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New Hybrids between *Saccharomyces* Sensu Stricto Yeast Species Found among Wine and Cider Production Strains

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Two yeast isolates, a wine-making yeast first identified as a Mel* strain (ex. *S. uvarum*) and a cider-making yeast, were characterized for their nuclear and mitochondrial genomes. Electrophoretic karyotyping analyses, restriction fragment length polymorphism maps of PCR-amplified *MET2* gene fragments, and the sequence analysis of a part of the two *MET2* gene alleles found support the notion that these two strains constitute hybrids between *Saccharomyces cerevisiae* and *Saccharomyces bayanus*. The two hybrid strains had completely different restriction patterns of mitochondrial DNA as well as different sequences of the *OLL1* gene. The sequence of the *OLL1* gene from the wine hybrid strain appeared to be the same as that of the *S. cerevisiae* gene, whereas the *OLL1* gene of the cider hybrid strain is equally divergent from both putative parents, *S. bayanus* and *S. cerevisiae*. Some fermentative properties were also examined, and one phenotype was found to reflect the hybrid nature of these two strains. The origin and nature of such hybridization events are discussed.

The genus *Saccharomyces* can be divided into two major groups: sensu stricto and sensu lato (2). The sensu stricto yeasts, which include *S. bayanus*, *S. cerevisiae*, *S. paradoxus*, and *S. pastorianus* (syn. *S. carlsbergensis*), represent a closely related biological complex (14). *S. cerevisiae* is the major species found among wine yeasts, while *S. bayanus* represents a small part of them. The sensu stricto yeasts contain at least 16 distinctive nuclear chromosomes of small, medium, and large sizes, and each species appears to contain a unique karyotype (27). Their mitochondrial DNA (mtDNA) molecules range in size from 64 to 85 kb and contain a number of G+C clusters, among them three to nine ori-rep sequences (27). Molecular polymorphism is widespread among the sensu stricto yeasts, especially among yeast strains associated with the wine industry (5, 30), and almost every isolate has a characteristic karyotype and restriction pattern of digested mtDNA (27). However, among isolates belonging to the same species, similar karyotypes and restriction patterns are observed. In the laboratory, members of the sensu stricto group can be mated at low frequency and can generate viable offspring (19).

The lager brewing strain *S. pastorianus* (syn. *S. carlsbergensis*) is a partial amphitrichoplax, which was generated upon an interspecific fusion-cross between two different yeasts (see, e.g., reference 11). One of the parental strains in this fusion-cross was *S. cerevisiae* and the second was a member of the *S. bayanus* species complex, possibly *S. monacensis* (8, 24, 27). In the characterized strains of *S. pastorianus* (syn. *S. carlsbergensis*), both sets of parental chromosomes are present (11), but the mtDNA molecule was inherited only from the non-*S. cerevisiae* parent (27). Initially, the hybrid zygot was possibly heteroplasmic regarding the mitochondrial genome, but apparently only one parental type was transmitted to the progeny.

In this report, two yeast isolates, a wine-making yeast first identified as Mel* (ex. *S. uvarum*) and a cider-making yeast, are characterized for their nuclear and mitochondrial genome and are shown to be hybrids. In addition, some fermentation properties such as production of aroma compounds of these two yeasts are studied.

**MATERIALS AND METHODS**

**Yeast strains and media.** The yeast strains used in this study are listed in Table 1. *S. cerevisiae* VKM Y-502 and *S. bayanus* VKM Y-1146 are monoporic cultures of reference strains (20). Except for strain S288C, which is a standard laboratory strain, all other strains of *S. cerevisiae* are industrial wine-making yeasts. *S. bayanus* CBS 380 and *S. paradoxus* CBS 432 are the type strains, and *S. carlsbergensis* Y385 was used for the mtDNA restriction fragment length polymorphism (RFLP) experiment is a beer production strain. Other strains of *S. bayanus* are wine yeasts from the collection of the Faculté d’Oenologie de Bordeaux. CID1 is a cider yeast which was isolated from a mixed culture from the bottom of a home-fabricated apple cider from Brittany, France. SEU is a wine-producing yeast. Yeast were grown on YPG medium (20 g of fructose/liter, 10 g of Bacto-Peptone/liter, 10 g of yeast extract/liter) at 20, 25, or 30°C, depending on the species. Fermentations were carried out in grape juice of *Vitis vinifera* var. Sauvignon. The must was sterilized by filtration (turbidity, <0.5 NTU).

**Contour-clamped homogeneous electric field gel electrophoresis.** Chromosomal DNA was prepared in agarose plugs (3) and separated on a 0.8% agarose gel (Agarose NA; Pharmacia) at 165 V and 10°C by using the following program (6): switch, 12.5 h, 40 to 90 s; switch, 16.5 h, 80 to 120 s.

**PCR-RFLP.** The PCR amplification reaction was carried out on entire yeast cells after cultivation on solid YPG medium until the stationary phase (17). Amplification reactions were performed with a Perkin-Elmer DNA thermal Cycler 480, using synthetic oligonucleotidc primers for *MET2* amplification as described by Hansen and Kielland-Brandt (8). PCR products were precipitated, and aliquots were digested with *EcoRI* or *PstI*. The resulting DNA fragments were analyzed by electrophoresis on a 1.8% agarose gel (Agarose NA). A Boehringer Mannheim DNA molecular weight marker VIII was used.

**Preparation and sequencing of *MET2* gene fragments.** For preparative purposes, *MET2* fragments from strains S6U and CID1 were amplified by PCR by using the primers 5′-CGGCTCCTAGACGAAAACGCTTACAGATTG-3′ and 5′-CGGCTCCTAGAACCAGACCGTCG-3′, containing at their ends XbaI restriction sites and four arbitrary bases to allow for restriction endonuclease digestion. Genomic DNA was prepared from 10-m1 liquid yeast cultures by the protocol of Hoffman and Winston (9). Ten microliters of a 100× dilution of each DNA preparation was used as template. The PCRs were performed on a Stratagene Robocycler 40 for 25 cycles of 1 min at 94°C, 2 min at 50°C, and 3 min at 72°C, followed by one cycle of 72°C for 10 min. Eight independent reactions for each DNA template were performed. Each series of identical reactions was pooled, and the amplified DNA was precipitated, washed, and resuspended in an appropriate volume of water and used for direct sequencing or cloning. Restriction digestions and ligation reactions were performed in

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TABLE 1. Yeast strains used in this study

<table>
<thead>
<tr>
<th>Strain</th>
<th>Origin or source</th>
<th>Species</th>
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<tbody>
<tr>
<td>VKM Y-502</td>
<td>VKM</td>
<td>S. cerevisiae</td>
</tr>
<tr>
<td>CBS 1171</td>
<td>CBS</td>
<td>S. cerevisiae (type strain)</td>
</tr>
<tr>
<td>CBS 380</td>
<td>CBS</td>
<td>S. bayanus (type strain)</td>
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<tr>
<td>VKM Y-1146</td>
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<tr>
<td>TB28</td>
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<td>S. paradoxa (type strain)</td>
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<td>S. carlsbergensis</td>
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<tr>
<td>Y385</td>
<td>Collection of J. Piskur</td>
<td>S. carlsbergensis</td>
</tr>
</tbody>
</table>

* CBS, Centraal Bureau voor Schimmelcultures, Baarn, The Netherlands; VKM, National Collection of Microorganisms, Moscow, Russia; CLIB, Collection de Levures d’Intérêt Biotechnologique, Paris, France.

***TABLE 1. Yeast strains used in this study***

accreditation with the recommendations of the manufacturers. DNA fragments were isolated from agarose by using Bio-Rad Prep-A-Gene purification matrix. The sequencing reactions were performed on a Perkin-Elmer DNA Thermocycler 480, and sequences were run on an Applied Biosystems Sequenator 373A or 310 in accordance with the recommendations of the manufacturers. Sequencing primers for direct sequencing were identical to the ones used for the PCR amplification except that no restriction sites or additional arbitrary bases were included. Sequencing of the cloned fragments was performed employing the same primers or standard m13 primers. Both strands of the DNA were sequenced in all cases.

**Cloning of MET2 DNA fragments.** PCR-amplified MET2 fragments from both strains were cloned into pUC19 as follows. Precipitated, redissolved DNA was restriction endonuclease digested with either PstI and XbaI or with EcoRI and XbaI. Uncut DNA fragments were purified as described above and ligated into pUC19 vector that was opened with XbaI and treated with calf intestine alkaline phosphatase. PCR-amplified DNA from several independent reactions were employed in these experiments. The resulting plasmids used for sequencing were either HplI digest (strain S6U; EcoRI uncut), pJH150 (strain CID1; EcoRI uncut), pJH153 (strain S6U; PstI uncut) and pJH156 and pJH157 (strain CID1; PstI uncut).

**Isolation and sequencing of mtDNA.** mtDNA was isolated from various yeasts by using a modification of the bisbenzimide-CsCl gradient method (25, 27). RFLP was studied on purified mtDNA by using GC-clutters as enzymes, i.e., *Hai*III and *Msp*I. The sequence of the mitochondrial *OLI1* gene was obtained by direct sequencing on purified mtDNA (7). The following two primers, with homology to the 5’ and 3’ ends of the *OLI1* coding region, were used in direct sequencing: *OLI1* YM1 (forward primer), 5’-GCAATTAAATTAGTATTAGCAGCT AATATATTGG-3’; and *OLI1* YM4 (reverse primer), 5’-AATAGAATATGAAACCATATAACACAG-3’. The open reading frame of *OLI1* is 228 bp long, and the sequence was determined on both strands except for the terminal 25 bases in the 5’ and 3’ ends, which were determined on only one strand.

**Fermentation experiments.** The yeast inocula were obtained from overnight cultures grown on diluted malt. The quantity of yeasts was measured by determining the optical density at 600 nm in order to inoculate the must at a level of 3 × 10⁹ cells/ml. The fermentation test was carried out in 375-ml bottles containing grape juice. The turbidity of the juice was adjusted to 200 NTU with insoluble material of the must to improve the fermentation velocity (22). At the midpoint of the fermentation, the control experiments were performed to ensure that the must had been inoculated with the correct yeast species and strains. The implantation of *S. cerevisiae* strains was verified by PCR amplification of delta sequences (17). For the *S. bayanus* strains, karyotyping by contour-clamped homogeneous electric field gel electrophoresis was used to confirm the inoculation when the sugars were below 2 glitler, bottles were placed at 10°C and SO₂ was adjusted to 60 mg/liter and the wines were rapidly analyzed for content of higher alcohols and esters by gas chromatography coupled with a flame ionization detector (CARBOWAX 20M capillary column, type BP20; length, 50 m; internal diameter, 0.25 mm; film thickness, 0.50 μm; VARIAN 3400 gas chromatograph; Merck D-2500 chromatointegrator).

**RESULTS**

**Electrophoretic karyotyping analyses.** Wine *S. cerevisiae* strains are characterized by important variations in chromosomal length whereas wine yeasts, genetically identified as *S. bayanus*, do not exhibit large chromosomal polymorphism (20, 30). However, many authors in previous works have shown that electrophoretic karyotyping analyses can be used to differentiate *S. cerevisiae* and *S. bayanus* (4, 13, 20, 24). Chromosomal DNA patterns of strains S6U and CID1 displayed similar but specific band patterns of more than 20 bands. When these patterns were compared to the reference strains of *S. cerevisiae* and *S. bayanus*, a high proportion of bands corresponded to one or the other reference strain (Fig. 1). Actually, the karyotypes of S6U and CID1 contained an almost complete set of *S. cerevisiae* and *S. bayanus* chromosomes, indicating the hybrid nature of these two yeasts. According to our previous studies of various yeast isolates which were either *S. cerevisiae*- or *S. bayanus*-like, the existence of hybrids in nature is quite rare (16, 18). Karyotypes of different isolates belonging to the same species exhibit polymorphism, like the isolates belonging to the presented *S. bayanus* and *S. cerevisiae* strains. On the other hand, the two hybrid isolates displayed a similar karyotype. Thus, it is likely that the hybrids have a similar origin for the nuclear genome.

**PCR-RFLP on the MET2 gene.** To substantiate the hypothesis that S6U and CID1 are hybrid yeasts containing genomic material related to that of *S. cerevisiae* and *S. bayanus*, we decided to employ RFLP on a 580-bp PCR-amplified fragment of a nuclear gene, MET2, as described previously (8, 18). The restriction endonuclease *Pst*I cuts this MET2 sequence of *S. bayanus*, and there is no *Pst*I site in the MET2 sequence of *S. cerevisiae*. On the contrary, *EcoRI* cuts the *S. cerevisiae* MET2 sequence but not the *S. bayanus* sequence (8, 18). The results obtained are shown in Fig. 2. *S. cerevisiae* VKM Y-502 and *S. bayanus* VKM Y-1146 and CBS 380 were used as reference strains. For the enzymes *EcoRI* and *Pst*I, the restriction fragment profiles are characteristic of the two species *S. cerevisiae* and *S. bayanus*; *EcoRI* cleaves the *S. cerevisiae* MET2 fragment (two bands of 211 bp and 369 bp) but does not cleave the *S. bayanus* MET2 fragment. The behavior of *Pst*I is different: for S6U and CID1, an *EcoRI* and a *Pst*I restriction frag-
A restriction fragment profile pattern of three bands was obtained, with the same intensity and the same length as those obtained for *S. cerevisiae* with EcoRI and for *S. bayanus* with PstI (Fig. 2). These restriction fragment profiles appeared to consist of a mix of the profiles seen from *S. cerevisiae* and *S. bayanus* with a given enzyme and were identified for different subclones of S6U and CID1 from vegetative cells, which were isolated with a micromanipulator.

Divergent MET2 sequences of the putative hybrid yeasts. To obtain the nucleotide sequence of the central 330 bp of the amplified MET2 DNA fragments, we employed two tech-

![Fig. 2. RFLP analysis of PCR-amplified MET2 gene fragment. Lanes 1 to 6, restriction analysis with EcoRI. Lanes 7 to 12, restriction analysis with PstI. Lanes 1 and 6, S6U; lanes 2 and 8, CID1; lanes 3 and 9, *S. cerevisiae* VKM Y-502; lanes 4 and 10, *S. cerevisiae* type strain 1171; lanes 5 and 11, *S. bayanus* VKM Y-1146; lanes 6 and 12, *S. bayanus* CBS 380. M, molecular weight marker (marker VIII from Boehringer Mannheim). Numbers and arrows are base pair markers.](image-url)

![Fig. 3. Partial nucleotide sequences of the MET2 genes from *S. bayanus* CBS 380 type strain (Bay T) (8), *S. cerevisiae* S288C (Cer-S288C) (15), the *S. bayanus*-like allele from S6U (S6U-1) and CID1 (CID1-1), and the *S. cerevisiae*-like allele from S6U (S6U-2) and CID1 (CID1-2). A dot denotes an identical nucleotide.](image-url)
techniques: direct sequencing of the PCR products, as described by Hansen and Kielland-Brandt (8) and sequencing of cloned PCR fragments. In the case of direct sequencing, the PCR fragments were thoroughly digested with either PstI or EcoRI. The DNA fragments in the restriction digests were separated on 2% agarose, and remaining uncut fragments were purified. In this manner, we were able to obtain unambiguous sequences from both fragments from strain S6U present after restriction digestion. Likewise, we obtained good sequences from the EcoRI-uncut MET2 fragment from strain CID1 and reasonably good sequences from the PstI-uncut MET2 fragment from the same strain. However, to resolve the identity of a few ambiguous nucleotides in the PstI-uncut MET2 from CID1, and in general to confirm the sequencing results, we decided to redo the sequencing on cloned MET2 PCR fragments. The cloning is described in the Materials and Methods section. The inserts of the plasmids pJH147, pJH150, pJH153, pJH156, and pJH157 were sequenced. The results from the direct sequencing of the S6U MET2 fragments were confirmed, and the sequences obtained from plasmids pJH156 and pJH157 were identical. As can be seen in Fig. 3, the 330 bp of the PstI-uncut MET2 alleles of both strains CID1 and S6U were completely identical to each other and to those of S. cerevisiae MET2 (15). The EcoRI-uncut MET2 alleles from both organisms were also identical, being 82% homologous to S. cerevisiae and 98.5% homologous to the MET2 allele of the S. bayanus type strain (8). We conclude that both strains are hybrid yeasts and that their genetic content may be regarded as derived, at least partially, from the genomes of S. cerevisiae and S. bayanus.

RFLP of mtDNA. When a cross between two yeast cells occurs, the zygote contains the nuclear and mitochondrial genetic material from both parents. However, while the nuclear chromosomes are transmitted almost equally to the daughter cells, the mtDNA molecules segregate and even exhibit a bias in transmission (26). Therefore, the progeny initially contains a mixture of daughter cells which have the mitochondrial genome from one or another parent, or a novel recombinant mtDNA molecule. As mentioned before, in the case of S. carlsbergensis, only the non-S. cerevisiae mtDNA molecule was inherited. When mtDNAs from the two putative hybrid yeasts, S6U and CID1, were digested, they provided two completely different restriction patterns (Fig. 4), composed of more than 20 distinctive bands. The two patterns were also different from the pattern characteristic for mtDNA from S. paradoxus, S. carlsbergensis (Fig. 4), S. cerevisiae, and S. bayanus (27). Therefore, these restriction patterns can be used as fingerprints for identification of these yeasts. In addition, these data suggest that the mtDNA molecules from these two yeasts may not be very closely related to each other and could have a different origin. This possibility was more closely examined by sequencing of the mitochondrial OLI1 gene.

Sequence analysis of the mitochondrial OLI1 gene. The OLI1 gene is one of the shortest and most conserved mitochondrial genes. In S. cerevisiae and S. paradoxus-S. douglasi, the open reading frame consists of 228 bp, which corresponds to 76 amino acids, and only three “silent” substitutions were found between these two species (21, 23). The open reading frames of the OLI1 gene originating from the two hybrid species, S6U and CID1, as well as from S. bayanus, were also found to be 228 bp long (Fig. 5). The amino acid sequence was identical in all cases, but several silent substitutions were observed among the analyzed species (Fig. 5). It is likely that these differences represent neutral mutations and can be directly used in reconstruction of the origin of the two hybrid yeasts. Nucleotide divergence within OLI1 among the tested species is shown in Table 2. Apparently, S. cerevisiae and S. douglasiobd

S. paradoxus are more closely related to each other than to S. bayanus. This observation fits well with the previous analysis of the mtDNA molecules from these three species (7, 27). The hybrid strains, S6U and CID1, show a divergence of 2.2%, which is almost as high as that between S. cerevisiae and S. bayanus, 2.6%. While the sequence of the OLI1 gene from S6U appears to be the same as for the S. cerevisiae gene, the CID1 OLI1 gene is equally divergent, 2.2%, from both putative parents, S. bayanus and S. cerevisiae (Table 2). These results unambiguously show that the mtDNA molecules of the two hybrid strains are different and are likely to have a different origin.

Phenotypic divergence: production of esters. Some strains of S. bayanus have been reported to have specific and somewhat unusual fermentation properties. Some are cryophilic, having a higher growth rate and a better fermentability at low temperatures as compared to S. cerevisiae strains (13), and these produce wines with higher than usual amounts of flavor-active esters, especially β-phenylethyl alcohol and β-phenylethyl acetate (12, 28). Moreover S. cerevisiae-S. bayanus hybrids obtained by hybridization in the laboratory exhibited such fer-
mentation characteristics at intermediate values (12, 28). We set out to investigate whether the hybrid nature of the genomes of S6U and CID1 was in any way phenotypically reflected in the production of these aroma compounds. The same Sauvignon blanc grape must was inoculated with four industrial S. cervisiae wine yeast strains (EG8, VL3c, VL1, and SIHA3), two indigenous S. bayanus wine yeast strains (P3 and TB28), and the strains S6U and CID1. At the midpoint of the alcoholic fermentation, the strain implantation was verified by PCR as-sociated with the delta sequence (S. cerevisiae strains and the strains S6U and CID1) (16) (data not shown) or electro-phoretic karyotyping (S. bayanus strains). The amounts of /H9252-phenylethyl alcohol and its acetate ester obtained for each strain are reported in Table 3. According to previous reports, the values obtained for the S. bayanus strains were 7.5 to 10 times higher for β-phenylethyl alcohol and 3 to 13 times higher for β-phenylethyl acetate compared to S. cerevisiae wine yeast (12). The wines produced by S6U and CID1 contain intermediate amounts of the two compounds, thus indicating that the genetic hybrid nature of these yeasts seems to be somewhat reflected in at least one phenotype of importance to the wine industry.

DISCUSSION

When the karyotypes of the two wine and cider yeasts, S6U and CID1, were compared to the karyotypes of some known yeast species, it was apparent that their nuclear genomes contain S. cerevisiae-like and S. bayanus-like chromosomes. Therefore, it seemed likely that the two yeast strains are hybrids.

| TABLE 3. Production of β-phenylethyl alcohol and β-phenylethyl acetate by different strains of S. cerevisiae and S. bayanus and by hybrid strains |
|---|---|---|---|
| Species or hybrid strain | Yeast strain | β-Phenylethyl alcohol (mg/liter) | β-Phenylethyl acetate (mg/liter) |
| S. cerevisiae | EG8 | 15.6 | 0.63 |
| S. cerevisiae | VL3c | 22.8 | 0.68 |
| S. cerevisiae | VL1 | 16.5 | 0.45 |
| S. cerevisiae | SIHA3 | 18.6 | 0.91 |
| S. bayanus | P3 | 256 | 12.3 |
| S. bayanus | TB28 | 118 | 1.31 |
| Hybrid | S6U | 47.6 | 2.17 |
| Hybrid | CID1 | 48.3 | 1.61 |

* The number of changes between each pair of species is shown, and the percentage of divergence is shown in parentheses. The data for S. paradoxus and S. cerevisiae are from Ooi et al. (23) and Nicoletti et al. (21). The S. bayanus strain was CBS 380.
between two species, *S. cerevisiae* and *S. bayanus*. This theory was substantiated by the analysis of the nuclear MET2 gene, the sequence of which differs characteristically between *S. cerevisiae* and *S. bayanus* (8, 18). RFLP maps of a PCR-amplified MET2 gene fragment appeared as mixes of the RFLP maps of *S. cerevisiae* and *S. bayanus*, thus supporting the notion that S6U and CID1 constitute hybrids between *S. cerevisiae* and *S. bayanus*. Verification of this theory was obtained by the sequence analysis of a part of the two MET2 gene alleles, supposedly present in both yeasts: in S6U and CID1, there are indeed two alleles of the gene, one identical to *S. cerevisiae* MET2, and one almost identical to *S. bayanus* MET2. It is furthermore interesting that both copies of this gene were almost identical in both hybrid strains. Therefore, it is likely that the nuclear genomes of both hybrids have a similar origin. The origin, a cross between *S. cerevisiae* and a *S. bayanus*-like yeast, is reminiscent of the situation found in *S. carlsbergensis* lager brewing yeast. However, while *S. carlsbergensis* inherited the non-*S. cerevisiae*-like mtDNA molecule (27), the mitochondrial inheritance pattern is different in S6U and CID1.

While in yeast crosses, nuclear DNA is inherited from both parents, mtDNA exhibits a non-Mendelian pattern of inheritance (26). In the progeny, only one or the other parental mtDNA molecule, or a recombinant one, is found. The S6U and CID1 hybrids contained two different mtDNA molecules. The mtDNA molecule in S6U appears to originate from the *S. cerevisiae*-like parent, whereas the CID1 mtDNA molecule differs from that of *S. cerevisiae* as well as that from *S. bayanus*. Phylogenetically, the latter mtDNA molecule could be positioned between the *S. cerevisiae* and *S. bayanus* mtDNA molecules. Therefore, it is likely that the two hybrid strains do not originate from a single hybridization event. While the nuclear backgrounds of the parents involved in both crosses were probably very similar, the mitochondrial backgrounds were likely to be different.

It appears that among Saccharomyces yeasts used in fabrication of wine, cider, and beer, stable interspecies hybrids are quite common. Whether such hybrids originate from events having taken place in the production environments or in nature is not known. As the genetic constitution of these yeasts seems having taken place in the production environments or in nature quite common. Whether such hybrids originate from events efficiently releases autonomous plasmids for transformation of *Escherichia coli*. Gene 57:267–270.


