isa0050	Putative molecular chaperone, small heat shock protein, Hsp20 family			-0.7
LSA0082	htrA	Serine protease HtrA precursor, trypsin family	0.6	-0.6	
LSA0207	clpL	ATPase/chaperone ClpL, putative specificity factor for ClpP protease	0.6		
LSA0358	groS	Co-chaperonin GroES (10 kD chaperonin) (protein Cpn10)			-0.5
LSA0359	groEL	Chaperonin GroEL (60 kDa chaperonin) (protein Cpn60)			-0.5
LSA0436	lsa0436	Putative peptidylprolyl isomerase (peptidylprolyl cis-trans isomerase) (PPlase)	0.7		-0.6
LSA0984	hslU	ATP-dependent Hsl protease, ATP-binding subunit HslU	0.7	0.6	0.7
LSA1465 LSA1618	clpE	ATPase/chaperone ClpE, putative specificity factor for ClpP protease	-0.7	-0.6 0.8	-0.6
	htpX to atypical	Membrane metalloprotease, HtpX homolog		0.0	
LSA0170	Isa0170	Putative general stress protein	0.5		-1.5
LSA0170 LSA0247		Similar to universal stress protein, UspA family	0.5		-0.5
LSA0247 LSA0264	usp2 Isa0264	Putative glycine/betaine/carnitine/choline transport protein	-0.6		-0.5
LSA0513	Isa0513	Putative stress-responsive transcriptional regulator	-0.0	-0.8	-0.0
LSA0552	Isa0513	Organic hydroperoxide resistance protein		0.6	
LSA0616	Isa0532	Putative glycine/betaine/carnitine/choline ABC transporter, ATP-binding subunit	0.9	0.0	
LSA0617	Isa0617	Putative glycine/betaine/carnitine/choline ABC transporter, membrane-spanning subunit	1.3		
LSA0618	Isa0618	Putative glycine/betaine/carnitine/choline ABC transporter, substrate-binding lipoprotein	0.6		
LSA0619	Isa0619	Putative glycine/betaine/carnitine/choline ABC transporter, membrane-spanning subunit	1.5	0.5	
LSA0642	usp3	Similar to universal stress protein, UspA	0.9	0.5	
LSA0768	csp1	Similar to cold shock protein, CspA family	2.1	0.6	1.8
LSA0706	usp6	Similar to universal stress protein, UspA family	0.6	0.0	1.0
LSA0946	csp4	Similar to cold shock protein, CspA family	0.6		
LSA1110	lsa1110	Putative NifU-homolog involved in Fe-S cluster assembly	0.0	0.6	
LSA1110	Isa1111	Putative cysteine desulfurase (class-V aminotransferase, putative SufS protein homologue)		0.7	
LSA1111	usp4	Similar to universal stress protein, UspA family	1.5	- 2.1	
LSA1173 LSA1694	usp 4 Isa1694	Putative glycine/betaine/carnitine ABC transporter, substrate binding lipoprotein precursor	-1.7	4.1	-1.1
LSA1695	Isa1695	Putative glycine/betaine/carnitine ABC transporter, substrate binding ipoprotein precursor	-2.1	-2.0	-1.9
ころ 1032	13U 1 U 9 J	i diative giyenie/betaine/caminine Abe tiansponter, membrane-spanning subunit	-2.1	-2.0	-1.9

Table 1 Genes with significant differential expression in three *L. sakei* strains grown on ribose compared with glucose, FDR adjusted p-value less than 0.01 and \log_2 of > 0.5 or < -0.5 (\log_2 values > 1.0 or < -1.0 are shown in bold) (Continued)

LSA1696	lsa1696	Putative glycine/betaine/carnitine ABC transporter, ATP-binding subunit	-1.6	-	-0.9
LSA1870	lsa1870	Putative glycine betaine/carnitine/choline ABC transporter, ATP-binding subunit	-0.6		-0.6
Protein m	odification				
LSA0865	Isa0865	Putative protein methionine sulfoxide reductase		-0.6	
LSA0866	msrA	Protein methionine sulfoxide reductase		-0.7	
LSA0934	IpIA	Lipoate-protein ligase	1.6	1.4	1.0
LSA0973	pflA	Pyruvate formate-lyase activating enzyme	1.7		
	•	diction only			
Miscellane				0.7	0.0
LSA0030	Isa0030	Putative aldo/keto reductase (oxidoreductase)	0.5	-0.7	-0.8
LSA0120	lsa0120	Putative GTP-binding protein	-0.5		
LSA0164	Isa0164	Putative serine/tyrosine protein phosphatase	0.2	-1.1	-1.2
LSA0165	Isa0165	Putative oxidoreductase, short chain dehydrogenase/reductase family		-0.9	-1.2
LSA0218	trxA1	Thioredoxin		-0.9	
LSA0258	lsa0258	Putative iron-containing alcohol dehydrogenase	1.6	0.5	1.6
LSA0260	lsa0260	Putative aldo/keto reductase (oxidoreductase)	1.9	1.2	1.7
LSA0312	lsa0312	Putative NADH oxidase	-0.9		-1.0
LSA0324	lsa0324	Putative hydrolase, haloacid dehalogenase family (N-terminal fragment), authentic frameshift	1.9		
LSA0325	lsa0325	Putative hydrolase, haloacid dehalogenase family (C-terminal fragment), authentic frameshift	1.8		
LSA0350	Isa0350	Putative N-acetyltransferase, GNAT family	-0.5		
LSA0369	lsa0369	Putative N-acetyltransferase, GNAT family	-0.5		-0.5
_SA0384	Isa0384	Putative phosphoesterase, DHH family	-0.5		
_SA0403	Isa0403	Putative thioredoxin reductase		0.9	
LSA0447	Isa0447	Putative hydrolase, haloacid dehalogenase family			0.6
LSA0475	Isa0475	Putative N-acetyltransferase, GNAT family		-0.6	
_SA0520	trxB2	Thioredoxin reductase	-0.8		
LSA0575	npr	NADH peroxidase	1.0	U	
LSA0802	nox	NADH oxidase	1.5		
LSA0806	Isa0806	Putative N-acetyltransferase, GNAT family	0.6		
LSA0831	Isa0831	Putative nitroreductase (oxidoreductase)		1.6	
LSA0896	sodA	Iron/Manganese superoxide dismutase	3.4	1.7	1.7
LSA0925	adh	Putative zinc-containg alcohol dehydrogenase (oxidoreductase)	0.5		
LSA0971	рра	Inorganic pyrophosphatase (pyrophosphate phosphohydrolase)	0.7		
LSA0994	lsa0994	Putative GTP-binding protein			0.6
_SA1016	engA	Putative GTP-binding protein	0.6		0.7
LSA1045	obgE	Putative GTP-binding protein	0.6		
LSA1153	lsa1153	Hypothetical protein, CAAX protease family	0.5		
LSA1311	lsa1311	Hypothetical protein containing a possible heme/steroid binding domain	0.7	-0.6	
LSA1320	lsa1320	Putative NADPH-quinone oxidoreductase		-0.8	
_SA1345	lsa1345	Putative hydrolase, haloacid dehalogenase family	0.5		
_SA1349	lsa1349	Putative N-acetyltransferase, GNAT family		-0.5	
LSA1365	lsa1365	Hypothetical protein		-0.5	-0.7
LSA1368	lsa1368	Hypothetical protein	0.9		0.6
LSA1371	lsa1371	Hypothetical membrane protein	0.6		
LSA1395	lsa1395	Putative zinc-containing alcohol dehydrogenase (oxidoreductase)	0.9		
LSA1427	lsa1427	Putative hydrolase, haloacid dehalogenase	1.3		0.6
_SA1472	lsa1472	Putative N-acetyl transferase, GNAT family	0.6		
_SA1535	lsa1535	Putative oxidoreductase	0.5	1.1	0.7
LSA1553	lsa1553	Putative hydrolase, haloacid dehalogenase family	-0.6		

Table 1 Genes with significant differential expression in three *L. sakei* strains grown on ribose compared with glucose, FDR adjusted p-value less than 0.01 and \log_2 of > 0.5 or < -0.5 (\log_2 values > 1.0 or < -1.0 are shown in bold) (Continued)

(Continue					
LSA1559	lsa1559	Putative oxidoreductase	0.6	1.1	0.7
LSA1702	lsa1702	Putative zinc-containing alcohol dehydrogenase (oxidoreductase)	1.1		
LSA1712	lsa1712	Putative nitroreductase (oxidoreductase)		-0.7	-0.8
LSA1832	lsa1832	Putative zinc-containing alcohol dehydrogenase (oxidoreductase)		1.0	
LSA1835	lsa1835	Putative zinc-containing alcohol dehydrogenase (oxidoreductase)	-0.7		-1.0
LSA1867	lsa1867	Putative acetyltransferase, isoleucine patch superfamily	-0.5	-0.6	-0.7
LSA1871	gshR	Glutathione reductase	-0.6		
Unknown					
		function that are similar to other proteins			
LSA0018	lsa0018	Hypothetical protein		0.5	
LSA0027	Isa0027	Hypothetical protein			-1.1
LSA0028	Isa0028	Hypothetical protein, DegV family	-0.5		
LSA0044	Isa0044	Hypothetical protein			-0.7
LSA0061	Isa0061	Hypothetical extracellular protein precursor	-0.5		
LSA0106	Isa0106	Hypothetical cell surface protein precursor	0.5		
LSA0160	Isa0160	Hypothetical protein	-0.7		
LSA0166	Isa0166	Hypothetical Integral membrane protein			-1.2
LSA0190	Isa0190	Hypothetical integral membrane protein	-0.7		-0.6
LSA0191	lsa0191	Hypothetical integral membrane protein	-0.6		-0.6
LSA0199	lsa0199	Hypothetical protein	1.1	1.0	1.1
LSA0208	Isa0208	Hypothetical integral membrane protein	0.7		
LSA0235	Isa0235	Hypothetical extracellular protein precursor	2.1	1.6	1.7
LSA0236	Isa0236	Hypothetical extracellular peptide precursor	2.0	1.3	1.5
LSA0244	Isa0244	Hypothetical integral membrane protein			-0.5
LSA0245	Isa0245	Hypothetical lipoprotein precursor	-0.9	-1.0	-1.1
LSA0249	lsa0249	Hypothetical protein	1.1	1.0	
LSA0263	Isa0263	Hypothetical integral membrane protein	-0.6		-0.9
LSA0300	Isa0300	Hypothetical protein			0.7
LSA0315	Isa0315	Hypothetical protein	-0.7		
LSA0319	lsa0319	Hypothetical protein		-0.8	-0.8
LSA0323	Isa0323	Hypothetical protein			-0.5
LSA0337	Isa0337	Hypothetical protein	-0.7		
LSA0348	lsa0348	Hypothetical integral membrane protein	-0.9		-0.7
LSA0352	lsa0352	Hypothetical integral membrane protein	-0.6		
LSA0354	lsa0354	Hypothetical integral membrane protein			-1.1
LSA0388	lsa0388	Hypothetical protein		-0.6	
LSA0389	lsa0389	Hypothetical protein		-0.7	-0.7
LSA0390	lsa0390	Hypothetical protein		-0.5	
LSA0409	lsa0409	Hypothetical integral membrane protein			-0.8
LSA0418	lsa0418	Hypothetical protein			-0.8
LSA0464	lsa0464	Hypothetical protein		-0.6	
LSA0470	Isa0470	Hypothetical protein	0.9		0.7
LSA0512	lsa0512	Hypothetical protein		-0.6	
LSA0515	Isa0515	Hypothetical integral membrane protein		-0.5	
LSA0536	Isa0536	Hypothetical protein		0.7	
LSA0716	Isa0716	Hypothetical protein		- **	0.6
LSA0752	Isa0752	Hypothetical protein	0.5		0.6
LSA0757	Isa0757	Hypothetical protein	0.5	0.8	5.0
LSA0773	Isa0773	Hypothetical protein	0.9	0.0	0.6
	.50.0775		5.7		5.0

Table 1 Genes with significant differential expression in three *L. sakei* strains grown on ribose compared with glucose, FDR adjusted p-value less than 0.01 and \log_2 of > 0.5 or < -0.5 (\log_2 values > 1.0 or < -1.0 are shown in bold) (Continued)

LSA0784	Isa0784	Hypothetical protein	-2.6		
LSA0786	Isa0786	Hypothetical protein	-2.0		
LSA0787	Isa0787	Hypothetical protein	-1.7		
LSA0790	Isa0790	Hypothetical protein, ATP utilizing enzyme PP-loop family	-2.5		
LSA0827	Isa0827	Hypothetical lipoprotein precursor	0.8		U
LSA0828	Isa0828	Hypothetical protein	0.7		
LSA0829	Isa0829	Hypothetical integral membrane protein			0.5
LSA0874	Isa0874	Hypothetical protein	0.5		
LSA0901	Isa0901	Hypothetical protein			0.5
LSA0913	Isa0913	Hypothetical extracellular protein precursor	0.5		0.7
LSA0919	lsa0919	Hypothetical protein			0.7
LSA0933	Isa0933	Hypothetical protein	0.6		0.6
LSA0961	Isa0961	Hypothetical protein, DegV family		-0.5	
LSA0968	Isa0968	Hypothetical integral membrane protein	0.7		
LSA0977	Isa0977	Hypothetical integral membrane protein	0.7		0.8
LSA0987	Isa0987	Hypotehtical protein, GidA family (C-terminal fragment)	0.5		
LSA0996	lsa0996	Hypothetical protein			0.5
LSA1003	lsa1003	Hypothetical protein	2.0		1.2
LSA1005	lsa1005	Hypothetical membrane protein	0.9	0.6	0.7
LSA1008	lsa1008	Putative extracellular chitin-binding protein precursor		0.9	1.2
LSA1027	lsa1027	Hypothetical protein			0.6
LSA1047	lsa1047	Hypothetical protein	3.5	1.2	1.3
LSA1064	lsa1064	Hypothetical protein	0.5		0.7
LSA1075	lsa1075	Hypothetical protein			0.5
LSA1078	lsa1078	Hypothetical protein			0.6
LSA1081	lsa1081	Hypothetical protein	1.0		1.0
LSA1091	lsa1091	Hypothetical protein			0.6
LSA1096	lsa1096	Hypothetical protein	0.6		
LSA1124	lsa1124	Hypothetical protein		-0.7	
LSA1154	lsa1154	Hypothetical protein	0.6		0.6
LSA1158	lsa1158	Hypothetical protein	1.7	1.4	
LSA1189	lsa1189	Hypothetical integral membrane protein	-1.6		-1.1
LSA1282	lsa1282	Hypothetical protein		-0.5	
LSA1296	lsa1296	Hypothetical integral membrane protein		-1.2	-0.8
LSA1342	lsa1342	Hypothetical protein		-0.7	
LSA1346	lsa1346	Hypothetical protein	0.8		
LSA1350	lsa1350	Hypothetical protein		-0.6	-1.0
LSA1353	lsa1353	Hypothetical integral membrane protein	-0.9	-0.5	
LSA1446	lsa1446	Hypothetical protein	-0.6	-0.6	-0.7
LSA1466	lsa1466	Hypothetical protein	0.6		
LSA1467	lsa1467	Hypothetical protein		-0.6	-1.1
LSA1524	Isa1524	Hypothetical protein	0.7		
LSA1540	Isa1540	Hypothetical extracellular protein precursor	0.7		
LSA1563	Isa1563	Hypothetical integral membrane protein	0.7	-0.6	-0.6
LSA1610	Isa1610	Hypothetical integral membrane protein	-0.7	0.0	-0.9
LSA1617	Isa1617	Hypothetical protein	0.7		-0.7
LSA1620	Isa1620	Hypothetical protein			-0.6
LSA1623	Isa1623	Hypothetical integral membrane protein	-0.5		-0.6
20111020		,,			
LSA1637	lsa1637	Hypothetical integral membrane protein, TerC family	-1.7	-1.0	-1.6

Table 1 Genes with significant differential expression in three *L. sakei* strains grown on ribose compared with glucose, FDR adjusted p-value less than 0.01 and \log_2 of > 0.5 or < -0.5 (\log_2 values > 1.0 or < -1.0 are shown in bold) (Continued)

(Commuc	,				
LSA1649	lsa1649	Hypothetical extracellular protein precursor			-0.5
LSA1659	lsa1659	Hypothetical protein	-0.5		
LSA1662	lsa1662	Hypothetical protein	-1.0	-0.6	-0.7
LSA1663	lsa1663	Hypothetical protein	-0.8		
LSA1678	lsa1678	Hypothetical protein	-0.6		
LSA1680	lsa1680	Hypothetical protein	-0.6		
LSA1716	lsa1716	Hypothetical protein		-0.5	
LSA1822	lsa1822	Hypothetical protein			-0.5
LSA1828	lsa1828	Hypothetical integral membrane protein	0.6	0.7	
LSA1850	lsa1850	Hypothetical protein		-0.6	
LSA1876	lsa1876	Hypothetical integral membrane protein			-0.6
LSA1877	lsa1877	Hypothetical protein			-0.6
Proteins o	f unknown	function only similar to other proteins from the same organism			
LSA1159	lsa1159	Hypothetical cell surface protein precursor	2.0		0.5
LSA1165	lsa1165	Hypothetical cell surface protein precursor	1.8		
LSA1700	lsa1700	Hypothetical protein	2.1	0.8	
LSA1814	lsa1814	Hypothetical protein			-0.5
Proteins o	f unknown	function. without similarity to other proteins			
LSA0065	Isa0065	Hypothetical integral membrane protein	-0.5		
LSA0093	Isa0093	Hypothetical integral membrane protein	-0.9		-1.2
LSA0121	Isa0121	Hypothetical small peptide	-0.7	-0.6	-0.5
LSA0163	Isa0163	Hypothetical protein		-1.1	-1.3
LSA0167	Isa0167	Hypothetical protein			-1.4
LSA0168	Isa0168	Hypothetical protein			-1.4
LSA0188	Isa0188	Hypothetical small peptide			-0.8
LSA0256_a	lsa0256_a	Hypothetical protein	2.3	1.0	2.2
LSA0257	Isa0257	Hypothetical protein	1.4		
LSA0281	Isa0281	Hypothetical lipoprotein precursor		-0.5	-0.6
LSA0301	Isa0301	Hypothetical protein			0.6
LSA0334	lsa0334	Hypothetical extracellular protein precursor	1.1		
LSA0339	lsa0339	Hypothetical protein	-0.5		
LSA0378	Isa0378	Hypothetical protein	-0.7		
LSA0514	lsa0514	Hypothetical small extracellular protein precursor		-0.8	
LSA0534	lsa0534	Hypothetical cell surface protein precursor (with LPQTG sorting signal)	1.0		D
LSA0576	Isa0576	Hypothetical protein	0.5	D	
LSA0641	Isa0641	Hypothetical extracellular peptide precursor		-0.5	
LSA0647	Isa0647	Hypothetical extracellular protein precursor	0.6		
LSA0667	Isa0667	Hypothetical protein	1.0		0.9
LSA0753	Isa0753	Hypothetical integral membrane protein			0.5
LSA0789	Isa0789	Hypothetical protein	-1.9		
LSA0837	Isa0837	Hypothetical protein	1.2	1.3	1.4
LSA0885	Isa0885	Hypothetical protein	1.8		
LSA0902	lsa0902	Hypothetical protein	0.7	D	
LSA0945	Isa0945	Hypothetical protein			0.9
LSA1019	lsa1019	Hypothetical cell surface protein precursor			0.8
LSA1035	lsa1035	Hypothetical small integral membrane protein			0.6
LSA1086	Isa1086	hypothetical protein	0.8		0.5
LSA1104	Isa1104	Hypothetical protein	-0.5		
LSA1155	lsa1155	Hypothetical integral membrane protein	0.5		
LSA1174	lsa1174	Hypothetical protein	1.0		

Table 1 Genes with significant differential expression in three *L. sakei* strains grown on ribose compared with glucose, FDR adjusted p-value less than 0.01 and \log_2 of > 0.5 or < -0.5 (\log_2 values > 1.0 or < -1.0 are shown in bold) (Continued)

LSA1176	lsa1176	Hypothetical protein		-1.0	U
LSA1319	lsa1319	Hypothetical small protein		-0.8	
LSA1408	lsa1408	Hypothetical protein			0.6
LSA1464	lsa1464	Hypothetical protein	-0.6		
LSA1478	lsa1478	Hypothetical protein	-0.7	-0.6	-0.6
LSA1480	lsa1480	Hypothetical membrane protein	0.5	D	
LSA1524	lsa1524	Hypothetical protein	0.8		
LSA1539	lsa1539	Hypothetical protein	0.9		
LSA1713	lsa1713	Hypothtical small peptide			-0.6
LSA1787	lsa1787	Hypothetical cell surface protein precursor	-0.5	U	
LSA1820	lsa1820	Hypothetical cell surface protein precursor			-0.6
LSA1821	lsa1821	Hypothetical cell surface protein precursor		-0.6	
LSA1845	lsa1845	Hypothetical small protein		0.8	
LSA1848	lsa1848	Hypothetical protein			-0.5
LSA1851	lsa1851	Hypothetical extracellular small protein	-0.6		-0.7
LSA1883	lsa1883	Hypothetical small protein	1.2		1.5
Bacterioci	n associate	d genes			
SKP0001	sppIP	Bacteriocin sakacin P inducing peptide	D	0.5	D
SKP0006	sppT	Sakacin P ABC transporter	D	0.6	D
SKP0007	sppE	Sakacin P accesory transport protein	D	0.6	D

The microarray used has been described previously [32]. Asterix (*) relates the gene to Table 2. D and U refer to genes classified as 'divergent' and 'uncertain', respectively, by CGH analysis [32]. Genes encoding proteins with a change in expression according to McLeod et al. [19], are underlined.

protein expression of glyceraldehyde-3-phosphate dehydrogenase, phosphoglycerate kinase and enolase previously seen in LS 25 [19]. The latter three enzymes are encoded from the central glycolytic operon (*cggR-gap-pgk-tpi-eno*) together with triose-phosphate isomerase and the putative central glycolytic genes regulator (CggR) [46]. Besides the *cggR* gene being down-regulated in MF1053 and LS 25, no change in gene expression was seen of these central glycolytic genes. Thus at the transcription level it is not obvious that the LS 25 strain

Up-regulated Down-regulated

23K

222

62

96

121

71

167

MF1053

LS 25

MF1053

LS 25

LS 25

Figure 1 Venn diagram showing the number of unique and common up- and down-regulated genes in *L. sakei* strains 23K, MF1053 and LS 25 when grown on ribose compared with glucose.

down-regulate the glycolytic pathway more efficiently than the other strains, as previously suggested [19].

Interestingly, all the strains showed an induction (1.4-2.3) of mgsA encoding methylglyoxal synthase, which catalyzes the conversion of dihydroxyacetone-phosphate to methylglyoxal (Figure 2). The presence of this gene is uncommon among LAB and so far a unique feature among the sequenced lactobacilli. The methylglyoxal pathway represents an energetically unfavourable bypass to the glycolysis. In *E. coli*, this bypass occurs as a response to phosphate starvation or uncontrolled carbohydrate metabolism, and enhanced ribose uptake was shown to lead to the accumulation of methylglyoxal [47,48]. As suggested by Chaillou et al. [7], such flexibility in the glycolytic process in L. sakei may reflect the requirement to deal with glucose starvation or to modulate carbon flux during cometabolism of alternative carbon sources. Breakdown of methylglyoxal is important as it is toxic to the cells [49]. An induction of the *lsa1158* gene contiguous with *mgsA* was seen for 23K and MF1053. This gene encodes a hypothetical protein, also suggested as a putative oxidoreductase, which may reduce methylglyoxal to lactaldehyde [7]. However, no induction of the adhE (lsa0379) gene encoding an iron-containing aldehyde dehydrogenase suggested to further reduce lactaldehyde to L-lactate [7] was seen. By CGH [32] lsa1158 and adhE were present in all

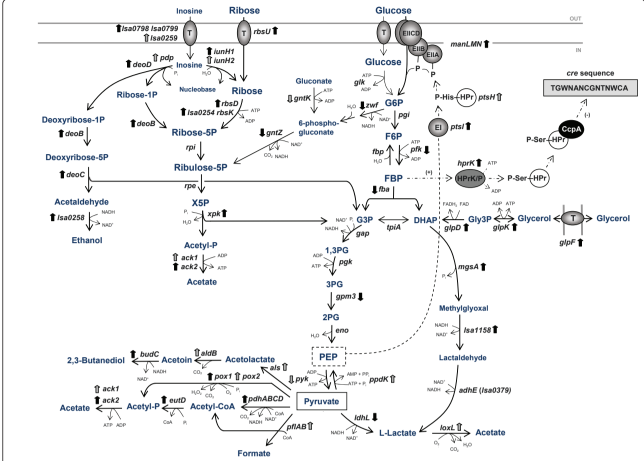


Figure 2 Overview of the glycolysis, phosphoketolase pathway and nucleoside catabolic pathway affected by the change of carbon source from glucose to ribose in three *L. sakei* strains in this study. Genes which expression is up- or down-regulated are indicated with upward and downward pointing arrows, respectively, and are listed in Table 1. Black arrows indicate regulation in all three strains, and grey arrows indicate regulation in one or two strains. Schematic representation of CcpA-mediated CCR pathway is shown in the upper right corner. Ell, enzyme II of the phosphotransferase system (PTS); El, enzyme I, HPr, Histidine-containing protein; T, transport protein; P, phosphate; HPrK/P, HPr kinase/phosphatase; G6P, glucose-6-phosphate; F6P; fructose-6-phosphate; FBP, fructose-1,6-bisphosphate; G3P, glyceraldehyde-3-phosphate; T,3PG, 1,3-phosphoglycerate; 3PG, 3-phosphoglycerate; G9P, glyceraldehyde-3-phosphate; G9P, glucokinase; *pgi*, phosphoglycerate; G9P, fructose-1,6-bisphosphatase; *tpi*, triose-phosphate isomerase; *gap*, glyceraldehyde-3-phosphate dehydrogenase; *pgk*, phosphoglycerate kinase; *eno*, enolase; *rpi*, ribose-5-phosphate isomerase; *rpe*, ribulose-phosphate 3-epimerase.

the *L. sakei* strains investigated, whereas *mgsA* was lacking in some strains, indicating that the MgsA function is not vital.

Pyruvate metabolism

Pyruvate is important in both glycolysis and PKP. It can be converted into lactate by the NAD-dependent L-lactate dehydrogenase, which regenerates NAD⁺ and maintains the redox balance. This enzyme is encoded by the *ldhL* gene which was down-regulated (0.7-1.4) in all three strains, in accordance with previous findings [50], and the down-regulation was strongest for the LS 25 strain. At the protein level, only LS 25 showed a lower expression of this enzyme during growth on ribose [19].

Genes responsible for alternative fates of pyruvate (Figure 2) were highly induced in all the strains, however with some interesting strain variation (Table 1). The shift in pyruvate metabolism can benefit the bacteria by generating ATP, or by gaining NAD⁺ for maintaining the redox balance and may lead to various end products in addition to lactate [51].

In all the strains, a strongly up-regulated (2.1-3.0) pox1 gene was observed, and in 23K an up-regulated pox2 (0.7), encoding pyruvate oxidases which under aerobic conditions convert pyruvate to acetyl-phosphate with hydrogen peroxide (H_2O_2) and CO_2 as side products. Accumulation of peroxide ultimately leads to aerobic growth arrest [52]. H_2O_2 belongs to a group of compounds known as reactive

oxygen species and reacts readily with metal ions to yield hydroxyl radicals that damage DNA, proteins and membranes [53]. Remarkable differences in redox activities exist among *Lactobacillus* species and *L. sakei* is among those extensively well equipped to cope with changing oxygen conditions, as well as dealing effectively with toxic oxygen byproducts [7]. 23K up-regulated npr (1.0) encoding NADH peroxidase which decomposes low concentrations of H_2O_2 to H_2O and O_2 , and all the strains upregulated the sodA gene (1.7-3.4) encoding a superoxide dismutase which produces hydrogen peroxide from superoxide (O_2^-) . Various oxidoreductases showed an up-regulation in all the strains (Table 1), indicating the need for the bacterium to maintain its redox balance.

The pdhABCD gene cluster encoding components of the pyruvate dehydrogenase enzyme complex (PDC) which transforms pyruvate into acetyl-CoA and CO₂ were among the strongly up-regulated (2.1-3.7) genes. The *eutD* gene encoding a phosphate acetyltransferase which further forms acetyl-phosphate from acetyl-CoA was also induced (1.0-2.0). Pyruvate can be transformed to acetolactate by acetolactate synthase and further to acetoin by acetolactate decarboxylase, before 2,3-butanediol may be formed by an acetoin recuctase (Figure 2). While the *budC* gene encoding the acetoin reductase showed a strong up-regulation in all three strains, the als-aldB operon was only strongly up-regulated in LS 25 (1.9). Pyruvate formate lyase produces acetyl-CoA and formate from pyruvate. Only in 23K, the *pflAB* genes encoding formate C-acetyltransferase and its activating enzyme involved in formate formation were strongly upregulated (4.0 and 1.7, respectively). This strain was the only one to strongly induce L-lactate oxidase encoding genes which are responsible for conversion of lactate to acetate when oxygen is present (Table 1). In 23K and LS 25, the *ppdK* gene coding for the pyruvate phosphate dikinase involved in regenerating PEP, was induced, as was also lsa0444 encoding a putative malate dehydrogenase that catalyzes the conversion of malate into oxaloacetate using NAD⁺ and vice versa (Table 1).

During growth on ribose, *L. sakei* was shown to require thiamine (vitamine B1) [15]. The E1 component subunit α of the PDC, as well as Pox and Xpk, require thiamine pyrophosphate, the active form of thiamine, as a coenzyme [54]. This could explain the induction of the *thiMDE* operon and *lsa0055* in LS 25, as well as *lsa0980* in 23K, encoding enzymes involved in thiamine uptake and biosynthesis (Table 1). The up-regulation of *lsa1664* (1.1-1.6) encoding a putative dihydrofolate reductase involved in biosynthesis of riboflavin (vitamin B2) in all the strains could indicate a requirement for flavin nucleotides as enzyme cofactors. Riboflavin is the precursor for flavin mononucleotide (FMN) and flavin adenine dinucleotide (FAD) redox cofactors in flavoproteins, and the

E3 component of PDC as well as glycerol-3-phosphate dehydrogenase encoded from the up-regulated *glpD*, are among enzymes requiring FAD. Another cofactor which seems to be important during growth on ribose is lipoate, essential of the E2 component of the PDC. An up-regulation of *lplA* (1.0 - 1.6) encoding lipoate-protein ligase, which facilitates attachment of the lipoyl moiety to metabolic enzyme complexes, was seen in all the strains, allowing the bacterium to scavenge extracellular lipoate [55,56].

Nucleoside catabolism

The L. sakei genome contains a multiplicity of catabolic genes involved in exogenous nucleoside salvage pathways, and the bacterium has been shown to catabolize inosine and adenosine for energy [7]. Three *iunH* genes are present in the 23K genome, which encode inosineuridine preferring nucleoside hydrolases responsible for conversion of inosine to ribose and purine base. The iunH1 gene was up-regulated in all the strains when grown on ribose (1.8-2.6), as was also the iunH2 gene in 23K (1.2). The deoC gene encodes a deoxyribose-phosphate aldolase, and is located in an operon structure preceding the genes deoB, deoD, lsa0798, lsa0799, deoR and pdp which encode phosphopentomutase, purine nucleoside phosphorylase, pyrimidine-specific nucleoside symporter, a putative purine transport protein, the deoxyribonucleoside synthesis operon transcriptional regulator (DeoR), and a pyrimidine-nucleoside phosphorylase, respectively. The complete operon was induced in all the strains, except for pdp only induced in 23K (Table 1). The phosphorylases catalyze cleavage of ribonucleosides and deoxyribonucleosides to the free base pluss ribose-1-phosphate or deoxyribose-1-phosphate. The bases are further utilized in nucleotide synthesis or as nitrogen sources. The pentomutase converts ribose-1phosphate or deoxyribose-1-phosphate to ribose-5-phosphate or deoxyribose-5-phosphate, respectively, which can be cleaved by the aldolase to glyceraldehyde-3-phosphate and acetaldehyde. Glyceraldehyde-3-phosphate enters the glycolysis, while a putative iron containing alcohol dehydrogenase, encoded by lsa0258 up-regulated in all the strains (0.5-1.6), could further reduce acetaldehyde to ethanol (Figure 2). The obvious induced nucleoside catabolism at the level of gene expression was not seen by proteomic analysis [19].

Genes involved in glycerol/glycerolipid/fatty acid metabolism

During growth on ribose, a strong induction of the *glpKDF* operon encoding glycerol kinase (GlpK), glycerol-3-phosphate dehydrogenase (GlpD), and glycerol uptake facilitator protein was observed (Table 1), which is in correlation with the over-expression of GlpD and GlpK seen by

proteomic analysis [19]. GlpD is FADH₂ linked and converts glycerol-3-phosphate to dihydroxyacetone-phosphate. An over-expression of GlpD was also reported when L. sakei was exposed to low temperature [57]. A glpD mutant showed enhanced survival at low temperature, and it was suggested that this was a result of the glycerol metabolism being redirected into phosphatidic acid synthesis which leads to membrane phospholipid biosynthesis [57]. Nevertheless, a down-regulation was observed of the lsa1493 gene (0.6-0.9) encoding a putative diacylglycerol kinase involved in the synthesis of phosphatidic acid, and of cfa (1.3-1.4) encoding cyclopropane-fatty-acyl-phospholipid synthase directly linked to modifications in the bacterial membrane fatty acid composition that reduce membrane fluidity and helps cells adapt to their environment [58]. Interestingly, LS 25 upregulated several genes (LSA0812-0823), including accD and accA encoding the α - and β -subunits of the multi-subunit acetyl-CoA carboxylase (Table 1). This is a biotin-dependent enzyme that catalyzes the irreversible carboxylation of acetyl-CoA to produce malonyl-CoA, an essential intermediate in fatty acid biosynthesis. In B. sub*tilis*, the malonyl-CoA relieves repression of the *fab* genes [59]. We observed that also acpP, fabZ1, fabH, fabD and fabI (Table 1) encoding enzymes involved in fatty acid biosynthesis were induced in LS 25. The altered flux to malonyl-CoA may be a result of the decreased glycolytic rate. MF1053, on the other hand, showed a down-regulation of several genes in the same gene cluster. A higher level of acetate is produced when the bacterium utilizes ribose, and acetate lowers the pH and has a higher antimicrobial effect than lactate. Changes in the phospholipid composition could be a response to changes in intracellular pH. Protons need to be expelled at a higher rate when the pH drops. The LS 25 strain which showed faster growth rates than the other strains [9], was the only strain to up-regulate the F_0F_1 ATP synthase (Table 1), which at the expense of ATP expels protons during low pH.

Regulation mechanisms

Little is known about the regulation of catabolic pathways in *L. sakei*. Starting from ribose uptake, the *rbs* operon may be both relieved from repression and ribose induced. Presumably, a dual regulation of this operon by two opposite mechanisms, substrate induction by ribose and CCR by glucose may occur in *L. sakei*. The *ccpA* gene was not regulated, consistent with this gene commonly showing constitutive expression in lactobacilli [42,60]. The local repressor RbsR is homologous with CcpA, both belonging to the same LacI/GalR family of transcriptional regulators. RbsR was proposed to bind a *cre*-like consensus sequence located close to a putative CcpA *cre* site, both preceding *rbsU* [28]. RbsR in the Gram-positive soil bacterium *Corynebacterium glutamicum* was shown to bind a *cre*-like

sequence, and using microarrays, the transcription of no other genes but the rbs operon was affected positively in an rbsR deletion mutant. It was concluded that RbsR influences the expression of only the rbs operon [61]. Similarily, in the $L.\ sakei$ sequence, no other candidate members of RbsR regulation could be found [28]. However, experiments are needed to confirm RbsR binding in

L. sakei. In Bacillus subtilis, RbsR represent a novel interaction partner of P-Ser-HPr in a similar fashion to CcpA [62]. The P-Ser-HPr interaction is possible also in L. sakei as the bacterium exhibits HPr-kinase/phosphatase activity.

A putative cre site is present in the promoter of lsa0254 encoding the second ribokinase (Table 2), and this gene is preceded by the opposite oriented gene *lsa0253* encoding a transcriptional regulator with a sugar binding domain which belongs to the GntR family. This family of transcriptional regulators, as well as the LacI family which RbsR and CcpA belong to, are among the families to which regulators involved in carbohydrate uptake or metabolism usually belong [63]. The GntR-type regulator could possibly be involved in regulating the expression of the second ribokinase, or of the inosine-uridine preferring nucleoside hydrolase encoding iunH1 gene which is located further upstream of lsa0254. C. glutamicum possesses an operon encoding a ribokinase, a uridine transporter, and a uridine-preferring nucleoside hydrolase which is co-controlled by a local repressor together with the RbsR repressor of the rbs operon [60,61,64]. It is possible that such co-control could exist also in L. sakei. Ribose as well as nucleosides are products of the degradation of organic materials such as DNA, RNA and ATP. The simultaneous expression of the *rbs* and *deo* operons as well as the other genes involved in ribose and nucleoside catabolism (Figure 2) allows the bacterium to access the different substrates simultaneously and use both ribose as well as nucleosides as carbon and energy source. DeoR shows 51% identity to the B. subtilis DeoR repressor protein [65,66]. Genes encoding deoxyribosephosphate aldolase, nucleoside uptake protein and pyrimidine nucleoside phosphorylase in B. subtilis are organized in a dra-nupC-pdp operon followed by deoR, and ribose was shown to release DeoR from DNA binding and thus repression of the operon genes are alleviated [65-67]. The *B. subtilis* pentomutase and purine-nucleoside phosphorylase are encoded from a drm-pupG operon which is not negatively regulated by DeoR, though both operons are subject to CcpA mediated CCR [65,66,68]. As a cre site is found preceding the L. sakei deoC (Table 2), the operon could be regulated by CcpA as well. It is interesting that deoR is the only strongly induced transcriptional regulator gene in all three strains, and the encoded regulator has sigma (σ) factor activity. We can only speculate whether it could function as

Table 2 Putative cre sites present in the promoter region of some L. sakei genes up-regulated in the present study

Gene locus	Gene	cre site sequence ^a	Position ^b	Co-transcribed genes/operon ^c	Gene locus
LSA0123	Isa0123	TGAAAG CG TTA <u>CA</u> A	-93		
LSA0185	galP	<u>GA</u> ACAT CG TTATCA	-46		
LSA0200	rbsU	<u>GT</u> AAAC CG TTTTCA	-113	rbsUDK	LSA0200-0202
LSA0254	Isa0254	TGTAAG CG TTTT <u>AT</u>	-56	lsa0254-lsa0255-lsa0256_a	LSA0254-0256_a
LSA0289	xpk	<u>CT</u> ATTA CG ATGACA	-8		
LSA0292	budC	tgtaac cg tttt <u>a</u> a	-51		
_SA0353	lsa0353	<u>a</u> gaaag cg ctta <u>t</u> a	-102		
_SA0370	arcA	TGAAAG CG ATTAC <u>C</u>	-58	arcA-arcB ^e -arcC-arcT ^e -arcD ^e	LSA0370-0374
SA0449	manL	tgttag cg tttt <u>t</u> a	-56	manL-manM-manN	LSA0449-0451
_SA0533	iunH2	<u>aa</u> aaag cg ttcaca	-35		
SA0572	tdcB	tgaaaa cg ttct <u>a</u> a	-134		
_SA0608	Glo AN	tgtaac cg tttt <u>a</u> a	-100	gloAN-gloAC	LSA0608-0609
_SA0649	glpK	<u>A</u> GGAAA CG TTTTC <u>C</u>	-42	glpK-glpD-glpF	LSA0649-0651
_SA0664	loxL1	<u>a</u> gaaag cg agtaca	-82	loxL1N-loxL1-loxL1C	LSA0664-0666
_SA0764	galK	TGAAAG CG ATTA <u>AT</u>	-30	galK-galE1-galT-galM	LSA0764-0767
_SA0795	deoC	TGAAAG CG TTAACA	-33	deoC-deoB-deoD-lsa0798-lsa0799-deoR-pdp	LSA0795-0801
_SA0974	pflB	TACGAA CG CTTACA	-147	pflB-pflA	LSA0974-0973
_SA1048	fruR ^e	TGTAAA CG ATGACA	-39	fruR ^e -fruK ^e -fruA	LSA1048-1050
SA1141	ррдК	<u>G</u> GTTAT CG ATAA <u>A</u> A	-29		
_SA1146	manA	<u>c</u> gaaat cg cttt <u>a</u> a	-98		
_SA1188	pox1	TGTAAT CG ATTTCA	-88		
SA1204	lsa1204	TGTAAT CG TTTT <u>TT</u>	-127		
_SA1343	eutD	<u>GT</u> AAAA CG CTCTCA	-94		
SA1399	loxL2	TGTAAA CG ATTTCA	-42		
_SA1457	lsa1457	TGATAA CG CTTACA	-85		
_SA1463 ^d	ptsH	TGAAAG CG GTAT <u>AG</u>	-161	ptsHl	LSA1463-1462
SA1641	nanE	TGTAAG CG GTTA <u>AT</u>	-85	nanE-nanA	LSA1641-1640
_SA1643	lsa1643	TGATAA CG CTTACA	-31		
_SA1651	lsa1651	<u>G</u> GTAAG CG GTTA <u>A</u> A	-148		
_SA1711	lacL	TGAAAC CG TTTT <u>A</u> A	-36	lacL-lacM	LSA1711-1710
_SA1792	scrA	TGTAAA CG GTT <u>GT</u> A	-78	scrA-dexB-scrK	LSA1792-1790
LSA1830	pox2	TTGTAA CG CTTACA	-70		

The identification is based on the genome sequence of L. sakei strain 23K, and the consensus sequence TGWNANCG NTNWCA (W = A/T, N = A/T/G/C), confirmed in Gram-positive bacteria [39] was used in the search, allowing up to two mismatches (underlined) in the conserved positions except for the two center positions, highlighted in boldface.

activator of transcription on some of the regulated genes in this study.

Expression of the Xpk encoding gene of *Lactobacillus pentosus* was reported to be induced by sugars fermented through the PKP and repressed by glucose mediated by CcpA [69]. Indeed, the *cre* site overlapping ATG start codon of *L. sakei xpk* (Table 2) indicates relief of CcpA-mediated CCR during growth on ribose. Also for several genes involved in alternative fates of pyruvate, putative *cre* sites were present (Table 2).

Several genes and operons involved in transport and metabolism of various carbohydrates such as mannose, galactose, fructose, lactose, cellobiose, N-acetylglucosamine, including putative sugar kinases and PTSs, were induced during growth on ribose (Table 1), and as shown in Table 2, putative *cre* sites are located in the promoter region of many of these up-regulated genes and operons. 23K showed an up-regulation of genes involved in the arginine deiminase pathway, and 23K and LS 25 showed an up-regulated threonine deaminase (Table 1). The *arcA* and *tdcB* both have putative *cre* sites in their promoter regions (Table 2). Thus ribose seems to induce a global regulation of carbon metabolism in *L. sakei*.

^a mismatch to consensus sequence is underlined

^b position of *cre* in relation to the start codon

^c suggested co-transcribed genes or genes organized in an operon

^d cre in preceding gene encoding hypothetical protein

e gene not regulated in this study

A putative *cre* site precedes the *glp* operon (Table 2), suggesting regulation mediated by CcpA. However, regulation of the L. sakei GlpK may also occur by an inducer exclusion-based CcpA-independent CCR mechanism as described in enterococci and B. subtilis [70,71], and as previously suggested by Stentz et al. [15]. By this mechanism, glycerol metabolism is regulated by PEPdependent, EI- and HPr-catalyzed phosphorylation of GlpK in response to the presence or absence of a PTS substrate. In the absence of a PTS sugar, GlpK is phosphorylated by P-His-HPr at a conserved histidyl residue, forming the active P-GlpK form, whereas during growth on a PTS sugar, phosphoryl transfer flux through the PTS is high, concentration of P-His-HPr is low, and GlpK is present in a less active dephospho form [20,70,71]. This conserved histidyl residue (His232) is present in L. sakei GlpK [20], and Stentz et al. [15] reported that whereas L. sakei can grow poorly on glycerol, this growth was abolished in ptsI mutants.

Mannose-PTS

As mentioned in the introduction, the PTS plays a central role, in both the uptake of a number of carbohydrates and regulatory mechanisms [20-22]. Encoding the general components, ptsH showed an up-regulation in MF1053 and LS 25 (1.2 and 0.9, respectively), while all the strains up-regulated ptsI (0.8-1.7). The manLMN operon encoding the EII^{man} complex was surprisingly strongly up-regulated during growth on ribose in all the strains (Table 1). By proteomic analysis, no regulation of the PTS enzymes was seen [19]. The expression of HPr and EI in L. sakei during growth on glucose or ribose was previously suggested to be constitutive [14], and in other lactobacilli, the EII^{man} complex was reported to be consistently highly expressed, regardless of carbohydrate source [72-74]. Notably, PEP-dependent phosphorylation of PTS sugars has been detected in ribose-grown cells, indicating that the EII^{man} complex is active, and since no transport and phosphorylation via EII^{man} occurs, the complex is phosphorylated, while it is unphosphorylated in the presence of the substrates of the EII^{man} complex [8,73]. The stimulating effect exerted by small amounts of glucose on ribose uptake in L. sakei, which has also been reported in other lactobacilli [74,75], was suggested to be caused by dephosphorylation of the PTS proteins in the presence of glucose, as a *ptsI* mutant lacking EI, as well as P-His-HPr, was shown to enhance ribose uptake [15,16,76]. Stentz et al. [15] observed that a *L. sakei* mutant (strain RV52) resistant to 2 deoxy-D-glucose, a glucose toxic analog transported by EIIman, and thus assumed to be affected in the EII^{man}, did not show the same enhanced uptake [15]. It was concluded that $\mathrm{EII}^{\mathrm{man}}$ is not involved in the PTS-mediated regulation of ribose metabolism in *L. sakei*. The mutation was though not reported verified by sequencing [15], and other mutations could be responsible for the observed phenotype. The *L.* sakei EIIAB^{man}, EIIC^{man} and EIID^{man} show 72, 81, and 82% identity, respectively, with the same enzymes in *L*. casei, in which mutations rendering the EII^{man} complex inactive were shown to derepress rbs genes, resulting in a loss of the preferential use of glucose over ribose [75]. Furthermore, in L. pentosus, EII^{man} was shown to provide a strong signal to the CcpA-dependent repression pathway [73]. The hprK gene encoding HPrK/P which controls the phosphorylation state of HPr was strongly upregulated (1.2-2.0) in all three strains. HPrK/P dephosphorylates P-Ser-HPr when the concentration of glycolytic intermediates drop, which is likely the situation during growth on ribose [20,22,24].

Numerous genes encoding hypothetical proteins with unknown function were also found to be differentially expressed (Table 1), as well as several other genes belonging to various functional categories. For most of these, their direct connection with ribose metabolism is unknown, and is likely an indirect effect.

Conclusions

The ability to ferment meat and fish is related to the capacity of the bacterium to rapidly take up the available carbohydrates and other components for growth. The importance of this process, especially to the meat industry, stimulates research aimed at understanding the mechanisms for transport and metabolism of these compounds, with the ultimate goal to be able to select improved strains. Genome-wide transcriptome analyses with DNA microarrays efficiently allowed the identification of genes differentially expressed between growth on the two carbohydrates which L. sakei can utilize from these substrates. Moreover, microarrays were a powerful tool to increase the understanding of the bacterium's primary metabolism and revealed a global regulatory mechanism. In summary, the ribose uptake and catabolic machinery is highly regulated at the transcription level, and it is closely linked with catabolism of nucleosides. A global regulation mechanism seems to permit a fine tuning of the expression of enzymes that control efficient exploitation of available carbon sources.

Additional material

Additional file 1: Table S3. Primer and probe sets used for qRT-PCR. Presents the primer and probe sets used for validation of microarray data by qRT-PCR analysis. Table S4. Comparison of microarray data with qRT-PCR results of *L. sakei* strain LS 25 grown on ribose compared with glucose. Presents gene regulation values (log₂) from the qRT-PCR analysis in comparison with microarray data.

Abbreviations

PKP: phosphoketolase pathway; PEP: phosphoenolpyruvate; PTS: PEP-dependent carbohydrate phosphotransferase system; CCR: carbon catabolite repression; *cre*: catabolite responsive element; RbsK: ribokinase; RbsD: D-Ribose pyranase; Xpk: xylulose-5-phosphate phosphoketolase; Ack: Acetate kinase, Pfk: 6-phosphofructokinase; Pyk: pyruvate kinase; PDC: pyruvate dehydrogenase complex; GlpD: glycerol-3-phosphate dehydrogenase; GlpK: glycerol kinase; ElI: enzyme II; El: enzyme I; HPr: histidine protein; HPrK/P: HPr kinase/phosphatase; DeoR: deoxyribonucleoside synthesis operon transcriptional regulator.

Acknowledgements and funding

This work was financially supported by Grant 159058/110 from the Norwegian Research Council. The authors would like to thank Monique Zagorec for helpful suggestions and critically reading the manuscript. We also thank Margrete Solheim, Mari Christine Brekke, and Signe Marie Drømtorp for their assistance during the experiments, and Hallgeir Bergum, the Norwegian Microarray Consortium (NMC), for printing the microarray slides.

Author details

¹Nofima Mat AS, Norwegian Institute of Food, Fisheries and Aquaculture Research, Osloveien 1, Ås, NO-1430, Norway. ²Department of Chemistry, Biotechnology and Food Science, Norwegian University of Life Sciences, P.O. Box 5003, Ås, NO-1432, Norway.

Authors' contributions

AM participated in the study design, conducted the experimental work, analyzed and interpreted data, and wrote the manuscript. LS conducted the statistical analysis. KN and LA conceived the study, participated in the study design process and reviewed the manuscript. All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

Received: 9 February 2011 Accepted: 24 June 2011 Published: 24 June 2011

References

- Hammes WP, Bantleon A, Min S: Lactic acid bacteria in meat fermentation. FEMS Microbiol Rev 1990, 87:165-174.
- 2. Hammes WP, Hertel C: **New developments in meat starter cultures.** *Meat Science* 1998, **49**:125-138.
- Bredholt S, Nesbakken T, Holck A: Protective cultures inhibit growth of Listeria monocytogenes and Escherichia coli O157:H7 in cooked, sliced, vacuum- and gas-packaged meat. Int J Food Microbiol 1999, 53:43-52.
- Bredholt S, Nesbakken T, Holck A: Industrial application of an antilisterial strain of Lactobacillus sakei as a protective culture and its effect on the sensory acceptability of cooked, sliced, vacuum-packaged meats. Int J Food Microbiol 2001, 66:191-196.
- Katikou P, Georgantelis D, Paleologos EK, Ambrosiadis I, Kontominas MG: Relation of biogenic amines' formation with microbiological and sensory attributes in *Lactobacillus*-inoculated vacuum-packed rainbow trout (*Oncorhynchus mykiss*) fillets. J Agric Food Chem 2006, 54:4277-4283.
- Vermeiren L, Devlieghere F, Debevere J: Evaluation of meat born lactic acid bacteria as protective cultures for biopreservation of cooked meat products. Int J Food Microbiol 2004, 96:149-164.
- Chaillou S, Champomier-Vergès MC, Cornet M, Crutz-Le Coq AM, Dudez AM, Martin V, Beaufils S, Darbon-Rongere E, Bossy R, Loux V, Zagorec M: The complete genome sequence of the meat-borne lactic acid bacterium *Lactobacillus sakei* 23 K. Nat Biotechnol 2005, 23:1527-1533.
- Lauret R, Morel-Deville F, Berthier F, Champomier-Vergès M, Postma P, Ehrlich SD, Zagorec M: Carbohydrate utilization in Lactobacillus sake. Appl Environ Microbiol 1996, 62:1922-1927.
- McLeod A, Nyquist OL, Snipen L, Naterstad K, Axelsson L: Diversity of Lactobacillus sakei strains investigated by phenotypic and genotypic methods. Syst Appl Microbiol 2008, 31:393-403.
- Chiaramonte F, Blugeon S, Chaillou S, Langella P, Zagorec M: Behavior of the meat-borne bacterium Lactobacillus sakei during its transit through

- the gastrointestinal tracts of axenic and conventional mice. *Appl Environ Microbiol* 2009, **75**:4498-4505.
- Dal Bello F, Walter J, Hammes WP, Hertel C: Increased complexity of the species composition of lactic acid bacteria in human feces revealed by alternative incubation condition. *Microb Ecol* 2003, 45:455-463.
- Walker A, Cerdeno-Tarraga A, Bentley S: Faecal matters. Nat Rev Microbiol 2006, 4:572-573.
- Chiaramonte F, Anglade P, Baraige F, Gratadoux JJ, Langella P, Champomier-Vergès MC, Zagorec M: Analysis of Lactobacillus sakei mutants selected after adaptation to the gastrointestinal tract of axenic mice. Appl Environ Microbiol 2010, 76:2932-2939.
- Stentz R, Lauret R, Ehrlich SD, Morel-Deville F, Zagorec M: Molecular cloning and analysis of the ptsHI operon in Lactobacillus sake. Appl Environ Microbiol 1997, 63:2111-2116.
- Stentz R, Cornet M, Chaillou S, Zagorec M: Adaption of Lactobacillus sakei to meat: a new regulatory mechanism of ribose utilization? INRA, EDP Sciences 2001, 81:131-138.
- Stentz R, Zagorec M: Ribose utilization in Lactobacillus sakei: analysis of the regulation of the rbs operon and putative involvement of a new transporter. J Mol Microbiol Biotechnol 1999. 1:165-173.
- Torriani S, Clementi F, Vancanneyt M, Hoste B, Dellaglio F, Kersters K: Differentiation of *Lactobacillus plantarum*, *L. pentosus* and *L. paraplantarum* species by RAPD-PCR and AFLP. Syst Appl Microbiol 2001, 24:554-560
- Claesson MJ, van Sinderen D, O'Toole PW: The genus Lactobacillus a genomic basis for understanding its diversity. FEMS Microbiol Lett 2007, 269:22-28.
- McLeod A, Zagorec M, Champomier-Vergès MC, Naterstad K, Axelsson L: Primary metabolism in *Lactobacillus sakei* food isolates by proteomic analysis. *BMC Microbiol* 2010, 10:120.
- Deutscher J, Francke C, Postma PW: How phosphotransferase systemrelated protein phosphorylation regulates carbohydrate metabolism in bacteria. Microbiol Mol Biol Rev 2006, 70:939-1031.
- Stulke J, Hillen W: Carbon catabolite repression in bacteria. Curr Opin Microbiol 1999, 2:195-201.
- 22. Titgemeyer F, Hillen W: Global control of sugar metabolism: a grampositive solution. *Antonie Van Leeuwenhoek* 2002, **82**:59-71.
- 23. Fujita Y: Carbon catabolite control of the metabolic network in *Bacillus subtilis*. *Biosci Biotechnol Biochem* 2009, **73**:245-259.
- Schumacher MA, Allen GS, Diel M, Seidel G, Hillen W, Brennan RG: Structural basis for allosteric control of the transcription regulator CcpA by the phosphoprotein HPr-Ser46-P. Cell 2004, 118:731-741.
- Obst M, Hehn R, Vogel RF, Hammes WP: Lactose metabolism in Lactobacillus curvatus and Lactobacillus sake. FEMS Microbiol Lett 1992, 97:209-214.
- 26. Montel MC, Champomier MC: Arginine catabolism in *Lactobacillus sake* isolated from meat. *Appl Environ Microbiol* 1987, **53**:2683-2685.
- Zuniga M, Champomier-Vergès M, Zagorec M, Pérez-Martinez G: Structural and functional analysis of the gene cluster encoding the enzymes of the arginine deiminase pathway of *Lactobacillus sake*. J Bacteriol 1998, 180:4154-4159.
- Rodionov DA, Mironov AA, Gelfand MS: Transcriptional regulation of pentose utilisation systems in the *Bacillus/Clostridium* group of bacteria. FEMS Microbiol Lett 2001, 205:305-314.
- Berthier F, Zagorec M, Champomier-Vergès MC, Ehrlich SD, Morel-Deville F: Efficient transformation of Lactobacillus sake by electroporation. Microbiol 1996, 142:1273-1279.
- 30. Hagen BF, Næs H, Holck AL: Meat starters have individual requirements for Mn2+. Meat Science 2000, 55:161-168.
- Møretrø T, Hagen BF, Axelsson L: A new, completely defined medium for meat lactobacilli. J Appl Microbiol 1998, 85:715-722.
- Nyquist OL, McLeod A, Brede DA, Snipen L, Nes IF: Comparative genomics of *Lactobacillus sakei* with emphasis on strains from meat. *Mol Genet Genomics* 2011, 285:297-311.
- Rud I, Naterstad K, Bongers RS, Molenaar D, Kleerebezem M, Axelsson L: Functional analysis of the role of CggR (central glycolytic gene regulator) in *Lactobacillus plantarum* by transcriptome analysis. *Microbial Biotechnology* 2011, 4:345-356.
- Vebø HC, Solheim M, Snipen L, Nes IF, Brede DA: Comparative genomic analysis of pathogenic and probiotic Enterococcus faecalis isolates, and

- their transcriptional responses to growth in human urine. *PLoS One* 2010, 5:e12489.
- 35. Smyth GK, Speed T: Normalization of cDNA microarray data. *Methods* 2003. **31**:265-273.
- Smyth GK: Linear models and empirical bayes methods for assessing differential expression in microarray experiments. Stat Appl Genet Mol Biol 2004. 3: Article3.
- Smyth GK, Michaud J, Scott HS: Use of within-array replicate spots for assessing differential expression in microarray experiments. Bioinformatics 2005, 21:2067-2075.
- Rode TM, Møretrø T, Langsrud S, Langsrud O, Vogt G, Holck A: Responses of Staphylococcus aureus exposed to HCl and organic acid stress. Can J Microbiol 2010, 56:777-792.
- Weickert MJ, Chambliss GH: Site-directed mutagenesis of a catabolite repression operator sequence in *Bacillus subtilis*. Proc Natl Acad Sci USA 1990, 87:6238-6242.
- Champomier-Vergès MC, Chaillou S, Cornet M, Zagorec M: Erratum to "Lactobacillus sakei: recent developments and future prospects". Res Microbiol 2002. 153:115-123.
- 41. Lorquet F, Goffin P, Muscariello L, Baudry JB, Ladero V, Sacco M, Kleerebezem M, Hols P: Characterization and functional analysis of the *poxB* gene, which encodes pyruvate oxidase in *Lactobacillus plantarum*. *J Bacteriol* 2004, **186**:3749-3759.
- Muscariello L, Marasco R, De Felice M, Sacco M: The functional ccpA gene is required for carbon catabolite repression in Lactobacillus plantarum. Appl Environ Microbiol 2001, 67:2903-2907.
- Branny P, De La Torre F, Garel JR: Cloning, sequencing, and expression in *Escherichia coli* of the gene coding for phosphofructokinase in *Lactobacillus bulgaricus*. J Bacteriol 1993, 175:5344-5349.
- 44. Viana R, Perez-Martinez G, Deutscher J, Monedero V: The glycolytic genes pfk and pyk from Lactobacillus casei are induced by sugars transported by the phosphoenolpyruvate:sugar phosphotransferase system and repressed by CcpA. Arch Microbiol 2005, 183:385-393.
- 45. Kandler O: Carbohydrate metabolism in lactic acid bacteria. *Antonie Van Leeuwenhoek* 1983, **49**:209-224.
- Naterstad K, Rud İ, Kvam I, Axelsson L: Characterisation of the gap operon from Lactobacillus plantarum and Lactobacillus sakei. Curr Microbiol 2007, 54:180-185.
- Kim I, Kim E, Yoo S, Shin D, Min B, Song J, Park C: Ribose utilization with an excess of mutarotase causes cell death due to accumulation of methylglyoxal. J Bacteriol 2004, 186:7229-7235.
- Weber J, Kayser A, Rinas U: Metabolic flux analysis of Escherichia coli in glucose-limited continuous culture. II. Dynamic response to famine and feast, activation of the methylglyoxal pathway and oscillatory behaviour. Microbiology 2005, 151:707-716.
- Totemeyer S, Booth NA, Nichols WW, Dunbar B, Booth IR: From famine to feast: the role of methylglyoxal production in *Escherichia coli*. Mol Microbiol 1998, 27:553-562.
- Malleret C, Lauret R, Ehrlich SD, Morel-Deville F, Zagorec M: Disruption of the sole *IdhL* gene in *Lactobacillus sakei* prevents the production of both L- and D-lactate. *Microbiology* 1998, 144:3327-3333.
- Axelsson L: Lactic acid bacteria: classification and physiology. In Lactic acid bacteria: microbiological and functional aspects.. Third revised and expanded edition. Edited by: Salminen S, von Wright A, Ouwehand A. New York, USA: Marcel Dekker, Inc./CRC Press; 2004:1-66.
- 52. Condon S: Responses of lactic acid bacteria to oxygen. FEMS Microbiol Rev 1987, 46:269-280.
- 53. Fridovich I: The biology of oxygen radicals. Science 1978, 201:875-880.
- Rodionov DA, Vitreschak AG, Mironov AA, Gelfand MS: Comparative genomics of thiamin biosynthesis in procaryotes. New genes and regulatory mechanisms. J Biol Chem 2002, 277:48949-48959.
- Jordan A, Reichard P: Ribonucleotide reductases. Annu Rev Biochem 1998, 67:71-98
- Keeney KM, Stuckey JA, O'Riordan MX: LplA1-dependent utilization of host lipoyl peptides enables *Listeria* cytosolic growth and virulence. *Mol Microbiol* 2007, 66:758-770.
- Marceau A, Zagorec M, Chaillou S, Mera T, Champomier-Vergès MC: Evidence for involvement of at least six proteins in adaptation of Lactobacillus sakei to cold temperatures and addition of NaCl. Appl Environ Microbiol 2004, 70:7260-7268.

- 58. Grogan DW, Cronan JE Jr: Cyclopropane ring formation in membrane lipids of bacteria. *Microbiol Mol Biol Rev* 1997, **61**:429-441.
- Schujman GE, Guerin M, Buschiazzo A, Schaeffer F, Llarrull LI, Reh G, Vila AJ, Alzari PM, de Mendoza D: Structural basis of lipid biosynthesis regulation in Gram-positive bacteria. Embo J 2006, 25:4074-4083.
- Mahr K, Hillen W, Titgemeyer F: Carbon catabolite repression in Lactobacillus pentosus: analysis of the ccpA region. Appl Environ Microbiol 2000. 66:277-283.
- Nentwich SS, Brinkrolf K, Gaigalat L, Huser AT, Rey DA, Mohrbach T, Marin K, Puhler A, Tauch A, Kalinowski J: Characterization of the Lacttype transcriptional repressor RbsR controlling ribose transport in Corynebacterium glutamicum ATCC 13032. Microbiology 2009, 155:150-164.
- Muller W, Horstmann N, Hillen W, Sticht H: The transcription regulator RbsR represents a novel interaction partner of the phosphoprotein HPr-Ser46-P in *Bacillus subtilis*. Febs J 2006, 273:1251-1261.
- Perez-Rueda E, Collado-Vides J: The repertoire of DNA-binding transcriptional regulators in Escherichia coli K-12. Nucleic Acids Res 2000, 28:1838-1847
- 64. Brinkrolf K, Ploger S, Solle S, Brune I, Nentwich SS, Huser AT, Kalinowski J, Puhler A, Tauch A: The Lacl/GalR family transcriptional regulator UriR negatively controls uridine utilization of Corynebacterium glutamicum by binding to catabolite-responsive element (cre)-like sequences. Microbiology 2008, 154:1068-1081.
- Saxild HH, Andersen LN, Hammer K: dra-nupC-pdp operon of Bacillus subtilis: nucleotide sequence, induction by deoxyribonucleosides, and transcriptional regulation by the deoR-encoded DeoR repressor protein. J Bacteriol 1996, 178:424-434.
- Zeng X, Saxild HH: Identification and characterization of a DeoR-specific operator sequence essential for induction of dra-nupC-pdp operon expression in Bacillus subtilis. J Bacteriol 1999, 181:1719-1727.
- Zeng X, Saxild HH, Switzer RL: Purification and characterization of the DeoR repressor of Bacillus subtilis. J Bacteriol 2000, 182:1916-1922.
- Schuch R, Garibian A, Saxild HH, Piggot PJ, Nygaard P: Nucleosides as a carbon source in *Bacillus subtilis*: characterization of the *drm-pupG* operon. *Microbiology* 1999, 145:2957-2966.
- 69. Posthuma CC, Bader R, Engelmann R, Postma PW, Hengstenberg W, Pouwels PH: Expression of the xylulose 5-phosphate phosphoketolase gene, xpkA, from Lactobacillus pentosus MD363 is induced by sugars that are fermented via the phosphoketolase pathway and is repressed by glucose mediated by CcpA and the mannose phosphoenolpyruvate phosphotransferase system. Appl Environ Microbiol 2002, 68:831-837.
- Charrier V, Buckley E, Parsonage D, Galinier A, Darbon E, Jaquinod M, Forest E, Deutscher J, Claiborne A: Cloning and sequencing of two enterococcal glpK genes and regulation of the encoded glycerol kinases by phosphoenolpyruvate-dependent, phosphotransferase systemcatalyzed phosphorylation of a single histidyl residue. J Biol Chem 1997, 272:14166-14174.
- Darbon E, Servant P, Poncet S, Deutscher J: Antitermination by GlpP, catabolite repression via CcpA and inducer exclusion triggered by P-GlpK dephosphorylation control *Bacillus subtilis glpFK* expression. *Mol Microbiol* 2002, 43:1039-1052.
- Barrangou R, Azcarate-Peril MA, Duong T, Conners SB, Kelly RM, Klaenhammer TR: Global analysis of carbohydrate utilization by Lactobacillus acidophilus using cDNA microarrays. Proc Natl Acad Sci USA 2006. 103:3816-3821.
- Chaillou S, Postma PW, Pouwels PH: Contribution of the phosphoenolpyruvate:mannose phosphotransferase system to carbon catabolite repression in *Lactobacillus pentosus*. *Microbiology* 2001, 147:671-679.
- Veyrat A, Gosalbes MJ, Perez-Martinez G: Lactobacillus curvatus has a glucose transport system homologous to the mannose family of phosphoenolpyruvate-dependent phosphotransferase systems. Microbiology 1996, 142:3469-3477.
- Veyrat A, Monedero V, Perez-Martinez G: Glucose transport by the phosphoenolpyruvate:mannose phosphotransferase system in *Lactobacillus casei* ATCC 393 and its role in carbon catabolite repression. *Microbiology* 1994, 140:1141-1149.
- Viana R, Monedero V, Dossonnet V, Vadeboncoeur C, Perez-Martinez G, Deutscher J: Enzyme I and HPr from Lactobacillus casei: their role in

sugar transport, carbon catabolite repression and inducer exclusion. *Mol Microbiol* 2000, **36**:570-584.

doi:10.1186/1471-2180-11-145

Cite this article as: McLeod *et al.*: Global transcriptome response in *Lactobacillus sakei* during growth on ribose. *BMC Microbiology* 2011

Submit your next manuscript to BioMed Central and take full advantage of:

- Convenient online submission
- Thorough peer review
- No space constraints or color figure charges
- Immediate publication on acceptance
- Inclusion in PubMed, CAS, Scopus and Google Scholar
- Research which is freely available for redistribution

Submit your manuscript at www.biomedcentral.com/submit

