Reconstruction of the Evolutionary History of Saccharomyces cerevisiae x S. kudriavzevii Hybrids Based on Multilocus Sequence Analysis

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Abstract

In recent years, interspecific hybridization and introgression are increasingly recognized as significant events in the evolution of Saccharomyces yeasts. These mechanisms have probably been involved in the origin of novel yeast genotypes and phenotypes, which in due course were to colonize and predominate in the new fermentative environments created by human manipulation. The particular conditions in which hybrids arose are still unknown, as well as the number of possible hybridization events that generated the whole set of natural hybrids described in the literature during recent years. In this study, we could infer at least six different hybridization events that originated a set of 26 S. cerevisiae x S. kudriavzevii hybrids isolated from both fermentative and non-fermentative environments. Different wine S. cerevisiae strains and European S. kudriavzevii strains were probably involved in the hybridization events according to gene sequence information, as well as from previous data on their genome composition and ploidy. Finally, we postulate that these hybrids may have originated after the introduction of vine growing and winemaking practices by the Romans to the present Northern vine-growing limits and spread during the expansion of improved viticulture and enology practices that occurred during the Late Middle Ages.

Introduction

The first evidence of production of fermented beverages dates back to 7000 BC in the Neolithic village of Jiahu in China [1], but the earliest evidence of winemaking is traced to Iran at the Hajji Firuz Tepe site (5400-5000 BC) [2]. From these origins in the slopes of northern Zagros, eastern Taurus and Caucasus Mountains, vineyards and grape wine production gradually spread to adjacent regions of the Fertile Crescent such as Mesopotamia and the Jordan Valley, and beyond, to the Eastern Mediterranean regions of Egypt, Phoenicia, Crete and Greece (5000 BC). Colonization by the Phoenicians, Carthaginians and Greek spread winemaking far across the Western Mediterranean regions of Southern Europe and Northern Africa. By 500 BC, wine was being produced in Italy, Sicily, Southern France, the Iberian Peninsula and the Maghreb. Vine cultivation was later extended by the Romans to the Northern limits of their empire (100 BC-100 AD). The next important expansion of winemaking was during the European colonization of America (16th century), South Africa (17th century), and Australia and New Zealand (18–19th centuries) [3,4].

On the other hand, beer elaboration is first recorded in the Mesopotamian region and in Egypt. Brewing diverged into two processes mainly differentiated by the prevailing fermentation temperature: ale, acquired from the Middle East by Germanic and Celtic tribes around the 1st century AD, and lager, which appeared during the Late Middle Ages in Europe [5,6].

A fortuitous domestication that acted on the S. cerevisiae populations is associated with wine and beer elaboration: it occurred as a consequence of the expansion of these fermentation processes. The first genetic diversity characterization of S. cerevisiae strains, isolated from different sources, showed clear differences between wild and domesticated strains [7]. Another study [8] evaluated the genetic variability of ~250 S. cerevisiae strains based on four nuclear gene sequences, and revealed for some genes the presence of two groups of alleles that differentiated wine strains from those isolated from other, non-wine, sources. Liti et al. [9] performed a genetic-population analysis based on whole genome sequences of 36 S. cerevisiae strains and reported the presence of five ‘clean’ (pure) lineages and different ‘mosaic’ (recombinant) strains. One of the ‘clean’ genotypic lineages comprises a number of wine strains from different geographic origins as well as European non-wine strains, and therefore, it was called wine/European population, the other lineages corresponded to strains isolated from other sources and origins [9].
In recent years, hybrids between *S. cerevisiae* and other *Saccharomyces* species such as the cryotolerant *S. uvarum* [10–12] and *S. kudriavzevii* [13–17] have been isolated from wine, cider and brewing fermentations, and other sources. These discoveries suggest that hybridization between different *Saccharomyces* species has been a frequent phenomenon in their evolution, particularly relevant during the adaption of *Saccharomyces* to fermentative conditions [18–20]. Some hybrids can be predominant even in the most Northern winemaking regions from Europe, very likely due to a better adaptation to growth at lower temperatures acquired from the non-*cerevisiae* parental, compared to *S. cerevisiae* [15,18,19,21].

Some reports carried out on a set of wine and beer *S. cerevisiae* and *S. kudriavzevii* hybrid strains suggested that those hybrids could be generated from hybridization between wine strains of *S. cerevisiae* and natural European strains of *S. kudriavzevii*; however, those results were not completely conclusive [17,22,23]. The aim of this study was to evaluate, by means of a multigeneic sequence approach, the potential origin of 24 *S. cerevisiae* x *S. kudriavzevii* and 2 *S. cerevisiae* x *S. kudriavzevii* x *S. uvarum* hybrid strains obtained from wine, beer and two other non-fermentative sources. The possible number of hybridization events that gave origin to the complete set of hybrids was also proposed based on the results obtained in this work and in previously reported data.

**Methods**

**Saccharomyces Strains, Culture Media and Nucleotide Sequences**

Twenty-six *S. cerevisiae* x *S. kudriavzevii* hybrid strains from different origins (Table S1) and seven strains belonging to *S. kudriavzevii* species (Table S2) were used in this study. Yeasts were grown at 28°C in GPY medium (2% glucose, 0.5% peptone, 0.5% yeast extract).

Nucleotide sequences corresponding to representative *S. cerevisiae* wine and non-wine alleles according to Arias [8] for genes BRE5, CAT8, EGT2 and GAL4 were also included in this study (Table S3 and Table S4).

Sequences for genes BRE5, CAT8, CYC3, CTR1, EGT2, CAT8, GAL4 and MET6 from *S. cerevisiae* strains (Table S2) representative of each pure population defined by Liti et al. [9] were obtained from SGRF (*Saccharomyces* Genome Resequencing Project, version 2 assemblies (20× coverage), except for strain RM11, which corresponded to version 1) (ftp://ftp.sanger.ac.uk/pub/dbmc/yeast/SGRF2/assembly/). In addition, sequences from wine strain EC1118 [24] were retrieved from GenBank database. Finally, *S. kudriavzevii* ZP591 and IFO 1802 sequences were downloaded from the *Saccharomyces sensu stricto* database (www.SaccharomycesSensuStricto.org).

**PCR Amplification and Sequencing**

DNA was extracted following the procedure described by Querol et al. [25]. Genes BRE5, CAT8, CYC3, CTR1, EGT2 and GAL4 were amplified by PCR, using primers CAT8_3, CYR1_5, MET6_5, MET6_3, MET6_3 K from González et al. [16] and newly designed primers (Table S5), obtained from the comparison among sequences from strains *S. cerevisiae* S288C and *S. kudriavzevii* IFO 1802 and ZP591.

Most primers were species-specific with the exception of those for genes CAT8, EGT2 and GAL4. The analysis of these genes required a previous step of cloning, performed by using a TOPO XL, PCR Cloning Kit (Invitrogen). To detect the *S. cerevisiae* alleles in clones, a screening was carried out by colony-PCR with the corresponding primers, and a subsequent digestion of the PCR fragments following the procedure described in González et al. [16].

PCR amplifications were performed by using conditions described in González et al. [16] in a G-Storm Thermocycler (G-Storm Ltd, UK). Amplification products were cleaned with a High Pure PCR Product Purification Kit (Roche Diagnostics, Mannheim, Germany) and both strands of the DNA were directly sequenced using the BigDyeTM Terminator V3.0 Cycle Sequencing Kit (Applied Biosystems, Warrington, UK), following the manufacturer’s instructions in an Applied Biosystems automatic DNA sequencer Model ABI 3730 (Applied Biosystems). Sequences were edited and assembled with Staden Package v1.5 [26] to be deposited in GenBank under accession numbers JN709116 to JN709440.

**Haplotype and Haplogroup Classification**

Gene sequences were aligned in MEGA 5 [27]. Haplotype classification was done in DnaSP v5 [28] using the previous haplotype number classification given by Arias [8]. New haplotypes were classified with consecutive Arabic numbers following the previous enumeration [8]. Median joining (MJ) networks [29] for BRE5, CAT8, EGT2, GAL4 were constructed using Network 4.5 (http://www.fluxus-engineering.com/).

**Phylogenetic Analysis and Supernetworks**

The neighbor-joining (NJ) and maximum-parsimony (MP) methods of phylogenetic reconstruction were applied to BRE5, CAT8, CYC3, CTR1, EGT2, GAL4 and MET6 separate sequence alignments of *S. cerevisiae* and *S. kudriavzevii* alleles from hybrid and reference strains described in Table S2. NJ trees were obtained with nucleotide distances corrected using the Maximum Composite Likelihood method. MP trees were obtained using the Close-Neighbor-Interchange algorithm in which the initial trees were obtained with the random addition of sequences (10 replicates). In all cases, a bootstrap analysis based on 2,000 pseudo-replicates was performed. For each gene, two NJ and MP phylogenetic trees were obtained, a tree based on *S. cerevisiae* alleles and another based on *S. kudriavzevii* alleles. Phylogenetic analyses were performed with MEGA 5 [27].

Two nexus files, with the collection of phylogenetic trees for *S. cerevisiae* and *S. kudriavzevii*, were created as an input of SPLITSTREE 4 package [30]. Two outputs corresponding to *S. cerevisiae* and *S. kudriavzevii* consensus super split networks (Supernetworks) were obtained, analyzing about 3.4 kb. For *S. cerevisiae* nexus file we reduced the number of splits setting maximum dimension parameter to 1, removing those splits in the network that are less supported. For the *S. kudriavzevii* nexus file we reduced the number of splits to simplify the final Supernetwork. For this simplification we applied the filtered Z-Closure method (filtering = 2). A filter of 2 takes into account those splits that are compatible in at least 2 input trees in the nexus file. The result is a network that summarizes the relationships found in at least two trees simplifying the network [31].

**Array Competitive Genomic Hybridization (aCGH) and Flow Cytometry**

Array competitive genomic hybridization (aCGH) experiments, scanning and data normalization were performed for IF6 and MR25 strains as previously described in Peris et al. [32]. A double-spotted array containing 6,240 ORFs of *S. cerevisiae* plus control spots totaling 6.4 K (Microarray Centre, University Health Network, Toronto, Canada) was used in aCGH assays. Raw and
normalized microarray data are available in ArrayExpress [33], under accession number E-MEXP-3375.

Caryoscopes were obtained using ChaRM v.1.1 [34]. Genome composition of IF6 and MR25 was inferred by combining aCGH (present study) and previous PCR-RFLP data [17]. aCGH was performed following the procedure described in Peris et al. [32].

The approximate locations of the recombination points in the mosaic chromosomes were determined from the up and down jump locations in the ORF’s mapping by microarray analysis of the hybrid yeast genomes. Collinearity between mosaic chromosomes were determined from the up and down intensity compared with the haploid [32]. Ploidy level was scored on the basis of the fluorescence enrichment with p-values, 0.05 were shown, after computing the Holm-Bonferroni for multiple hypothesis test correction. Significant GO terms were sorted according with their corresponding GO category.

The DNA content (C-value) of IF6 and MR25 was assessed by flow cytometry using a Beckman Coulter FC 500 (Beckman Coulter, USA) following the methodology described in Peris et al. [32]. Ploidy level was scored on the basis of the fluorescence intensity compared with the haploid S. cerevisiae S288c and diploid S. cerevisiae FY1679 reference strains.

Maximum Parsimony Tree of Chromosomal Rearrangements

A list of minimal number of chromosomal rearrangements, chromosomal losses and restriction site changes for IF6 and MR25 strains obtained in this work as well as data obtained from Belloch et al. [37] and Peris et al. [17,32] were included in the maximum parsimony analysis. A binary matrix was constructed to codify each particular event and these data were used to generate parsimony trees using MIX program from Phylip 3.66 package [38]. For this analysis, both chromosomal rearrangements and chromosomal gain/losses were considered as irreversible events (Camin-Sokal criterion), but data obtained from PCR-RFLP or sequence analyses were considered reversible events (Wagner criterion). The consensus tree was obtained by using the majority rule in the Consense program.

This binary matrix was also used to reconstruct a Median Joining Network, using Networks 4.5 [http://www.fluxus-engineering.com/], and a NeighborNet Phylowebnet, using SPLITSTREE 4 package [30].

Results

Phylogenetic Analysis of S. cerevisiae Genes from Hybrids

Phylogenetic relationships between S. cerevisiae x S. kudriavzevi natural hybrids obtained from several origins and a set of pure strains of the two parental species were analyzed to decipher possible common origins of these hybrids.

Nucleotide sequence data for both S. cerevisiae and S. kudriavzevi alleles of seven nuclear genes (BRE5, CAT8, CYC3, CYR1, EGT2, GAL4, and MET6) were obtained from a total of 24 natural S. cerevisiae x S. kudriavzevi and 2 S. cerevisiae x S. kudriavzevi x S. uvarum hybrid strains from several origins (Table S1). In a first phylogenetic analysis, we compared the S. cerevisiae sequences obtained for genes BRE5, CAT8, EGT2 and GAL4 from hybrids and from a representative selection, at the genotypic level, of 65 wine and 19 non-wine S. cerevisiae strains previously analyzed in our laboratory (Table S3 and Table S4). These genes were selected because they had shown high variability among S. cerevisiae strains from different origins [9]. Additionally, sequences from eight S. cerevisiae strains, five representative of the different “pure” lineages proposed by Liti et al. [9] and those from the completely sequenced genome of wine strain EC1118 [24] were also included in this study (Table S4).

Median-Joining networks (Figure 1) for all genes, except GAL4, showed two clearly differentiated groups of alleles or haplogroups. One haplogroup comprises those alleles present only in non-wine strains (so called non-wine alleles) and the second haplogroup includes alleles present in both wine and non-wine strains; however they are the only alleles exhibited by wine strains, and hence, they were called wine alleles. These wine alleles, when present in non-wine strains, are mainly found in heterozygosis with non-wine alleles. GAL4 is the exception because non-wine alleles were clustered into two haplogroups. The first group is characterized by the presence of a common deleted region of 15 bp, and the second comprises different lineages and appears to be closer to the wine alleles than to haplogroup 1 (Figure 1D).

Fourteen BRE5 alleles were present in hybrids (Figures 1 and S1), six are haplotypes already described in wine strains (32, 35, 41, 58, 66 and 95) and the other eight were new alleles (96, 97, 98, 99, 100, 108, 109 and 110). MR25, CECT 1388 and CECT 1990 are heterogeneous for this gene, exhibiting two wine S. cerevisiae alleles differing in one single nucleotide substitution (Figure S1). In the case of CAT8, 5 alleles from hybrids were present in wine yeasts (26, 33, 55, 57 and 91) and 2 were new (92 and 93). For EGT2, 2 alleles correspond to very common alleles in wine strains (3 and 5) and two were new (62 and 63), and finally GAL4 showed a higher diversity in hybrids with 3 already known alleles (1, 18 and 27) and 6 new [34, 89, 90, 92, 93 and 94]. These new alleles, found in hybrids for the first time, are indicated with asterisks in Figure 1. In general, alleles present in hybrids show few nucleotide differences (Figure S1) and are grouped together within the wine allele group for the four genes under analysis, with the exception of the BRE5 new allele 98 from the brewing strain CECT11011 which is located within the non-wine haplogroup, probably due to the presence of 2 convergent nucleotide substitutions.

Strains DBVPG6044, Y12, YPS128 and UWOPS03-461.4 were selected as representative strains of the West African, Sake, North American and Malaysian pure populations of S. cerevisiae, respectively, as defined by Liti et al. [9]. Sequences from these strains (indicated in bold in Figure 1) always clustered within the non-wine group for the four genes analyzed. To the contrary, L1528, EC1118 and RM11, three wine strains representative of the pure Wine/European genotypic lineage defined by Liti et al. [9], always appear within the wine allele group (alleles indicated in italics in Figure 1). The laboratory strain S288c clustered within the wine (for BRE5, CAT8 and GAL4) or non-wine groups (for EGT2) in accordance with its mosaic nature according to Liti et al. [9].

Because most S. cerevisiae alleles from hybrids are included within the wine allele group, the possible geographical origin of the hybrids was evaluated by analyzing the presence of these hybrid alleles in a set of 142 wine strains isolated from 8 different geographical areas, previously studied by Arias [8]. Table 1 shows the frequency of wine strains from each particular country sharing haplotypes with hybrids. The new alleles detected only in hybrids were not included in this analysis. As a general rule, the most frequent alleles in hybrids also corresponded to the most frequent alleles present in wine strains from several winemaking countries. For this reason, it is difficult to identify a specific geographic origin where hybridization processes may have occurred according to these comparisons (Table 1). Alleles 58 and 18 for BRE5 and GAL4...
respectively were not found among the *S. cerevisiae* wine strains analyzed (Table 1), but they were detected in some non-wine strains (Table S4 and ref. [8]). However, these two alleles clustered within the wine allele groups (Figure 1 A and D).

To identify how many putative *S. cerevisiae* parental strains were potentially involved in the origin of *S. cerevisiae* x *S. kudriavzevii* hybrids, we increased the number of genes analyzed in a second phylogenetic analysis. For this new analysis we included sequence data previously reported by Liti et al. [9] and Novo et al. [24] for comparative purposes.

Initial phylogenetic analyses on yeast were based on single gene sequences [39], but several times they failed to establish the overall history of these organisms. As an improvement, multigene sequence approaches using a concatenation of genes were proposed to construct the phylogenetic tree [40,41]; however, they would represent an oversimplified version of the genetic history [30]. As an alternative, the construction of consensus trees has also been proposed, but this method can be only used when each gene tree has the same taxa representation [42]. In this work, because some hybrid strains have lost some particular *S. kudriavzevii* genes, both concatenated or consensus trees would oversimplify the results. Recently, a Z-closure method has been proposed to overcome this kind of problem [30,31,43,44]. With this methodology, several gene trees with different taxa representation can be used as input files and a supernetwork with the complete set of taxa is obtained as [43]. However, one of the limitations of the Supernetwork analysis is the absence of statistical support, for this reason we interpreted our results according to a complementary phylogenetic analysis of the individual genes based on both Maximum Parsimony and Neighbor Joining. Both methods gave very similar or identical phylogenetic reconstructions (Figures S2 and S3).

A supernetwork, containing the information of 7 *S. cerevisiae* nuclear genes (Figure 2A), showed two well defined groups of strains: a group comprising non-wine strains Y12, DBVPG6044, YPS128 and UWOPS03-461.4 and a group containing wine strains RM11, L1528, EC1118 and all hybrids (Figure 2A). The position of strain S288c in this supernetwork proved again ambiguous due to the mosaic nature of this strain.

According to this supernetwork analysis of *S. cerevisiae* gene sequences, hybrid strains appear clustered in two main subgroups (C1 and C2) and several independent lineages (Figure 2A).
Table 1. Frequency of wine strains isolated from different countries showing the same alleles found in hybrids.

| Country     | Total number of strains | BRE5 | CAT8 | EGT2 | GAL4 |
|-------------|-------------------------|------|------|------|------|
| Argentina   | 37                      | 32*  | 49*  | 60*  | 81*  |
| Austria     | 30                      | 30*  | 10   | 40*  | 60*  |
| Chile       | 23                      | 23   | 10   | 60*  | 70*  |
| France      | 13                      | 23   | 46*  | 38*  | 92*  |
| Slovenia    | 5                       | 20   | –    | 80*  | –    |
| South Africa| 15                      | 33*  | –    | 33*  | 33*  |
| Spain       | 14                      | 21*  | 14   | 43*  | 93*  |
| Switzerland | 5                       | 40*  | –    | 80*  | 40*  |

*The most frequent haplotype.

aOnly those alleles present in more than one strain were included.

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Figure 2. Supernetworks obtained using data from seven nuclear genes (BRE5, CAT8, CYC3, CYR1, EGT2, GAL4 and MET6) for both *Saccharomyces cerevisiae* (A) and *Saccharomyces kudriavzevii* (B) alleles from hybrids, from reference *S. cerevisiae* [9,24] and *S. kudriavzevii* strains. Scale bar represents the edge’s weights inferred using the tree size weighted means options, a measure similar to those from branches in a phylogenetic tree.

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Subgroup C1 comprises Austrian (HA strains) and 3 other wine hybrids (PB7, SOY3 and Vin7), and the triple hybrids CID1 and CBS 2834, and subgroup C2 includes Swiss wine hybrids and Trappist beer strain CECT11003. The other hybrids appear in independent lineages (AMH, CECT 1990, 11002, 11004 and 11011) or in an ancestral position with respect to the two main subgroups (CECT1388, IF6 and MR25).

The supernetwork reconstruction method takes as input a set of complete or partial gene trees and produces a split network with the signals present in the gene trees, but it doesn’t allow to test the reliability of the the phylogenetic relationships. Therefore, bootstrap analyses for each individual gene Maximum-Parsimony and Neighbor-Joining gene trees, even when nucleotide diversities are lower than in the case of S. cerevisiae alleles. In those trees based on variable genes BRE5, CAT8, CYC3 and CYR1, Swiss wine hybrids and beer hybrids CECT 11003 and 11004 always appear within subgroup K2 (indicated in blue in Figure S2) and the wine hybrids from Austria (HA strains), VIN7 and SOY3 within subgroup K1 (indicated in yellow in Figure S2). In the case of hybrid IF6, this strain has lost two genes (CAT8 and CYC3), but for the other genes it shares the same alleles than hybrids from subgroup K1 (Figure S1).

The positions of the remaining strains change from one subgroup to the other, or to independent positions (Figure S2, alleles in green) depending on the gene (summarized in Table S6). Thus, brewing hybrids CECT1388, 1990, and 11002, and the clinical isolate MR25 lost 1–2 genes (including the shared loss of BRE5). In the CAT8 and CYC3 trees, these strains appear within subgroup K1, but for CYR1 they are included in a separate subgroup (indicated in green in Figure S2) due to the presence of allele 7, which differs from subgroup K1 allele 8 in a nucleotide substitution (Figure S1). Hybrid CECT11011 shares with the previous strains the CYR1 allele 7 and their inclusion within subgroup K1 in the CAT8 and CYC3 trees, but within subgroup K2 in the BRE5 tree, because maintains an allele identical to that from subgroup K2 strains. A similar situation is observed for triple hybrids CBS2834 and CID1, they appear within subgroup K2 in the BRE5 tree but within subgroup K1 in the other gene trees, including CYR1. Finally, the Spanish wine hybrid PB7 appears within subgroup K1 in two gene trees (CYC3 and CYR1), within subgroup K2 in other two (BRE5 and CAT8), and it exhibits a different allele for EGT2.

Genotypes of the Putative Parents of Hybrids Based on the Sequence Analysis of Seven Nuclear Genes

We tried to infer how many S. cerevisiae and S. kudriavzevii parents may have been involved in the generation of hybrids according to the phylogenetic analyses of the seven gene sequences. According to these sequences, the 24 double and 2 triple hybrids exhibit 20 different S. cerevisiae genotypes (allelic combinations) and 11 different S. kudriavzevii genotypes (Figure S1). These S. cerevisiae and S. kudriavzevii genotypes are found in 22 different combinations in hybrids. However, this does not mean that 22 different hybridization events occurred because hybrids are evolving after their origins. As seen before, the phylogenetic analysis of the sequences discriminate groups of alleles with putative common origins from an ancestral parental strain. In fact, the presence of rare alleles differing in few unique nucleotide substitutions (singleton) from the most common alleles in hybrids supports that these changes occurred after the hybridization process.

By considering the phylogenetic relationships among alleles and their combinations in hybrids (summarized in Table S6), we could infer 6 S. cerevisiae and 6 S. kudriavzevii putative ancestral genotypes (parental strains) that are arranged in 10 hybrid combinations (possible hybridization events). The first main hybrid combination is present in 6 wine hybrids, four from Austrian (HA strains), one from South Africa (VIN7, likely of European origin) and another from Croatia (SOY3). This SOY3 strain shares identical or closely related S. cerevisiae and S. kudriavzevii alleles with the other strains of this group for all genes except BRE5, which shows 4 nucleotide differences. This allele appears in the BRE5 gene as closer to alleles from other hybrids (Figure 2B). These similarities could be
explained by convergent substitutions, but we cannot rule out the possibility that the parental strain were originally heterozygous for BRE5 and suffered a subsequent differential loss of heterozygosity in each derived hybrid lineage.

The second main combination is found in the 8 wine double hybrids from Switzerland and the Trappist beer hybrids CECT11003 and 11004 from Belgium. In this group, a slight discrepancy is also observed in strain CECT11004. This strain exhibits a MET6 allele (allele 1) different to that present in other strains of this group (allele 2), but identical to that exhibited by strains from other groups (Figure S1). However, these MET6 alleles 1 and 2 differ in one single synonymous substitution and a simple convergent change may explain this difference. An alternative explanation would be to consider allele MET6-1 as the ancestral one present in the S. cerevisiae parent of this group of hybrids later originating the derived allele MET6-2 shared by the Swiss and CECT11003 hybrids.

In the remaining hybrid combinations, both S. cerevisiae and S. kudriavzevii genotypes basically correspond to different arrangements of the alleles present in the first and second hybrid combinations described before. One explanation is that these recombining genotypes, generated by sexual mating at the within species level, were already present in the S. cerevisiae and S. kudriavzevii population before the hybridization events occurred. In this case, a minimum of 10 hybridization events would be necessary to explain the origin of these hybrids. However, another compatible explanation is that some hybrids may have originated by rare mating between diploid heterozygous cells, and a subsequent segregation of alleles due to chromosome loss (most hybrids are triploid [21,32]), or random loss of heterozygosity due to recombination and/or gene conversion would generate the different mosaic hybrids. In this case, the number of hybridization events would be smaller than ten. This could be the case of brewing strains CECT1388, 1990, 11002, 11011 and the clinical isolate MR25. These strains exhibit similar S. kudriavzevii genotypes (including the specific allele CTIR1-7), but different S. cerevisiae allele combinations, including wine and non-wine alleles (CTIR1-2 and -4 in strains CECT1990 and 11011).

The Genome Constitution of Non-fermentative Hybrids IF6 and MR25

In previous studies, we analyzed the genetic diversity of S. cerevisiae x S. kudriavzevii hybrids by RFLP analysis of 35 nuclear genes [16,17] combined with array comparative genome hybridization (aCGH) [32,37]. These analyses provided us information on the genome rearrangements occurred in the hybrids after their origins. Most of these rearrangements are non-reversible events that can complement the information obtained with the phylogenetic analysis of gene sequences to unveil the origin and evolution of these S. cerevisiae x S. kudriavzevii hybrids.

However, the genome constitutions of hybrids IF6 and MR25 were not characterized in our previous studies, and therefore, they were subjected to aCGH and flow cytometry analyses to assess their genome compositions. Our results indicated that DNA content of IF6 and MR25 were 3.25 and 2.92 times that of the reference haploid strain S288c, respectively. These DNA content values, together with the aCGH analysis and PCR-RFLP data for 35 nuclear genes previously reported [17], allowed us to detect the presence of three chimerical chromosomes in hybrid IF6 (chr. X, XII and XIII) and five in MR25 (chr. IV, VII, IX, XII and XIV (Figure 3). The hypothetical recombination points were mapped according to the Saccharomyces genome described in the SGD database [http://db.yeastgenome.org] using a window size of 15–20 Kb [four genes in the left and right of the most plausible recombination point]. These recombination points were located in sequences corresponding to Ty LTRs, ARS and rRNAs (Table S7). RFLP analysis of genes located at the end of chromosomes [17] confirmed the presence of S. kudriavzevii segments in chromosomes VII and IX from IF6, and chromosomes X and XIII from MR25, however, their putative chimerical nature could not be detected by the aCGH analysis (Figure 3).

Following the same methodology used in our previous study [32], we obtained a list of S. cerevisiae genes lost in both hybrids IF6 and MR25. Both IF6 and MR25 have depleted a similar number of genes classified as retrotransposons as well as genes belonging to the ASP3, CUP1 and ENA clusters (Table S8). In particular, hybrid IF6, obtained from a dietary supplement, exhibited a deleted region (YLR155C-YLR256W) in its S. cerevisiae chromosome XII (Figure 3). This region is adjacent to the rDNA repeat region located between YLR154C and YLR155C, which is not included in the microarray platform. A PCR amplification of the 5.8S-ITS region and the subsequent restriction analysis [45], revealed the absence of S. cerevisiae rDNA genes in this region (data not shown).

With respect to their S. kudriavzevii subgenome, IF6 and MR25 hybrids lost ~33% and ~18% of the total S. kudriavzevii genes, respectively. Gene Ontology (GO) enrichment analysis applied to the common set of S. kudriavzevii genes maintained by the two hybrids, demonstrated a high frequency of stress response genes among those S. kudriavzevii genes conserved in both hybrids (Table S9). Some of the significant GO terms shared by MR25 and IF6 are “response to stimulus” with p-values <0.05. In the case of MR25 is also important to note the significant GO term “cellular lipid metabolic process” and “response to stress” (p-value <0.05).

Analysis of the Number of Hybridization Events

Genome composition data obtained for the 26 S. cerevisiae x S. kudriavzevii hybrids from this study as well as from previous studies [32,37] were used to reconstruct a parsimony tree based on the presence of chimerical chromosomes, on the absence of chromosomes from one or another parental strain and the presence of specific allelic variants. Using the information from this parsimony tree together with the putative genetic constitution of the hypothetical parental strains obtained from the phylogenetic analysis of nuclear gene sequences, as well as from COX2 sequences also obtained in our previous studies [17], allowed us to reduced the number of hybridization events to a minimum of six for the S. cerevisiae x S. kudriavzevii hybrids under analysis, and two additional events for the origin of the S. cerevisiae x S. kudriavzevii x S. warau triple hybrids. The putative ploidies of the parental cells involved in hybridization were also estimated by analyzing the genomic constitution of the hybrids derived from each event.

Figure 4 shows five out of the six different origins for double hybrids proposed according to this study. AMH is not included due to its complex genome structure, because it is a tetraploid hybrid that lost most of the S. kudriavzevii subgenome [17,32]. Independent origin for AMH is clearly supported by the different sets of data used in this analysis.

Wine hybrid strains from Switzerland (W27, SPG14–91, SPG16–91,126, 172, 319 and 441), and the Trappist brewing strains CECT11003 and CECT11004 share a common origin. Their nuclear genomes derive from a hybridization event between the hypothetical S. cerevisiae CG2 and S. kudriavzevii KG2 parents (Figure S1). They inherited their mtDNA type K2 from S. kudriavzevii [17]. Hybrid W46 was also included in this group although it exhibits a mitochondrial type K3 (Figure 4), derived from K2 by a single nucleotide difference [16].

The group of Austrian hybrids HA as well as wine hybrids VIN7 and SOY3 have also a common origin in a hybridization
event involving hypothetical parents *S. cerevisiae* CG1 and *S. kudriavzevii* KG1 (Figures 2, 4 and S1), sharing the mitochondrial type K4 from *S. kudriavzevii* [17].

A third group includes the brewing triploid hybrids CECT1388, CECT1990, CECT11002, CECT11011 and the clinical isolate MR25, sharing several genome rearrangements and restriction patterns as well as a recombinant mtDNA type K6 [17]. According to the seven gene sequence analysis, these strains seem to have independent hybridization origins from crosses between different *S. cerevisiae* parents (CG3, CG4 and CG5) but the same *S. kudriavzevii* strain KG3 characterized by a specific *CYR1* allele (Figure S1). These contradictory results may be explained by considering a heterozygous *S. cerevisiae* diploid cell containing wine and non-wine alleles as the parental strain, as mentioned above.

Wine strain PB7 from Leon, Spain, was included in the same subgroup than the Austrian wine strains according to the supernetwork analyses, due to network simplification (Figure 2). However, this strain likely originated in an independent hybridization event because it derives from different parents, the mosaic *S. cerevisiae* CG6 and *S. kudriavzevii* KG5 genotypes (Figure S1), exhibits a recombinant mtDNA K10 [17], and finally, it possesses a tetraploid genome [32].

In the case of IF6, although it shares the same *S. kudriavzevii* KG1 ancestor with Austrian hybrids, its *S. cerevisiae* parental strain is clearly different: a mosaic CG5 genotype closer to the *S. cerevisiae* parent of the brewing hybrids. The different hybrid combination of parental genotypes supports an independent origin for this strain.

**Discussion**

By analyzing the sequences of four nuclear genes from a total of more than 250 *S. cerevisiae* yeast isolates from wine (Europe, South America and South Africa) and non-wine origins (wild, brewing, cider, sake and traditional beverage fermentations mainly from
Latin America, but also from Africa and Asia), Arias [8] demonstrated the existence of two groups of alleles, those present only in strains isolated from non-wine sources, called non-wine alleles, and another group of alleles that, while they also appear in non-wine alleles, they are the only alleles present in wine strains (wine alleles). These wine alleles are much less frequent in non-wine strains, and they mainly appear in heterozygosis with non-wine alleles. Liti et al. [9] obtained the complete genome sequences of 37 *S. cerevisiae* strains from different sources of isolation and geographic origins. The phylogenetic analysis of nucleotide polymorphisms showed a complex *S. cerevisiae* population structure. Liti et al. [9] observed five genotypic lineages, called according to their origins or source of isolation as Malaysia, West Africa, sake, North America and ‘Wine/European’, which exhibited the same phylogenetic relationships across their entire genomes. The strains from these five lineages were considered as ‘clean’, pure strains, representative of diverged populations. The other strains evidenced variable phylogenetic relationships depending on the genome region analyzed, and were considered as ‘mosaics’ with a mixed genome architecture that could be due to human traffic in yeast strains and subsequent recombination between them. The analysis of the sequences of the same four gene regions used by Arias [8] indicated that alleles present in the four non-wine lineages fell within the group of non-wine alleles; alleles present in strains of the wine/European lineage were included within the ‘wine allele’ group, and the locations of the ‘mosaic’ alleles were variable depending on the gene. Because Liti et al. [9] sequenced derivative monosporic cultures, some of the ‘mosaic’ parental strains could be heterozygous for wine and non wine alleles for many genes, as observed by Arias [8]. High levels of heterozygosity for non-wine yeast were also observed by Fay and Benavides [7], and for ale strains by Dunn and Sherlock [46].

The accessibility to such a collection of sequences (including genome sequences) from *S. cerevisiae* strains from different sources of isolation and geographic origins was an excellent opportunity to decipher the nature of the *S. cerevisiae* parents involved in the origin of hybrids. This way, for all genes under analysis, *S. cerevisiae* alleles from hybrids were always clustered within the wine allele group, with the exception of the *BRE5* allele from the brewing strain CECT11011, which clustered in the non-wine group, and *CYR1* allele from the brewing CECT1388 and CECT11011, which clustered with non-wine strains from Liti et al. (2009) in the
individual gene trees. Moreover, the phylogenetic supernetwork analysis of *S. cerevisiae* alleles from hybrids identified two main subgroups of *S. cerevisiae* parental strains, and due to its simplification it failed to detect mosaic *S. cerevisiae* genotypes. It followed that the *S. cerevisiae* CG2 parental strain was involved in the hybridization event that originated the complete group of wine Swiss hybrids and *S. cerevisiae* CG1 was involved in the origin of the Austrian wine hybrids, SOY3 and Vin7.

The aCGH analyses of hybrid genome composition (present study and [32]) showed the depletion or underrepresentation of certain *S. cerevisiae* genes (Ty retrotransposons and *EM1* and *JSP* gene families), which were proposed as genomic signatures for wine *S. cerevisiae* yeasts [47,48], which is in agreement with the postulated wine origin of the *S. cerevisiae* parental strains involved in the generation of these hybrids. The maintenance of *S. kudriavzevii* genes related to stress response, in MR25 and IF6, and lipid metabolism, in MR25, also confirms the importance of *S. kudriavzevii* subgenome in cold stress resistance, postulated in previous studies [32].

In the case of IF6, aCGH and PCR confirmation of 5.8S-ITS regions support the loss of heterozygosity (LOH) of the rDNA region in chromosome XII, maintaining only the *S. kudriavzevii* sequences for this region. This region has been characterized, in plants and animals, to be under concerted evolution [49–51]. This has been also observed in a natural hybrid *S. pastorianus* (CBS 1538 strain), where the *S. cerevisiae* tDNA region of chromosome XII has been lost [52].

The wine origin of the *S. cerevisiae* parent of most *S. cerevisiae* x *S. kudriavzevii* hybrids has already been postulated in previous works based on genomic composition data inferred by aCGH and PCR-RFLP analysis [32], as well as by microsatellite analysis [21]. The use of a multilocus sequence analysis approach certainly confirms the wine origin of the *S. cerevisiae* strains involved in the generation of most *S. cerevisiae* x *S. kudriavzevii* hybrids.

The exceptions are the brewing hybrid CECT11011, in which a possible recombinant *BRE5* allele is present, and CECT11900 and CECT11011, which contain *CYR1* non-wine alleles, and hence, a heterozygous non-wine *S. cerevisiae* strain, with both wine and non-wine alleles, could be involved in their origin. Dunn and Sherlock [46] demonstrated that *S. pastorianus* hybrids, responsible of lager beer fermentations, very likely derived from a cross between a haploid *S. bayanus*-like strain, later identified as belonging to the new species *S. eubayanus* [53], and a diploid *S. cerevisiae* strain, related to ale brewing strains, which are characterized by a high heterozygosity. Arias [8] also included in his study several ale strains that showed as heterozygous, for wine and non-wine alleles. Therefore, the parental *S. cerevisiae* involved in the origin of brewing hybrids CECT11900 and CECT11011 could be an ale strain originally heterozygous for wine and non-wine alleles. Another brewing hybrid, strain CECT11002, appeared as related to the brewing hybrids and the clinical isolate, but it did not contain non-wine alleles for the genes under analysis; all these hybrids may also have been originated from a similar ale parental strain. Erny *et al.* [21] included in their microsatellite analysis a Chimay strain which clusters with the *S. cerevisiae* brewing strains. We do not know whether their Chimay strain and our CECT11002 (also from Chimay) is the same or not, but at least they should be related, which could corroborate the ‘ale’ origin of their *S. cerevisiae* parent. Genome sequencing of one of these strains will elucidate this hypothesis.

By using the population genetic information from Arias [8], we also tried to determine the exact geographic origin of the parental *S. cerevisiae* strains. We looked for particular *S. cerevisiae* strains from different wine regions possessing the combination of alleles present in the hypothetical parental *S. cerevisiae* strains. With the exception of one *CAT8* allele, genotype CG1 was present in strains from Chile, South Africa, Switzerland and Spain; and genotypes CG2 and CG3, with the exception of *BRE5*, were found in strains from Argentina, Chile, Italy, Japan, South Africa, Austria, France and Spain. Other genotypes, with slight differences were found in Argentina, Chile, South Africa, Austria, Slovenia, Switzerland, Italy, Japan, France and Spain. As strains from the new winemaking regions (South America and South Africa in this case) were introduced from Europe with vines and winemaking tools, the most probable geographic origin for hybridization, according to the *S. cerevisiae* hypothetical parental genotype, is Europe.

The European origin of hybrids is also supported by the phylogenetic analysis of *S. kudriavzevii* alleles. Alleles present in hybrids were detected among European *S. kudriavzevii* pure strains. Three of seven alleles of *S. kudriavzevii* KG1, were found in 3 *S. kudriavzevii* strains from Ciudad Real (Spain), Castellon (Spain) [23] and Portugal [22]. However, other genotypes have not been found among the few *S. kudriavzevii* pure strains available. Future surveys on the genetic variability of European populations of *S. kudriavzevii* may be of interest to decipher the geographic origin of hybridization, because this wild species has not been subjected to human traffic and it may preserve its original population structure in the same way than *S. paradoxus* [54,55]. A recent study [21], complementary to the present one, on the possible origin of a different set of European *S. cerevisiae* x *S. kudriavzevii* hybrids from winemaking (only four Swiss hybrids and VIN7 are in common), carried out by means of microsatellite information, also confirmed the European origin of the putative parental strains of hybrids.

By combining the phylogenetic analysis of gene sequences with all the available information on genetic and genomic characterization of *S. cerevisiae* x *S. kudriavzevii* hybrids [16,17,21,32,37], a total of six potential hybridization events were determined. The first hybridization event involved a haploid *S. kudriavzevii* parent KG2 with mtDNA K2 and a diploid *S. cerevisiae* parental CG2. This event originated all Swiss hybrids and the related Trappist brewing strains CECT11003 and 11004. This clearly independent origin for Swiss wine hybrids is in accordance with the microsatellite phylogenetic analysis of hybrids performed by Erny *et al.* [21].

A second hybridization event involving a haploid *S. kudriavzevii* KG1 with mtDNA type K4 (found in all hybrids from this group) and a diploid *S. cerevisiae* CG1 originated a lineage of hybrids widely distributed in different wine regions such as Austrian hybrids, the Croatian strain SOY3, and the South African hybrid VIN7 of putative European origin according to Erny *et al.* [21]. These authors observed in their study that VIN7 is included in the same group as other Abatian and German wine hybrids and bears a close relationship to Hungarian wine hybrids, confirming an European origin for VIN7. Therefore, this is a lineage of wine hybrids widely distributed from the Rhine valley (Alsace and Germany) to the Danube valley (Pannonian region: Austria, Croatia and Hungary).

A third hybridization event was involved in the origin of a lineage of brewing strains also widely distributed in ale breweries from England, Germany, Belgium (Chimay Trappist Abbey), New Zealand and the clinical isolate MR25. This hybridization event involved a haploid *S. kudriavzevii* parental close to K2, KG3 strain, and probably a heterozygous diploid *S. cerevisiae* parental. An “ale” *S. cerevisiae* strain heterozygous for wine and non-wine alleles could be involved in the origin of this group of hybrids.
Hybrid PB7 was probably originated from two diploid cells derived from mosaic strains *S. cerevisiae* CG6 and *S. kudriavzevii* KG5. Its tetraploidy [32] and the presence in this hybrid of a recombinant mtDNA [17] supports an independent hybridization event.

Independent origins are postulated for hybrids IF6 and AMH. In the case of AMH, its complex tetraploid genome [32], in which most of the *S. kudriavzevii* subgenome is lost [17], led us to suspect a possible scenario in which a diploid *S. cerevisiae* crossed with a haploid *S. kudriavzevii* strain and, after sporulation or a drastic *S. kudriavzevii* genome reduction, a diploid spore or an evolved derivative backcrossed with a diploid *S. cerevisiae*. IF6 was originated from a cross between a diploid *S. cerevisiae* CG5 mosaic genotype and a haploid *S. kudriavzevii* KG1, identical to the one involved in the origin of Austrian hybrids. Therefore, the possibility of a common origin with Austrian HA, VIN7, and SOY3 hybrids cannot completely ruled out if a heterozygous *S. cerevisiae* ancestor were involved in the hybridization event. However, this hypothesis not only requires the differential loss or segregation of alleles in the IF6 and Austrian lineages, but also the independent acquisition of the mitochondrial genome from the hybrid zygote, *S. cerevisiae* type C2 in IF6 and *S. kudriavzevii* type K4 in the Austrian lineage. This is possible in hybrid zygotes where three types of mitochondrial genomes may be present: two from each parental and a recombinant, generated after mitochondria fusion [56], but mitochondrial sorting occurs from the first budding formation [57], generating independent lineages that are difficult to distinguish from independent hybridizations in which parental relatives were involved.

Finally, triple hybrids *S. cerevisiae* x *S. kudriavzevii* x *S. warau* are not shown in Figure 4, also due to to their complex origins, in which a secondary hybridization was involved. However, the supernetwork analysis and gene trees information indicates that CBS2834 and CID1 were probably derived from the same (or similar) *S. kudriavzevii* parent (KG6) but different *S. cerevisiae* parental strains, the same than the Austrian strains (CG1) for CBS2834 and similar to PB7 (CG6) for CID1.

Finally, the origin of the triple hybrids CID1 and CBS2834 is not clear due to the additional occurrence of a secondary hybridization event either between a *S. cerevisiae* x *S. kudriavzevii* hybrid or derivative with a *S. warau* strain or between a *S. cerevisiae* x *S. warau* hybrid or derivative with a *S. kudriavzevii* strain. However, CID1 and CBS2834 were probably originated from independent hybridization events.

Most hybrids seem to have been generated by rare-mating events involving a diploid *S. cerevisiae* strain and a haploid strain of *S. kudriavzevii* generating different chimerical genomes with ploidy values close to 3 n. This is most clear for brewing strains (CECT1388, CECT1990, CECT11011 and MR25) where heterozygous genes could be observed. In PB7, which exhibited a ploidy value of 3.96, two diploid parents could be involved. Rare-mating has already been proposed as a mechanism for natural hybrid generation [58]. Additionally, artificial hybrids generated by rare mating are easily obtained in laboratory conditions [59].

Hybrid distribution and their physiological properties, together with the conclusions of recent studies on the population-genetic structure of *S. cerevisiae* [7,9] as well as the phylogenetic analyses performed in the present study, can be used to speculate a possible scenario for the hybridization process. Grapevine [60] and barley [61] domestication mainly occurred in the Middle East, where the earliest archaeological evidence of winemaking [62] and brewing [63] have been discovered. From these areas of domestication, there was a gradual radiation to adjacent areas of the Mediterranean, European regions of Europe and Africa, following the spread of Phoenician, Greek and Carthaginian civilizations. Finally, the expansion of grape growing and winemaking to temperate regions of Oceanic, Greek and Carthaginian civilizations. Finally, the expansion of grape growing and winemaking to temperate regions of Oceanic and Continental climates of Europe, following the main trade fluvial routes, was performed under the influence of the Romans, who would take vine-growing to the limits of their empire, the Rhine and Danube Rivers. By the end of the Roman Empire, grape growing was common in most European locations. In the Middle Ages, viticulture and enology were improved and expanded by Christian monks.

Recent studies on the genetic diversity of *S. cerevisiae* populations [7–9,64] show that wine strains constitute a genetically differentiated population that could have appeared during the process of adaptation to winemaking conditions, a process of fortuitous domestication of a *S. cerevisiae* wine strain. The microsatellite population analysis of *Saccharomyces* strains [65] also suggests that this population likely originated in the North East and spread during the expansion of grapevine and winemaking.

About 2,000 years ago, wine *S. cerevisiae* yeasts were likely taken by the Romans, together with the vines and winemaking tools, to the Northern limit of grapevine distribution. There, *S. cerevisiae* wine strains, even nowadays, have problems when performing wine fermentations at the lower temperatures to which other *Saccharomyces* species are better adapted [66]. In these regions, cryotolerant species, such as *S. bayanus* var. *warau*, may out-compete *S. cerevisiae* [11,67–69]. Under such circumstances, however, hybrids may have advantages over the parental species [19,70,71]. This is due to the acquisition of physiological properties from both parents, which provide a mechanism for selection of hybrids [10,18,72,73]. In the case of *S. cerevisiae* x *S. kudriavzevii* hybrids, they acquired good alcohol and glucose tolerances and fast fermentation performances from *S. cerevisiae* [19,74] and a better adaptation to low and intermediate temperatures from *S. kudriavzevii* [18,19,71].

These *S. cerevisiae* x *S. kudriavzevii* hybrids likely appeared several times, according to this study, and became frequent in some areas of the Northern limit of vine growing, but they could probably spread in Central Europe with the expansion of vine growing and winemaking practices that occurred during the Middle Ages [4]. Winemaking was preserved and improved during the Middle Ages by Christian monks. Benedictine abbeys were the main wine producers and traders, but the Cistercian reformation made possible the main revolution in winemaking improvements and vine growing extension [75].

From their original abbeys in Burgundy, Cistercians spread across Europe during the 11th and 12th centuries to establish more than 300 abbeys. During this expansion, the white monks spread the viticulture and enology practices to the Rhine and Danube valleys and the Pannonian basin of Central Europe [75]. They extended the Burgundian family of grape varieties, mainly Chardonnay and Pinots, as well as German varieties, and with them likely the hybrid yeasts responsible for wine fermentation.

In the regions where the main lineages of *S. cerevisiae* x *S. kudriavzevii* wine hybrids have been found, winemaking was introduced or improved on by Cistercian monks. In fact, the Cistercian order is given credit for planting in the French regions of Burgundy, Chablis, Loire, Rhone, Champagne (where the *S. cerevisiae* x *S. kudriavzevii* hybrid EPII, also called Epernay 2, was isolated [76]), Alsace (where many hybrids are also present and predominant [21]) and in several other wine regions in Central Europe. Some of these regions are: Rheingau Wine Region in Germany, where hybrid AMH (Assmannshausen) and those from Geisenheim [21] were isolated; Thermanregion, Austria, where HA hybrids, characterized in this study, were found as pre-
dominant in vineyards [15]; Slavonian Croatia, where SOY3 was isolated; and Hungary, where these hybrids have also been found [21].

Supporting Information

Figure S1 Polymorphic sites of the S. cerevisiae alleles (A) and S. kudriavzevii alleles (B) present in double and triple hybrids, as well as representative strains of the parental species. Groups of hybrid alleles are colored according to their phylogenetic relationships based on the Maximum-Parsimony and Neighbor-Joining gene trees depicted in Figures S2 and S3, respectively. An asterisk indicates new allele not described in previous studies.

Figure S2 Maximum-Parsimony phylogenetic trees of 7 nuclear gene partial sequences of the S. cerevisiae (C) and S. kudriavzevii (K) subgenomes present in hybrids. Sequences from pure strains representative of the parental species were also included in the analysis. S. cerevisiae reference strains isolated from wine and non-wine sources are indicated in red and blue, respectively. Numbers on the branches are nucleotide substitutions. Bootstrap values (in %) based on 2000 pseudo-replicates are given between parentheses. Groups of related hybrid alleles are highlighted in different colors, groups C1 and K1 in yellow, C2 and K2 in blue and C3 and K3 in green.

Figure S3 S. cerevisiae (C) and S. kudriavzevii (K) phylogenetic gene trees reconstructed using the Neighbor-joining method of the seven nuclear partial genes. Scales are given in nucleotide substitutions per site, numbers on the nodes are bootstrap values (in %) based on 2000 pseudo-replicates. In the S. cerevisiae phylogenetic gene trees, representative wine and non-wine S. cerevisiae strains are colored in red and blue, respectively. In the case of S. kudriavzevii phylogenetic gene trees, Japanese and European S. kudriavzevii strains are indicated in red and blue, respectively. Groups of related hybrid alleles are indicated in different colors, groups C1 and K1 in yellow, C2 and K2 in blue and C3 and K3 in green.

Table S1 Geographic origins, source of isolation and genetic constitution of Saccharomyces cerevisiae x Saccharomyces kudriavzevii hybrids.

Table S2 Geographic origins, source of isolation and genetic constitution of reference Saccharomyces kudriavzevii and Saccharomyces cerevisiae strains. S. cerevisiae strains included in this Table correspond to representative strains belonging to the 'pure' lineages described by Liti et al. [9] based on their genome sequences, as well as to wine strain EC1118 [24].

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