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Molecular Characterization of a Chromosomal Rearrangement Involved in the Adaptive Evolution of Yeast Strains

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Wine yeast strains show a high level of chromosome length polymorphism. This polymorphism is mainly generated by illegitimate recombination mediated by Ty transposons or subtelomeric repeated sequences. We have found, however, that the *SSUI-R* allele, which confers sulfite resistance to yeast cells, is the product of a reciprocal translocation between chromosomes VIII and XVI due to unequal crossing-over mediated by microhomology between very short sequences on the 5' upstream regions of the *SSUI* and *ECM34* genes. We also show that this translocation is only present in wine yeast strains, suggesting that the use for millennia of sulfite as a preservative in wine production could have favored its selection. This is the first time that a gross chromosomal rearrangement is shown to be involved in the adaptive evolution of *Saccharomyces cerevisiae*.

[The sequence data from this study have been submitted to EMBL under accession nos. AF239757, AF239758, and AJ458340–AJ458367. The following individual kindly provided reagents, samples, or unpublished information as indicated in the paper: N. Goto-Yamamoto.]

The unaware use of yeast for winemaking by the first agricultural civilizations has been reported as far back as 7400 years ago. Until the middle of the last millennium, wines were mainly produced around the Mediterranean Sea and the Caucasus. Since then, winemaking has spread with the European colonizers throughout the temperate regions of the world (Pretorius 2000).

Although different genera and species of yeasts are found in musts, the species *Saccharomyces cerevisiae* is mainly responsible for the transformation of musts into wines. The origin of *S. cerevisiae* is controversial. Some authors propose that this species is a "natural" organism present in plant fruits (Mortimer and Polsinelli 1999). Others argue that *S. cerevisiae* is a domesticated species originated from its closest relative *S. paradoxus*, a wild species found all around the world (Vaughan-Martini and Martini 1995). This debate is important in postulating the original genome of *S. cerevisiae* and how the strong selective pressure applied since its first unconscious use in controlled fermentation processes has reshaped it. Useful phenotypic traits such as fast growth in sugar-rich media, high alcohol production and tolerance, and good flavor production selected for billions of generations have had strong influences on the *S. cerevisiae* genome.

In contrast to most *S. cerevisiae* strains used in the laboratory, which are either haploid or diploid and have a constant chromosome electrophoretic profile, wine yeast strains are mainly diploid, aneuploid, or polyploid, homothallic, and

highly heterozygous (Bakalinsky and Snow 1990; Barre et al. 1993; Codón et al. 1995), and show a high level of chromosome length polymorphisms (Bidenne et al. 1992; Rachidi et al. 1999). Moreover, wine yeast strains seem not to remain genetically uniform (Pretorius 2000). Their exacerbated capacity to reorganize its genome by chromosome rearrangements such as Ty-promoted chromosomal translocations (Longo and Vézinhel 1993; Rachidi et al. 1999), mitotic crossing-over (Aguilera et al. 2000), and gene conversion (Puig et al. 2000) promotes a faster adaptation to environmental changes than spontaneous mutations, which occur at comparatively very low rates. The ploidy of the wine yeasts may confer advantages in adapting to variable external environments or increasing the dosage of some genes important for fermentation (Bakalinsky and Snow 1990; Salmon 1997).

In addition, the possibility of adaptive gross genomic changes occurring during laboratory growth conditions has been demonstrated with DNA chip technology by Hughes et al. (2000). Those authors showed in multiple cases that the deletion of a gene strongly favors the acquisition of a second copy of a whole chromosome or a chromosomal segment containing a compensatory copy of a close homolog of the deleted gene.

In a comparative study of transcriptomes, we found that *SSUI*, a gene that mediates sulfite efflux in *S. cerevisiae* and, hence, confers sulfite resistance (Park and Bakalinsky 2000), showed a significantly higher expression in the T73 wine yeast strain than in a laboratory strain (Hauser et al. 2001). In contrast to the allele present in the laboratory strains, a highly sulfite-resistant wine strain exhibited a translocation involving the promoter region of the gene (*SSUI-R* allele), which produces an increase in the sulfite resistance (Goto-Yamamoto et al. 1998). In the present study, we explored the organization of this gene at the molecular level in different wine yeast strains.

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RESULTS

Genome Organization of the *SSU1* Locus in T73 Wine Yeast Strain

Sulfite-generating compounds are widely used during wine making as bacterial inhibitors (Pretorius 2000). Therefore, sulfite resistance is a desired trait for wine yeast strains. Interestingly, our comprehensive study of gene expression in the T73 wine yeast strain grown under standard laboratory conditions (Hauser et al. 2001) revealed that the *SSU1* gene, which mediates sulfite efflux in *S. cerevisiae* (Park and Bakalinsky 2000), is expressed more in the wine strain than in a reference laboratory strain (S288c background). High levels of *SSU1* mRNA have also been observed in other wine yeast strains, such as Y-9, and this has been correlated with increased sulfite resistance (Goto-Yamamoto et al. 1998). Y-9 strain possesses an *SSU1* allele (*SSU1-R*) with an upstream sequence completely replaced due to a putative translocation (Goto-Yamamoto et al. 1998).

To understand the mechanisms underlying the increased expression of *SSU1* in wine yeast strains, we decided to investigate the promoter sequence of this gene. For this purpose, PCR primers for the selective amplification of the *SSU1* region were designed according to the sequences of the laboratory S288c background (*SSU1* allele) and Y-9 (*SSU1-R* allele) wine yeast strain (Fig. 1). Both primer pairs (*SSU1MD/SSU1R* for the *SSU1* allele, and *ECM34D/SSU1R* for the *SSU1-R* allele) amplified in T73 fragments of the predicted lengths, 569 and 573 bp, respectively (data not shown). Their sequences were in both cases 100% identical to those described for S288c background (from MIPS database) and Y-9 strain (Goto-Yamamoto et al. 1998). Consequently, we can conclude that the T73 wine yeast strain is a heterozygote containing both *SSU1* and *SSU1-R* alleles.

The promoter sequence of the *SSU1-R* allele presents a very high similarity with the promoter sequence of *ECM34*, a gene of unknown function from chromosome VIII (this study: accession no. AF239758, and Goto-Yamamoto et al. [1998]: accession no. AB002531). It is worth noting that the sequence of the *SSU1-R* allele contains four repeats of a 76-bp sequence, which is a single copy of 77 bp in *ECM34* from S288c background (Fig. 2). These results strongly suggest that a reciprocal

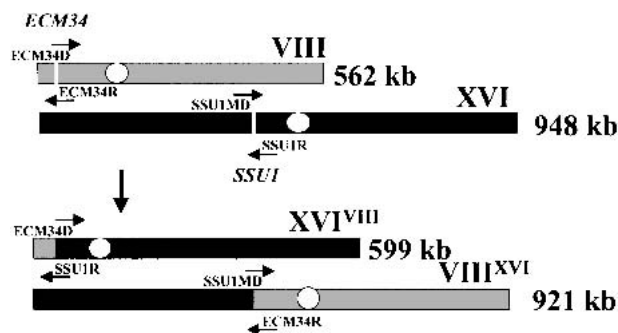


Figure 1 Diagram representing the reciprocal translocation between chromosomes VIII and XVI observed in wine yeast strains. This translocation was mediated by crossing-over between microhomology regions of the promoters of the *ECM34* and *SSU1* genes, the locations of which, on chromosomes VIII and XVI, respectively, are indicated by white bars. Small arrows indicate the PCR primers used to amplify those regions involved in the recombination.

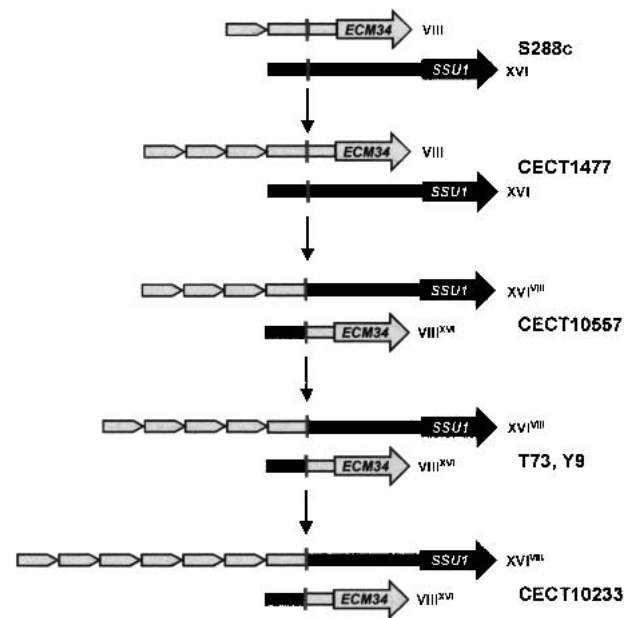


Figure 2 Diagrams representing the organization of the *ECM34* (in gray) and *SSU1* (in black) nonrecombinant alleles and their corresponding recombinant variants obtained by an illegitimate crossing-over of a microhomology region (see Fig. 3), indicated by a vertical line, located in the promoters of both genes. This illegitimate crossing-over was involved in the generation of the reciprocal translocation between chromosomes VIII and XVI found in *Saccharomyces* strains (Fig. 1). Thick arrows represent the protein-coding regions of each gene. The pentagon block corresponds to a 76-bp sequence that has been found several times repeated in the promoters of both nonrecombinant *ECM34* (strain CECT 1477) and recombinant *SSU1-R* (several strains). Strains bearing these sequences are indicated for each diagram.

translocation between chromosomes VIII and XVI could have occurred in the wine yeasts (Fig. 1).

By designing a new primer from the *ECM34* coding sequence (*ECM34R*), two new PCR amplifications were performed with the *ECM34D+ECM34R* and *SSU1MD+ECM34R* primer pairs (Fig. 1). The first primer pair allowed us to amplify from T73 a band of 207 bp corresponding to the “standard” *ECM34* locus, and the second pair, a band of ~450 bp (accession number AF239757), which corresponded with a putative reciprocal translocation between chromosomes VIII and XVI at the 5' upstream regions of the *SSU1* and *ECM34* genes (30 bp upstream from the *ECM34* ATG start codon) (Fig. 2).

This putative translocation explains the contoured homogeneous electrical field electrophoresis (CHEF) analysis we previously observed for the T73 strain (Puig et al. 2000). That analysis revealed the existence of anomalously sized bands in the region of chromosomes VIII and XVI (Figs. 1 and 2 from Puig et al. 2000). A probe from chromosome VIII (*CUP1* sequence) hybridized in a Southern blot with two chromosomal bands of ~560 (chromosome VIII) and ~920 Kb (chromosome VIII^{XVI}), and a probe from chromosome XVI (*CARI* sequence) hybridized with bands of ~920 (chromosome VIII^{XVI}) and ~950 Kb (chromosome XVI). This strain had a sporulation efficiency of 60% and a spore viability of 70%. However, only 13% of the dissected tetrads produced four viable spores (Puig et al. 2000). In all complete viable

tetrads analyzed, the polymorphic bands observed after hybridization with probes from chromosomes VIII and XVI segregated at 2:2 (Puig et al. 2000), indicating that each pair of bands is allelic. These results confirm that a reciprocal translocation involving chromosomes VIII and XVI occurred (Fig. 1), and that the T73 strain is heterozygous for the reciprocal translocation, containing both the translocated and the “standard” chromosomal arrangements. Because translocated chromosomes contain all of the original genes from chromosomes VIII and XVI, spores are viable only if they contain either both “standard” chromosomes or both translocated ones.

The reciprocal translocation between chromosomes VIII and XVI could be originated by different molecular mechanisms. We wondered whether a short region of sequence homology could mediate a heterologous recombination between the promoter sequences of *ECM34* and *SSU1* genes. When comparing the sequences of both gene promoters at the translocation breakpoint, we realized that the best alignment contained a short sequence of 9–13 bp of microhomology (Fig. 3), which includes the recombination site as deduced from the sequences of the recombinant and nonrecombinant alleles (Fig. 2). Taken together, these results suggest that the *SSU1-R* allele present in T73 and Y-9 wine yeast strains was generated by a reciprocal translocation between chromosomes VIII and XVI due to unequal crossing-over between a short region of microhomology located in the promoter region of *SSU1* and *ECM34* genes.

Frequency and Origin of the Translocation (VIII;XVI) in *S. cerevisiae*

The fact that two geographically distant but naturally occurring wine yeast strains, Y-9 and T73, exhibited the same translocation prompted us to investigate whether this rearrangement was present in other *S. cerevisiae* strains isolated from different sources (wine and nonwine), in diverse geographic origins, and during several periods of time. A total of 30 strains (Table 1, 18 isolated from wine and 12 from other sources) were analyzed by PCR amplification with the different combinations of primers. The characteristic 450-bp PCR fragment of the recombinant *SSU1-R* allele (*ECM34*-derived promoter region+*SSU1* coding region) was found in five additional strains, all of them also isolated from wines in different geographical areas. Four of them (CECT 10120, 1485, 10557, and 11827) were homozygous for the translocation as Y-9,

and the other one (CECT 10233) corresponded to a translocation heterozygote as T73 (Table 1). The recombinant *SSU1-R* promoters from four of these additional strains were sequenced (EMBL accession numbers AJ458364–AJ458367), and the results showed they were all identical, except for the number of 76-bp repeats, indicating that the translocation (VIII;XVI) was a rare and unique event.

The 76-bp repeats present in the recombinant *SSU1-R* promoters contain some nucleotide substitutions and insertions/deletions (indels) compared to the original promoter of the *ECM34* allele (see Fig. 2). Therefore, to deduce the origin of the recombinant promoter, we decided to amplify and sequence the nonrecombinant *ECM34* promoter region from the two heterozygous strains T73 and 10233, from 23 additional *S. cerevisiae* strains, and from eight strains from other *Saccharomyces* “sensu stricto” species (Table 1, EMBL accession numbers AJ458340–AJ458363). The *ECM34* promoter region could only be amplified in the *S. cerevisiae* strains and in the two *S. pastorianus* strains. This species is a partial allotetraploid originated from an *S. cerevisiae* × *S. bayanus* hybridization (Vaughan-Martini and Kurtzman, 1985; Casaregola et al. 2001; de Barros Lopes et al. 2002), and the amplified *ECM34* promoter region probably corresponds to the *S. cerevisiae* fraction of the *S. pastorianus* genome. The results showed seven different sequence variants of the nonrecombinant *ECM34* promoter (Fig. 4, variants A to G). The most frequent variants, A and B, and the related variants E to G, are mainly present in wine strains, with the exception of strains 10131 and 10392, which were isolated from the plant *Centaurea alba* and from alpechin (olive residues after oil extraction), respectively, in Spain, a wine-producing country, and 10691, isolated from palm wine in West Africa. In contrast, variants C and D are present in laboratory strains, but also in strains 1462, 11837, and 11838 isolated from ale beer, bili wine and grapes, respectively.

We obtained a maximum parsimonious tree that minimizes the number of changes (nucleotide substitutions, indels, and repeat insertions) required to connect these sequences (Fig. 5). In this tree, variant A occupies a central position from which all the other variants can be derived. Variant A and its closest related variant B, which differs in a single nucleotide substitution, are exhibited by a heterogeneous group of wine-related strains and by the hybrid *S. pastorianus* strains.

Variant C, exhibited by three laboratory and three natural strains, differs from variant A by a single G nucleotide insertion. Variant D differs in a single nucleotide substitution from variant A and is present only in laboratory strains S288c and its derivative W303 (Rogowska-Wrzesinska et al. 2001). S288c, the most popular genetic background in yeast research laboratories, is a derivative of a natural heterothallic diploid strain isolated from rotting figs in California in 1938 (Mortimer and Johnston 1986). It is quite likely that this yeast strain was carried from cellars by insects. Taken together, these observations suggest that wine strains could probably be the ancestors of the domesticated laboratory strains (Mortimer and Johnston 1986).

The other variant sequences (E, F, G, and the recombinant R, S, and T), which include strains containing the promoter of the recombinant *SSU1-R* allele, differ from variant A by the presence of a series of tandem repeats of 1, 47, or 76 bp (see Fig. 4). The nonrecombinant allele of the translocation heterozygote T73 (variant E, T73NR) exhibits an insertion of 10 T that was probably generated by replication slippage

Non-recombinant

ECM34 (S288c) gttgtAAC•tgCAAAAaAaTgtCacCggg
ECM34 (CECT1477) gttgtAAC•tACAAAaaAaTgtCacCggg
SSU1 (S288c) tagacAACacACAAAAtTa•CAgCttt



Recombinant

pSSU1-ECM34 (T73) tagacAACacACAAAaaA•TgtCacCggg
pECM34-SSU1 (T73) gttgtAAC•tACAAAaaAtTa•CAgCttt

Figure 3 Microhomology regions, located in the *ECM34* and *SSU1* promoters, that were involved in the crossing-over generating the reciprocal translocation between chromosomes VIII and XVI. Strains where sequences were obtained from are indicated in parentheses. Black and gray lines highlight *ECM34* and *SSU1* promoter sequences, respectively. Perfect sequence matches are shown in capitals, and dots correspond to gaps required to align the sequences.

Table 1. *Saccharomyces* “sensu stricto” Strains Analyzed, Origins, Sources Whence They Were Isolated, PCR Patterns Obtained by Amplification With Primers Specific of the *SSU1* and *ECM34* Regions, and *ECM34* Promoter Sequence Type Exhibited

Strain designation	Other designations	Source, origin, and year of isolation	PCR pattern	pECM34 type
<i>S. cerevisiae</i> strains				
A364a		Laboratory	207, (568)	C
CEN-PK 113 17A		Laboratory	207, (568)	C
S288c		Laboratory	207, 568	D
SK1		Laboratory	207, (568)	C
W303		Laboratory	207, (568)	D
T73	CECT 1894	Wine, Spain (1987)	220, 450, 568, 573	E and S
Y-9		Wine, Japan	450, 573	S
CECT 1383	CBS 2978	Distiller's yeast	207, (568)	A
CECT 1462	NCYC 963	Ale beer, UK	207, (568)	C
CECT 1475	UCD 519	Sherry wine, Jerez, Spain	207, 568	B
CECT 1476	UCD 522	Montrachet, California, USA	207, 568	A
CECT 1477	UG5	Sparkling wine, Bordeaux, France	380, 568	G
CECT 1485		González Byass wineries, Spain	450, 573	S
CECT 1882		Wine pellicle, Huelva, Spain	207, 568	B
CECT 10009	CBS 4054	Red wine, Spain (1958)	207, (568)	B
CECT 10095	CBS 5835	Wine, Spain (1959)	207, (568)	n.d.
CECT 10120		Fruit of <i>Arbutus unedo</i> , Spain	450, 573	S
CECT 10131		Flower of <i>Centaurea alba</i> , Spain	207, (568)	A
CECT 10233	CBS 2247	Grape must, South Africa (1955)	207, 450, (568), 750	D and T
CECT 10392	CBS 3081	Alpechin, Spain (1958)	220, (568)	E
CECT 10557	CBS 5112	Grape must, Spain (1962)	450, 497	R
CECT 10691	CBS 400	Palm wine, from Ivory Coast (1927)	207, 568	A
CECT 10692	CBS 429	Fermenting Champagne grapes (1899)	207, 280, (568)	B
CECT 11032	CBS 459	Grape must, Italy (1938)	220, (568)	F
CECT 11827		Dry wine yeast, Switzerland (1984)	450, (573)	n.d.
CECT 11833	CBS 423	Wine, Switzerland (1924)	207, (568)	C
CECT 11834	CBS 4070	Red wine, Spain (1958)	207, (568)	D
CECT 11835	CBS 1250	Sherry, Spain (1936)	207, (568)	D
CECT 11837	CBS 405	Bili wine, West Africa (1925)	207, (568)	B
CECT 11838	CBS 5287	Grape, Russia (1961)	207, (568)	A
Other <i>Saccharomyces</i> “sensu stricto” species				
CECT 1969	CBS 395	<i>S. bayanus</i> (type of <i>S. uvarum</i>)	No amplification	–
CECT 12635		<i>S. bayanus</i> isolated from wine	No amplification	–
CECT 12636		<i>S. bayanus</i> isolated from wine	No amplification	–
CECT 1939 ^{NT}	CBS 432	<i>S. paradoxus</i> neotype strain	568	–
CECT 11152	IFO 1804	<i>S. paradoxus</i> from tree exhudate, in Japan	568	–
CECT 11158	CBS 2980	<i>S. paradoxus</i> from <i>Drosophila</i> , in California	568	–
CECT 1970	CBS 1503	<i>S. pastorianus</i> (type of <i>S. monacensis</i>)	207	A
CECT 11037	CBS 1513	<i>S. pastorianus</i> (type of <i>S. carlsbergensis</i>)	207	A

Between parentheses are indicated those expected fragments that did not amplify. 450 is the diagnostic band for the translocation, n.d., not determined. CBS (Centraalbureau voor Schimmelcultures, Utrecht, The Netherlands), CECT (Spanish Type Culture Collection, University of Valencia, Spain), IFO (Institute for Fermentation, Osaka, Japan), UCD (Hermann Phaff Collection, University of California, Davis, USA), NCYC (National Collection of Yeast Cultures, Norwich, UK).

within a region of 3 T. The same insertion is also present in strain 10392. Strain 11032 (variant F) contains two tandem repeats of a 47-bp region that could have been generated either by unequal crossing-over between two almost identical 7-bp regions flanking the repeated region (Fig. 4), or by replication slippage also favored by the pairing of the 7-bp flanking regions and the possible secondary structure of the repeat. A different repeated region, although overlapping with that from strain 11032, is present in the promoter region of the recombinant *SSU1-R* alleles (variants R, S, and T). These promoter variants contain three, four, or six tandem repeats of a 76-bp region, respectively. The first duplication event, which occurred either by unequal crossing-over or by replication slippage, could be favored by the presence of a 6-bp identical sequence flanking the repeated region (Fig. 4), and also by a potential secondary structure of three hairpin-loops.

The nonrecombinant *ECM34* promoter from strain 1477 (variant G) also contains three tandem repeats of the same

76-bp region and a G to A substitution located at the putative crossing-over site shared with the recombinant promoter variants of the *SSU1-R* allele, R, S, and T (Figs. 2–4). This sequence organization strongly suggests that the first duplication of the 76-bp region occurred at the *ECM34* promoter before the illegitimate crossing-over between *ECM34* and *SSU1* promoters produced the translocation (*VIII;XVI*). Once the rare first duplication event took place, subsequent duplications could, with a higher probability, be extended by unequal crossing-over (either meiotic or mitotic) between any repeats, or by slipped-strand mispairing between contiguous repeats. It is likely that one subsequent duplication also occurred before the translocation.

The translocation between chromosomes VIII and XVI by the illegitimate crossing-over between *ECM34* and *SSU1* promoters took probably place in a strain with three 76-bp repeats in its *ECM34* promoter, similar to 1477, giving rise to a recombinant promoter with three repeats as that found in

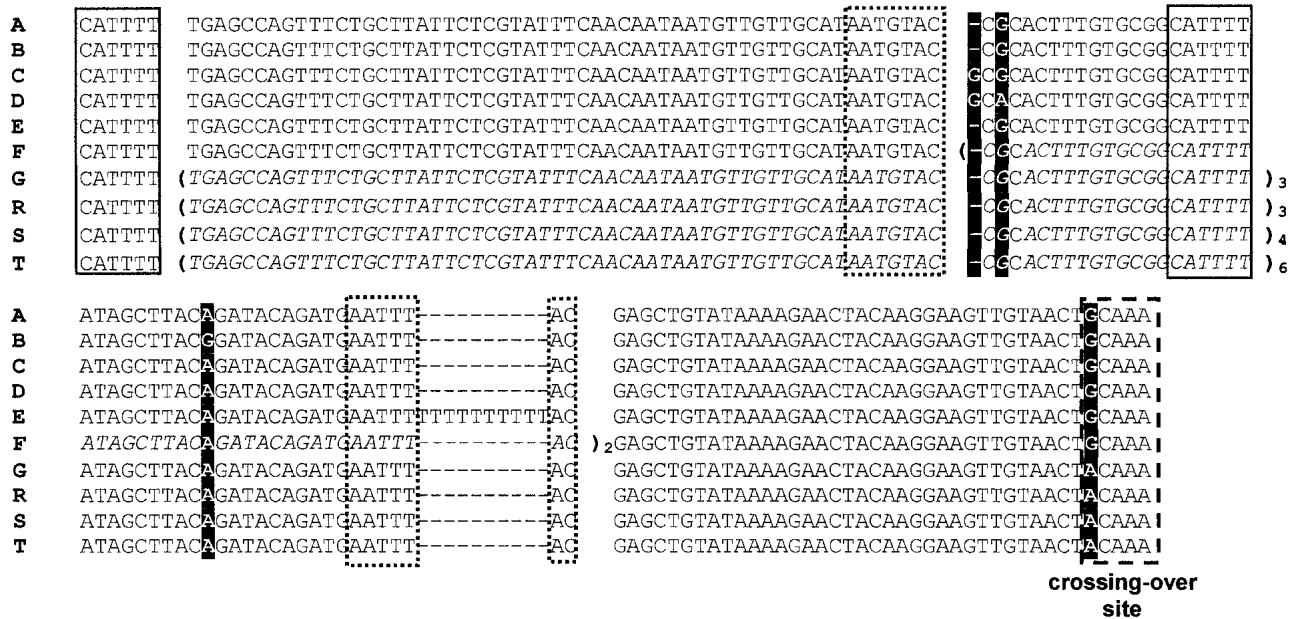


Figure 4 Sequence alignment of the seven nonrecombinant and three recombinant variants of the *ECM34* promoter region found in *Saccharomyces* strains (see Table 1). Variable positions are shown in negative. Italic sequences in parentheses correspond to repeated regions, and subscript numbers after the parentheses indicate the number of repeats. Continuous rectangles highlight a small direct repeated sequence flanking a large 76-bp sequence repeat that could be involved in its duplication. Dotted rectangles indicate a small imperfect repeat that could be involved in the generation of a 47-bp duplication found in the CECT11032 strain. The discontinuous rectangle indicates the crossing-over site involved in the reciprocal translocation t(VIII,XVI).

strain 10577 (Fig. 2). However, the presence of heterozygotes for the translocation (strains 10233 and T73), which exhibit nonrecombinant alleles (variants B and E, respectively) quite different from that of strain 1477 (variant G), can only be explained by subsequent sexual reproduction. Convergent

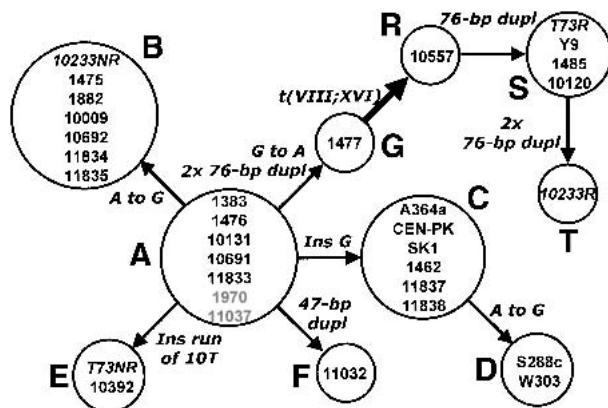


Figure 5 Maximum parsimony tree that minimizes the number of mutational events required to connect all the sequence variants of the *ECM34* gene promoter from different *Saccharomyces* strains N to N, nucleotide substitutions; ins, insertions; dupl; sequence duplications or repeats; 2x, a double event. A to G are nonrecombinant sequence variants, and R to T corresponds to the recombinant variants (*pECM34-SSU1* or *SSU1-R*) generated by the unequal crossing-over involved in the translocation t(VIII;XVI). The translocation event due to unequal crossing-over between microhomology regions located in the *ECM34* and *SSU1* promoters is indicated by a thick arrow. Strains in gray correspond to *S. pastorianus*. Strains in italics are heterozygotes for the translocation and contain both a recombinant (R) and a nonrecombinant (NR) variant.

evolution as an alternative explanation is discarded because it implies convergent changes in three characters: the A-G substitution, the gain of a run of Ts, and the loss of 76-bp repeats. Finally, a heterozygote for the translocation should also give rise to homozygotes such as strains Y-9, CECT1485, 10120, 10557, and 11827. The simplest explanation is that the fixation in homozygosis of this advantageous translocation conferring a higher resistance to sulfite occurred after sporulation and homothallic conjugation.

The observed selection for strains that contain several repeats of the 76-bp sequence suggests that the presence of these repeats increases sulfite tolerance. This fact was demonstrated by Goto-Yamamoto et al. (1998) with natural strains, and by Park and Bakalinski (2000) with genetically modified strains. In the present study, we corroborated this hypothesis by measuring sulfite tolerance of several wine strains containing different numbers of 76-bp repeats in their recombinant *SSU1* promoters (Table 2). From these results, it can be concluded that there is a direct relationship between the number of 76-bp repeats and sulfite tolerance, irrespective of the homozygotic or heterozygotic nature of the locus.

DISCUSSION

Translocations have been shown to be very common in *S. cerevisiae*; however, most of them are mediated through Ty or subtelomeric Y' element recombination (Kupiec and Petes 1988; Casaregola et al. 1998), especially in wine strains (Bidenne et al. 1992; Rachidi et al. 1999). Ectopic translocations through subtelomeric repetitive elements have also been proposed as the mechanism involved in the origin of some subtelomeric gene families such as *SUC*, *MAL*, *RTM*, or *MEL*, and have been correlated with the improvement of the features of some industrial yeast strains (discussed in Ness and

Table 2. Sulfite Tolerance of Yeast Strains Exhibiting Different Numbers of Repeats of a 76-bp Region in the Recombinant *SSU1* Promoter

Strain	Number of 76-bp repeats in recombinant <i>SSU1</i> promoter	Sulfite concentration (mM)				
		0	1	3	6	8
S288c	0 (nonrecombinant control)	+	+	–	–	–
T73	4	+	+	+	–	–
CECT10233	6	+	+	+	+/-	–
CECT10557	3 × 2 ^a	+	+	+	+/-	–
Y-9	4 × 2 ^a	+	+	+	+	–

Sulfite sensitivity was determined on YPD+TA plates containing 0–8 mM Na₂SO₃ as described in Methods. +, growth; –, no growth; +/-, poor growth scored after 24 h. Similar results were obtained in liquid medium in microtiter plates after 40 h.

^aThese strains are homozygous for the recombinant *SSU1* promoter.

Aigle 1995). However, in the present study we showed that the most probable cause of the reciprocal translocation between chromosomes VIII and XVI in wine yeasts is an illegitimate recombination mediated by microhomology. This kind of nonhomologous recombination is extremely rare in wild-type strains, occurring at a frequency of 3.5×10^{-10} . This frequency is only increased in double-strand break (DSB) repair-deficient mutants (Chen and Kolodner 1999). Thus, it seems that the new VIII^{XVI} and XVI^{VIII} chromosomes in wine yeast strains were probably generated by a spontaneous reciprocal translocation mediated by the fortuitous appearance of a broken chromosome end produced by a DSB in either of the two gene promoters, *ECM34* or *SSU1*. This end likely facilitated recombination with the other promoter through a very short homologous region. Whether this presumed recombination has been produced during mitosis or meiosis is currently unknown. This is the first time that such a rare event has been described in the evolution of yeast strains in the wild.

The enhanced expression of *SSU1* gene enabled wine yeast strains carrying the translocation to resist higher sulfite concentrations (Goto-Yamamoto et al. 1998, Park and Bakalinsky 2000; our results, Table 2). The new chromosomes, which did not lack any essential element, contained a new *SSU1* promoter with putative binding sites for the Fzf1p transcription activator within the 76-bp repeats (Avram et al. 1999). According to this hypothesis and the evolutionary analysis of DNA sequences performed, it seems likely that the 76-bp sequence was already repeated before the translocation event, giving rise to a higher expression of *SSU1* since the very beginning. We observed a clear relationship between the number of 76-bp repeats in the *SSU1* promoter and the level of sulfite resistance. This 76-bp sequence is partially palindromic and has a direct 6-bp repeat at both ends that may easily promote tandem repeat formation (Fig. 4). Similarly, a 147-bp repeated element found in *MAL* promoters from baker's yeast strains also alters gene expression (Bell et al. 1997).

The equilibrated chromosome pair VIII^{XVI} and XVI^{VIII} is quite frequent in wine yeasts. Thus, Goto-Yamamoto et al. (1998) and Bidden et al. (1992) also observed that among different wine yeast strains other than those studied here, eight were heterozygous and two homozygous for the *SSU1-R* allele (and hence, as demonstrated in the present study, heterozygous for the translocation), and that the *SSU1-R* allele (and hence, the translocation) was absent in the different nonwine strains analyzed. This higher frequency of the *SSU1-R* allele generated by the *t*(VIII;XVI) translocation in

wine yeasts can be explained by its adaptive value in wine-making environments where sulfite is widely used as a preservative. Wine strains of *S. cerevisiae* tolerate relatively high concentrations of sulfur dioxide as a result of adaptation and natural selection. This is due to the fact that sulfur dioxide is an antioxidant and antimicrobial agent that has been used in winemaking for millennia (Romano and Suzzi 1993). The Egyptians, and later the Greeks and Romans, made use of burning sulfur fumes to clean their wine containers. During the Middle Age, SO₂ became a widely used preservative, obtained originally by

burning sulfur but later by adding sulfite or bisulfite to musts. Nowadays, the use of SO₂ in winemaking is a common practice that is permitted by all wine-producing countries, in concentrations varying from 160 to 400 mg/L of total SO₂ or 20 to 100 mg/L of free SO₂. Moreover, sulfur dioxide resistance is an enological character used for the selection of commercial wine yeast, which is why many of the commercial strains analyzed exhibit the *t*(VIII, XVI) translocation (T73, this study and Goto-Yamamoto et al. 1998).

Finally, another conclusion that can be drawn from the present study is the role of sexual reproduction during wine fermentation. Sexual reproduction in wine *S. cerevisiae* yeasts has been a matter of controversy. Wine yeasts are prototrophic, homothallic, highly heterozygous and aneuploid, and exhibit low sporulation rates and spore viability (Bakalinsky and Snow 1990; Barre et al. 1993; Guijo et al. 1997). These characteristics, along with the observation of sexual isolation in yeast population during wine production (Guijo et al. 1997), are evidence favoring sexual reproduction as very rare, or even absent, in wine yeasts. However, the observation in the present study of translocation heterozygotes with very different nonrecombinant alleles supports the conclusion that sexual reproduction may be present in natural *S. cerevisiae* strains.

METHODS

Yeasts Strains and Culture Conditions

Forty-four strains of genus *Saccharomyces* were examined, all of them obtained from the Spanish Type Culture Collection (CECT). Three strains belong to the species *S. bayanus*, 30 to *S. cerevisiae*, three to *S. paradoxus* and two to *S. pastorianus*. The sources from which they were isolated are shown in Table 1. For laboratory culture, yeast cells were grown at 30°C in YPD (1% yeast extract, 2% bacteriological peptone, 2% glucose).

Sulfite tolerance was scored in YPD+TA (tartaric acid) agar plates as described by Park et al. (1999) by replicating cells grown in YPD plates. Alternatively, liquid YPD+TA containing 0–12 mM Na₂SO₃ was distributed in 100 µL aliquots in multiwell plates, and 2 µL of YPD-exponentially growing cells (0.2 OD₆₀₀) was added to every well. Growth was scored after 40 h.

PCR Reaction and Sequencing

A single yeast colony, taken using a micropipet yellow tip, was suspended in 100 µL of a PCR reaction mix containing 100 µM

deoxynucleotides, 1× reaction buffer, and 1 μM of these primers: ECM34D (5'-TCGAACATCGAGCATGCA-3'), ECM34R (5'-CCATATTGTGATGATATCG-3'), SSU1MD (5'-ACCTATCGAGTCTCCCAC-3'), SSU1R (5'-GACACCCATGACCATCAC-3'). The mixture was heated at 95°C for 15 min in a thermocycler, and 1 U of Biotools II DNA Polymerase (Biotools) was added to each tube. The PCR conditions were as follows: denaturation at 95°C for 5 min, followed by 30 cycles of denaturation at 94°C for 1 min, annealing at 52.5°C for 1 min, and polymerization at 72°C for 1 min. The polymerization was completed by one additional cycle of 5 min at 72°C.

The PCR product was separated on 3% (w/v) agarose gels with 1×TAE (40 mM Tris-Acetate, 1mM EDTA) buffer. After electrophoresis, gels were stained with ethidium bromide, visualized under UV light, and photographed. Molecular weights were estimated by comparison against a 100-bp DNA ladder. PCR bands were purified with a GeneClean II Kit (Bio101) and directly sequenced using the Ready Reaction DyeDeoxy Terminator Cycle Sequencing Kit (Perkin Elmer), following the manufacturer's instructions, in an Applied Biosystems automatic DNA sequencer model 3700.

Sequences were deposited in the EMBL database under accession numbers AF239757, AF239758, and AJ458340 to AJ458367.

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