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Functional Divergence in the Genus *Oenococcus* as Predicted by Genome Sequencing of the Newly-Described Species, *Oenococcus kitaharae*

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Abstract

*Oenococcus kitaharae* is only the second member of the genus *Oenococcus* to be identified and is the closest relative of the industrially important wine bacterium *Oenococcus oeni*. To provide insight into this new species, the genome of the type strain of *O. kitaharae*, DSM 17330, was sequenced. Comparison of the sequenced genomes of both species show that the genome of *O. kitaharae* DSM 17330 contains many genes with predicted functions in cellular defence (bacteriocins, antimicrobials, restriction-modification systems and a CRISPR locus) which are lacking in *O. oeni*. The two genomes also appear to differentially encode several metabolic pathways associated with amino acid biosynthesis and carbohydrate utilization and which have direct phenotypic consequences. This would indicate that the two species have evolved different survival techniques to suit their particular environmental niches. *O. oeni* has adapted to survive in the harsh, but predictable, environment of wine that provides very few competitive species. However *O. kitaharae* appears to have adapted to a growth environment in which biological competition provides a significant selective pressure by accumulating biological defence molecules, such as bacteriocins and restriction-modification systems, throughout its genome.

Introduction

*Oenococcus kitaharae* is a lactic acid bacterium (LAB) that was recently isolated from composting distilled Shochu residue [1]. This species represents only the second member of the genus *Oenococcus* to be identified, with *Oenococcus oeni*, the founding member of this genus, being reclassified from *Leuconostoc oenos* by Dicks et al in 1995 [2]. Whereas little is known regarding the biology or ecology of *O. kitaharae*, *O. oeni* plays a pivotal role in the production of wine (its almost exclusive habitat) where it is involved in defence against both bacteriophage (restriction-modification and CRISPR) and other microorganisms (bacteriocins), and has had its genome populated by at least two conjugative transposons, which is in contrast to currently available genome sequences of *O. oeni* which lack the vast majority of these defence proteins. It therefore appears that the genome of *O. kitaharae* has been shaped by its need to survive in a competitive growth environment that is vastly different from that encountered by *O. oeni*, where environmental stresses provide the greatest challenge to growth and reproduction.

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**Abstract**

*Oenococcus kitaharae* is only the second member of the genus *Oenococcus* to be identified and is the closest relative of the industrially important wine bacterium *Oenococcus oeni*. To provide insight into this new species, the genome of the type strain of *O. kitaharae*, DSM 17330, was sequenced. Comparison of the sequenced genomes of both species show that the genome of *O. kitaharae* DSM 17330 contains many genes with predicted functions in cellular defence (bacteriocins, antimicrobials, restriction-modification systems and a CRISPR locus) which are lacking in *O. oeni*. The two genomes also appear to differentially encode several metabolic pathways associated with amino acid biosynthesis and carbohydrate utilization and which have direct phenotypic consequences. This would indicate that the two species have evolved different survival techniques to suit their particular environmental niches. *O. oeni* has adapted to survive in the harsh, but predictable, environment of wine that provides very few competitive species. However *O. kitaharae* appears to have adapted to a growth environment in which biological competition provides a significant selective pressure by accumulating biological defence molecules, such as bacteriocins and restriction-modification systems, throughout its genome.
Results

The *O. kitaharae* DSM 17330 genome was assembled from 1×10^6 Illumina paired-end reads (500 bp spacing) into 17 contigs, comprising 12 single-copy contigs in addition to five contigs that were present in two copies each (with three of these having 100% identity). Through the application of paired-end information and the precise order of the unique sequences bounding these repeats, these 17 contigs were able to be manually arranged into two replicons (Fig. 1). The first of these is a 1.84 Mb circular chromosome which, due to the presence of a highly repetitive repeat cluster that is associated with the coding region of a serine-repeat protein, contains one assembly gap. The second replicon is predicted to be an 8.3 kb plasmid, which, on the basis of sequencing coverage, is predicted to be present in low copy number (~2 copies per cell, data not shown).

Genome annotation using a combination of Glimmer [7] and RAST [8] identified a total of 1833 predicted open reading frames (ORFs), four rRNA genes (two copies of each of the large and small ribosomal subunits), which are identical in sequence to the previously published ribosomal sequences for *O. kitaharae* [1], and 44 tRNA genes (see Table S1).

Comparative genomics of *O. kitaharae* and LAB

While the genome size of *O. kitaharae* is similar to that of *O. oeni* PSU1 (1.84 Mb vs 1.78 Mb), initial estimates of DNA-DNA relatedness were only 25% to 30% [1]. By direct comparison of the available genome sequences of *O. kitaharae* and *O. oeni*, a similar picture emerged. As expected, the majority of the ORFs predicted in the *O. kitaharae* genome (62%) have their closest homolog in *O. oeni* supporting the classification of *O. kitaharae* DSM 17330 within the *Oenococcus* genus (Fig. 2A). Of the remaining 703 *O. kitaharae* ORFs, 487 have at least one match in the Genbank non-redundant protein database, with LAB such as *Lactobacillus* spp, *Leuconostoc* spp, *Weissella* spp and *Lactococcus* spp predominating (Fig. 2A), while the remainder display no recognizable homolog.
and either represent novel protein sequences or false positives of the ORF prediction methodology applied.

In order to compare the phylogenetic position of O. kitaharae as determined from 16S rDNA sequencing [1] with that based upon whole genome data, homologs of each of the predicted ORFs of O. kitaharae were sought from whole genome sequence annotations of thirteen strains of LAB representing the genera Enterococcus, Lactobacillus, Lactococcus, Leuconostoc, Oenococcus, Pediococcus and Weissella. A total of 561 of the O. kitaharae ORFs were shown to be conserved across all thirteen strains (BLAST e-value<10^{-20}, minimum 50% coverage of the query protein). Of these, 95 were then selected based on high sequence conservation and lack of potential paralogous sequences (see Table S2 for a complete list of the protein sequences used). Each of the subsequent individual protein alignments

Figure 2. Evolutionary relationship of Oenococcus kitaharae and members of the LAB family. (A) The distribution of BLAST best-hits by genus for each ORF predicted in the O. kitaharae genome. (B) Whole genome phylogenetic relationship between O. kitaharae and other LAB based upon a conserved group of 95 proteins.

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produced from these conserved groups of ORFs were then concatenated and used to construct a single maximum-likelihood phylogeny (Fig. 2B). The result of this analysis was consistent with phylogenies based upon 16S rDNA [1,9,10] and positions. O. kitaharae as a clear sister species to O. oeni with both Leuconostoc spp. and Weissella spp. being the next closest evolutionary relatives.

Chromosomal elements of “foreign” origin. The majority of the O. kitaharae genome is conserved with that of O. oeni with the exception of several large islands (Fig. 3). In all but a limited number of cases, these O. kitaharae-specific islands were also shown to lack identifiable homology with other species of LAB used in the phylogenetic constructions. In addition, the majority of these regions display a high probability of being acquired by horizontal gene transfer (HGT) [11] and would be expected to have entered the O. kitaharae genome via genetic elements such as bacteriophage or conjugative plasmids.

As previous investigations between strains of O. oeni have shown that non-conserved genomic islands can often be attributed to the differential presence of prophage elements [12], homology searches were used to identify classical prophage genes, such as those that encode conserved bacteriophage integrase and lyase proteins, in the genome of DSM 17330. While several O. kitaharae ORFs were found to be homologous to phage proteins (Table S1), each potential prophage region lacked the repertoire of proteins which would be expected for the presence of functional prophage elements [12]. In addition, in all but one case (the genomic region from 382388 bp–404717 bp), these genomic islands were not found downstream of tRNA genes as is observed for O. oeni phage, which use tRNA genes as attachment sites [12].

Whereas the non-conserved genomic islands in the O. kitaharae genome do not appear to encode active phage, two of these islands have the hallmarks of genically-integrated conjugative transposons (Fig. 4). Conjugative transposons are DNA elements that combine features of bacteriophage, transposons and conjugative plasmids [13]. These elements are often found integrated into the genome and are able to transpose to new sites in the host genome via the formation of a circular intermediate, which can sometimes replicate as a plasmid. As the name suggests, conjugative transposons also encode the ability to be horizontally transferred to new cells (and across species boundaries) via the classical conjugation pathway [13]. As yet, there have been no documented cases of conjugative transposons found in the genome of any strain of O. oeni.

The two potential conjugative transposons of O. kitaharae share a conserved core of 12 proteins. These proteins are predicted to encode the machinery necessary for conjugation and are highly homologous to a 12 kb region of the conjugative plasmid pWCS103 from Lactobacillus plantarum [14]. In addition to the conserved conjugative core, one transposon is predicted to encode transposase-based integrative functions and the ability to replicate as a plasmid via the RepA and RepB plasmid replication proteins, whereas the other appears to use bacteriophage-based proteins for integration into the genome.
Accompanying the replicative functions of the *O. kitaharae* conjugative transposons, each element is predicted to encode several proteins that can be broadly categorised as functioning as part of “cellular defence” from either foreign DNA or from competition imposed by other microorganisms. One conjugative transposon is predicted to encode two separate restriction-modification (RM) systems (one combined Type III enzyme, and one Type II RM pair) and the other contains at least five ORFs which are potentially involved in the production of the antibacterial compounds (Fig. 4). In addition to the gene content of these potentially transmissible elements, the *O. kitaharae* DSM 17330 genome encodes another twelve proteins with putative roles in bacteriocin/antibacterial manufacture, transport or detoxification, four proteins involved in DNA RM and one CRISPR pathway array which, in other bacterial species, has been shown to provide a memory-based immunity to bacteriophage infection and possibly to the transmission of plasmid DNA [reviewed in (15)] (Table 1). All of these cellular defence mechanisms are lacking conserved homologs in the sequenced strains of *O. oeni*, with the exception of a single Type III RM enzyme which is found specifically in *O. oeni* strain AWRI429 [6].

**Phenotypic differences attributable to genomic variation**

In the initial characterization of *O. kitaharae* [1] several phenotypic traits were noted that differentiate this new species from *O. oeni*. Comparative genomics reveals a basis for some of these known differences while also suggesting several additional points of phenotypic differentiation.

**Sugar utilization.** One of the defining biochemical differences between *O. kitaharae* and *O. oeni* that was noted in its original isolation was the ability of *O. kitaharae* to produce acid from maltose [16,17]. By comparing available whole-genome annotations for *O. oeni* with *O. kitaharae* DSM 17330 [8], it was possible to identify several genes associated with sugar utilization that are differentially present across the species (Table 2). Of these, at least four genes which are present in *O. kitaharae*, but absent in the *O. oeni* genomes, are predicted to be involved in the utilization of maltose, providing a direct genetic basis for this phenotype. In addition to genes predicted to be involved in the species-specific utilization of maltose, there are several ORFs predicted to be involved in the metabolism of trehalose, D-glucuronate, D-ribose and fructose that are specifically present in *O. kitaharae*. While the assimilation of these sugars is often carried out by specific strains of *O. oeni* [17], this genotypic data agrees well with biochemical tests performed previously that indicated that *O. kitaharae* was able to utilize all of these various carbon sources [1,17].

In addition to those genes that are specifically present in *O. kitaharae* DSM 17330, several were identified that were present only in strains of *O. oeni* and which are predicted to be involved in the uptake and metabolism of arabinose and xylose (Table 2). This is consistent with the inability of *O. kitaharae* to produce acid from either L-arabinose or D-xylose, two biochemical reactions that many strains of *O. oeni*, including those for which genome sequence are available often perform [1,17].

**Amino acid biosynthesis.** Both *O. oeni* and *O. kitaharae* are fastidious microorganisms that require many exogenous vitamins and amino acids. However, it appears that the *O. kitaharae* genome encodes biosynthetic pathways for at least two amino acids, arginine and histidine, which are lacking in *O. oeni* [18].

The *O. kitaharae* DSM 17330 genome encodes the six genes necessary for the production of arginine from glutamate via the ornithine/carbamoyl-phosphate (CP) pathway, in addition to encoding a second set of carbamoyl-phosphate synthase (CPSase) enzymes that are predicted to be involved in the synthesis of histidine from the pentose phosphate pathway intermediate 5-phosphoribosyl 1-pyrophosphate (PRPP) (Table 3). As CP is an important intermediate in both the arginine and pyrimidine biosynthetic pathways, many bacteria, such as *Lactobacillus plantarum*, contain two completely separate sets of CPSase proteins [19]. In this situation, one protein is encoded in an operon with genes involved in arginine biosynthesis and regulated by arginine, while a second gene is located in the pyrimidine biosynthetic operon and regulated by exogenous pyrimidines. *O. kitaharae* contains both sets of CPSase enzymes while the *O. oeni* genomic sequences are predicted to encode only the single pyrimidine-associated CPSase [19].

*O. kitaharae* also appears to encode all of the enzymes necessary for the synthesis of histidine from the pentose phosphate pathway intermediate 5-phosphoribosyl 1-pyrophosphate (PRPP) (Table 3). These 11 genes lie adjacent to each other in the *O. kitaharae* DSM 17330 genome while *O. oeni* genome sequences lack this entire complement of enzymes.

**Organic acid metabolism.** Many strains of *O. oeni* are capable of fermenting citrate when additional sources of fermentable carbon sources are also present [20,21]. Several...
genes in the *O. oeni* PSU-1 genome have been identified previously that would provide this strain with the ability to convert citrate to pyruvate [4] (Table 4). These genes are absent from the *O. kitaharae* DSM 17330 genome leading to the prediction that, unlike *O. oeni*, this strain would lack the ability to ferment this organic acid.

One of the key defining biochemical features that separates *O. oeni* from *O. kitaharae* is the ability to perform malolactic fermentation. Malolactic fermentation has been shown to require the action of three proteins, a malate permease, which transports malate into the cell, the malolactic enzyme, which is responsible for converting malic acid into lactic acid, and a regulatory protein for these two downstream genes [22]. Surprisingly, the *O. kitaharae* genome was shown to contain genes that are orthologous to those which encode all three of these activities in *O. oeni* (Fig. 5A). It was subsequently shown that, while the sequences of all three genes are present in the *O. kitaharae* genome, the gene encoding malolactic enzyme contained a nonsense mutation that would prematurely truncate the protein coding region (Fig. 5B). The alteration of a single base in this premature stop codon would be sufficient to restore the full-length malolactic enzyme coding region (Fig. 5B) that is highly conserved with malolactic enzymes from many bacteria (Fig. S1). Furthermore, the *O. kitaharae* gene was shown to have a low ratio of non-synonymous to synonymous mutations (*dN/dS* = 0.0123) when compared with its *O. oeni* homologue. This would indicate that there has been limited opportunity for the unconstrained accumulation of synonymous mutations in the two fragments of the malolactic enzyme coding region in *O. kitaharae* (as would be expected in a non-functional gene undergoing random drift). It is therefore likely that the conversion of the malolactic enzyme to a pseudogene is a very recent event in *O. kitaharae* and it may be possible to obtain a functional enzyme through reversion of the nonsense mutation or to find a functional malate pathway in strains of *O. kitaharae* other than DSM 17330.

**Discussion**

*O. kitaharae* and *O. oeni* comprise the only known members of their genus. Sequencing of the *O. kitaharae* DSM 17330 genome has provided important insights into the genetic diversity across this genus. These two species of *Oenococcus* appear to inhabit significantly different ecological niches, with *O. oeni* being found almost exclusively in the highly stressful environment of wine whereas *O. kitaharae* was isolated from a composting shochu residue of unknown nutrient composition. Accordingly, the two species have accumulated genetic adaptations that reflect different metabolic needs and environmental constraints.

Although little is known regarding the exact nutrient profile of the residue from which *O. kitaharae* was isolated, the average composition of the major wine metabolites are well known. Finished wine has little or no glucose, fructose or maltose but does contain

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**Table 1. Genes from *Oenococcus kitaharae* predicted to be involved in cellular defence.**

| Function                          | Description                               | ORF(s)     |
|-----------------------------------|-------------------------------------------|------------|
| Bacteriocin production or immunity| Putative bacteriocin ABC transporter       | OKIT_0291  |
|                                   | Bacteriocin immunity-associated integral  | OKIT_0292  |
|                                   | membrane protein                          | OKIT_0293  |
|                                   | Bacteriocin, lactococcin 972 family       | OKIT_0294  |
|                                   | Bacitracin transport ATP-binding protein   | OKIT_0298  |
|                                   | Putative blasticidin S deaminase 2C       | OKIT_0304  |
|                                   | Putative bacteriocin transport accessory protein | OKIT_0665 |
|                                   | Streptolysin S biosynthesis protein B (SagB) | OKIT_0885  |
|                                   | Streptolysin S biosynthesis protein C (SagC) | OKIT_0886  |
|                                   | Streptolysin S biosynthesis protein D (SagD) | OKIT_0887  |
|                                   | Phenazine biosynthesis protein PhzF        | OKIT_0912  |
|                                   | Lactococcin A immunity protein             | OKIT_0790  |
|                                   | Nisin transport protein                    | OKIT_0796  |
|                                   | Bacitracin export permease protein         | OKIT_1723  |
|                                   | Putative bacteriocin ABC transporter       | OKIT_1725  |
| Restriction modification           | Type III restriction enzyme 2C res subunit: DEAD/DEAH box helicase 2C N-terminal | OKIT_0515  |
|                                   | Type II modification methyltransferase     | OKIT_0538  |
|                                   | Type II restriction endonuclease           | OKIT_0539  |
|                                   | Type I restriction-modification system 2C restriction subunit R (EC 3.1.21.3);Ontology_term = KEGG_ENZYME:3.1.21.3 | OKIT_0971  |
|                                   | Type II restriction modification enzyme methyltransferase | OKIT_0974  |
|                                   | Type III restriction enzyme, restriction subunit | OKIT_0978  |
|                                   | S-methylcytosine-specific restriction enzyme* | OKIT_1348  |
| CRISPR                            | CRISPR-associated protein 2C SAG0894 family | OKIT_1269  |
|                                   | CRISPR-associated protein Cas1             | OKIT_1270  |
|                                   | CRISPR-associated protein Cas2             | OKIT_1271  |
|                                   | CRISPR-associated Csn2 family protein      | OKIT_1272  |

*a homolog found in *O. oeni* AWRIB429.

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*Comparative Genomics of Oenococcus kitaharae*
significant quantities of arabinose and xylose [23,24]. Many strains of O. oeni are capable of exploiting these carbohydrates, and contain genes whose biochemical functions are consistent with this ability, but, in all but a limited number of cases [16,17], cannot utilize maltose. In comparison, O. kitaharae DSM 17330 lacks the genes required for the use of arabinose and xylose but has the ability to utilize the maltose that would be present in the feedstocks, such as barley, which are used in the production of shochu.

In addition, while little is known regarding the amino acid profile of the shochu residue in which O. kitaharae was isolated, it would be predicted that these feedstocks would generally be lower in arginine and histidine than wine (where they are often amongst the most prevalent amino acids [24]) given the presence of the biosynthetic pathways for both of these amino acids in O. kitaharae. Interestingly, both the histidine and arginine biosynthetic pathways display a scattered pattern of presence throughout the LAB phylogeny with only a limited number of species within a genus possessing these pathways (Fig. S2). It appears that there must be significant selective pressure working for and against these biosynthetic pathways in an environmentally-dependent manner across the LAB. For O. kitaharae, the evolutionary origins of both pathways are more consistent with loss of these enzymes followed by horizontal gain from other species of bacteria (bacteriocin immunity proteins) and to also defend against invasion by foreign DNA, such as that introduced by bacteriophage (restriction-modification systems and the CRISPR element). Although the biological diversity of composting residue in which O. kitaharae was not formally evaluated, it can be assumed that there was sufficient microbiological competition for resources to justify the selective advantage for the presence of these defence compounds. This argument is further supported by the fact that at least two other novel species of LAB have been isolated from this environment in addition to many other species of LAB that have been shown to be present during the shochu production process [1,26–29]. O. kitaharae has therefore evolved to compete in a mixed-species environment whereas O. oeni has adapted to a niche in which the extreme nature of the growth substrate has removed the majority of biological competition.

Whereas the applicability of O. kitaharae for use as an industrial species is yet to be determined, it could prove useful for the development of improved strains of its relative O. oeni. Despite the environment of wine providing protection from competition for O. oeni, it is still argued that many instances of failed malolactic fermentations are due to the action of bacteriophage on susceptible strains [30,31]. If it were possible to move genes, such as those of the CRISPR array, from O. kitaharae into O. oeni via the conjugation machinery which is predicted to be present in O. kitaharae, this could provide a non-GM means of equipping industrial O. oeni strains with general resistance to bacteriophage infection. Likewise, if the genes involved in antimicrobial production can be transferred to O. oeni, these strains could limit potential negative impacts on wine quality.
due to the growth of spoilage bacteria such as *Pediococcus* spp, *Lactobacillus* spp and acetic acid bacteria, and may provide the means to reduce the amount of sulfite that is currently used for the microbial stabilization of wine [32]. The use of genes from *O. kitaharae* may therefore allow the production of strains of *O. oeni* which are not only able to thrive in the harshness of the wine environment, but are more resistant to potential biological competition from bacteriophage or other microorganisms.

**Materials and Methods**

**DNA isolation and sequencing**

*O. kitaharae* DSM 17330 was obtained from DSMZ (Germany) and was grown in modified MRS media (Amyl, Australia). Genomic DNA was isolated using standard phenol-chloroform extractions. Sequencing was performed on an Illumina GAIIx using 2×100 bp paired-end ends with an average library size of 500 bp (Ramaciotti Centre, NSW, Australia).

**Genome assembly**

A total of 990,000 reads (~50-fold expected genome coverage) were randomly selected and assembled using MIRA (version 3.2.1). The MIRA output was imported into Seqman Pro (DNAstar, Madison, WI) for manual alignment and editing of the assembly. This Whole Genome Shotgun project has been deposited at DDBJ/EMBL/GenBank under the accession AFVZ00000000. The version described in this paper is the first version, AFVZ01000000.

**Genome Annotation**

Gene predictions were made using Glimmer 3.02 [7]. Gene functional annotations were performed using the RAST server [8].

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**Table 3.** Genes involved in arginine and histidine biosynthesis in *O. kitaharae*.

| RAST Pathway            | Description                                           | ORF(s)       |
|-------------------------|-------------------------------------------------------|--------------|
| Arginine biosynthesis   | Acetylglutamate kinase (EC 2.7.2.8)                   | OKIT_0634    |
|                         | Acetylornithine aminotransferase (EC 2.6.1.11)        | OKIT_0630    |
|                         | Glutamate N-acetyltransferase (EC 2.3.1.35)           | OKIT_0629    |
|                         | N-acetyl-gamma-glutamyl-phosphate reductase (EC 1.2.1.38) | OKIT_0628 |
|                         | N-acetylglutamate synthase (EC 2.3.1.1)               | OKIT_0629    |
|                         | Ornithine carbamoyltransferase (EC 2.1.3.3)           | OKIT_0631    |
|                         | Carbamoyl-phosphate synthase small chain (EC 6.3.5.5) | OKIT_0632    |
|                         | Carbamoyl-phosphate synthase large chain (EC 6.3.5.5) | OKIT_0633    |
| Histidine biosynthesis  | phosphoribosyl-AMP cyclohydrolase( EC:3.5.4.19 )     | OKIT_1691    |
|                         | Phosphoribosyl-ATP pyrophosphatase (EC 3.6.1.31)      | OKIT_1690    |
|                         | Imidazoleglycerol-phosphate dehydratase (EC 4.2.1.19)| OKIT_1695    |
|                         | Imidazole glycerol phosphate synthase cyclase subunit (EC 4.1.3.1) | OKIT_1692 |
|                         | Imidazole glycerol phosphate synthase amidotransferase subunit (EC 2.4.2.1) | OKIT_1694 |
|                         | Histidinol-phosphate aminotransferase (EC 2.6.1.9)    | OKIT_1689    |
|                         | Histidinol-phosphatase (EC 3.1.3.15)                  | OKIT_1699    |
|                         | Histidinol dehydrogenase (EC 1.1.1.23)               | OKIT_1696    |
|                         | ATP phosphoribosyltransferase regulatory subunit (EC 2.4.2.17) | OKIT_1698 |
|                         | ATP phosphoribosyltransferase catalytic subunit (EC 2.4.2.17) | OKIT_1697 |
|                         | PhosphoribosylformiminonoS-aminomimidazole carboxamide ribotide isomerase (EC 5.3.1.16) | OKIT_1693 |

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**Table 4.** Genes involved in citrate utilization in *O. oeni*.

| RAST Pathway            | Description                                           | ORF(s)       |
|-------------------------|-------------------------------------------------------|--------------|
| Citrate Metabolism, Transport, and Regulation | 2-(5’-triphosphorylsilyl)-3’-dephosphoenzyme-A synthase (EC 2.7.8.25) | YP_810049.1* |
|                         | Apo-citrate lyase phosphoribosyl-dephospho-CoA transferase (EC 2.7.7.61) | YP_810048.1* |
|                         | Citrate lyase alpha chain (EC 4.1.3.6)                | YP_810047.1* |
|                         | Citrate lyase beta chain (EC 4.1.3.6)                | YP_810046.1* |
|                         | Citrate lyase gamma chain, acyl carrier protein (EC 4.1.3.6) | YP_810045.1* |
|                         | Citrate lyase transcriptional regulator CitI          | YP_810041.1* |
|                         | Oxaloacetate decarboxylase involved in citrate fermentation (EC 4.1.1.3) | YP_810042.1* |
|                         | [Citrate [pro-3S]-lyase] ligase (EC 6.2.1.22)         | YP_810044.1* |

*O. oeni* Genbank protein ID from genome accession number NC_008528.1.

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and BLAST [33] with comparisons to the non-redundant Genbank database. Predictions of genomic regions likely to have been acquired by horizontal gene transfer were calculated using Alien Hunter [11]. dN/dS ratios were calculated using Pal2nal [34]. Circular genome plots were compiled using Artemis [35] and DNAplotter [36]. Comparisons to the various LAB genomes were performed using BLAST [33] and custom written scripts. For phylogenetic analysis, proteins used for the analysis were first screened to ensure that they were conserved (minimum 60% identity when compared to the homologous \(O.\) kitaharae protein) in all of the LAB genomes used in this study (See Table S2). Next, proteins which had potential paralogs (which could confound the phylogeny) were identified by assigning each protein to specific orthoMCL [37] clusters and then only retaining those groups of orthologs in which each protein was the only member of a particular orthoMCL group. Individual protein alignments were then performed on each set of homologous sequences using Muscle [38]. These individual alignments were then concatenated into a single large sequence for each strain which was used to construct a maximum-likelihood phylogenetic tree using PhyML [39].

**Supporting Information**

**Figure S1** Amino acid alignment of malolactic enzymes from various species of lactic acid bacteria. Amino acid sequences were aligned using ClustalX and conserved residues (>60%) are highlighted (black shading). The position of the in-frame stop-codon in \(O.\) kitaharae is also highlighted (red shading). (TIF)

**Figure S2** Phylogenetetic relationship of the histidine and arginine operons of \(O.\) kitaharae. ORFs from each pathway in each species (if present) were concatenated prior to alignment. Each maximum-likelihood tree is presented for comparison at the same scale as the full tree which is comprised of 95 conserved ORFS from 13 species of lactic acid bacteria. (TIF)

**Table S1** \(Oenococcus\) kitaharae genome annotation. \(O.\) kitaharae ORFs as predicted by Glimmer [7] are matched against an automated annotation and functional prediction performed using RAST [8]. Comparative analysis of each ORF was also performed using BLASTp comparisons against \(O.\) oeni PSU1 [33] with the comparisons parameters listed for each \(O.\) kitaharae ORF and its closest \(O.\) oeni PSU1 match (if present). (XLS)

**Table S2** Evolutionarily conserved ORFs in lactic acid bacteria (LAB). Homologs of each of the predicted \(O.\) kitaharae ORFs were sought from thirteen strains of LAB using BLASTp [33]. Evolutionarily conserved proteins were classified as exhibiting sequence conservation across all fourteen species (minimum 60% identity when compared to the homologous \(O.\) kitaharae protein). This list was further refined by mapping each ORF from each species to an orthoMCL group [37] and only retaining those ORFs which were found be be unique within their particular orthoMCL cluster (removes the potential for paralogous ORFs to interfere with the phylogeny). (XLS)

**Author Contributions**

Conceived and designed the experiments: AB PC EB. Performed the experiments: AB JM. Analyzed the data: AB. Contributed reagents/materials/analysis tools: AB JM. Wrote the paper: AB PC EB.
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