



Genomic and Transcriptomic Basis of *Hanseniaspora vineae*'s Impact on Flavor Diversity and Wine Quality

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ABSTRACT *Hanseniaspora* is the main genus of the apiculate yeast group that represents approximately 70% of the grape-associated microflora. *Hanseniaspora vineae* is emerging as a promising species for quality wine production compared to other non-*Saccharomyces* species. Wines produced by *H. vineae* with *Saccharomyces cerevisiae* consistently exhibit more intense fruity flavors and complexity than wines produced by *S. cerevisiae* alone. In this work, genome sequencing, assembling, and phylogenetic analysis of two strains of *H. vineae* showed that it is a member of the *Saccharomyces* complex and it diverged before the whole-genome duplication (WGD) event from this clade. Specific flavor gene duplications and absences were identified in the *H. vineae* genome compared to 14 fully sequenced industrial *S. cerevisiae* genomes. The increased formation of 2-phenylethyl acetate and phenylpropanoids such as 2-phenylethyl and benzyl alcohols might be explained by gene duplications of *H. vineae* aromatic amino acid aminotransferases (*ARO8* and *ARO9*) and phenylpyruvate decarboxylases (*ARO10*). Transcriptome and aroma profiles under fermentation conditions confirmed these genes were highly expressed at the beginning of stationary phase coupled to the production of their related compounds. The extremely high level of acetate esters produced by *H. vineae* compared to that by *S. cerevisiae* is consistent with the identification of six novel proteins with alcohol acetyltransferase (AATase) domains. The absence of the branched-chain amino acid transaminases (*BAT2*) and acyl coenzyme A (acyl-CoA)/ethanol *O*-acyltransferases (*EEB1*) genes correlates with *H. vineae*'s reduced production of branched-chain higher alcohols, fatty acids, and ethyl esters, respectively. Our study provides sustenance for understanding and potentially utilizing genes that determine fermentation aromas.

IMPORTANCE The huge diversity of non-*Saccharomyces* yeasts in grapes is dominated by the apiculate genus *Hanseniaspora*. Two native strains of *Hanseniaspora vineae* applied to winemaking because of their high oenological potential in aroma and fermentation performance were selected to obtain high-quality genomes. Here, we present a phylogenetic analysis and the complete transcriptome and aroma metabolome of *H. vineae* during three fermentation steps. This species produced significantly richer flavor compound diversity than *Saccharomyces*, including benzenoids, phenylpropanoids, and acetate-derived compounds. The identification of six

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proteins, different from *S. cerevisiae* ATF, with diverse acetyltransferase domains in *H. vineae* offers a relevant source of native genetic variants for this enzymatic activity. The discovery of benzenoid synthesis capacity in *H. vineae* provides a new eukaryotic model to elucidate an alternative pathway to that catalyzed by plants' phenylalanine lyases.

KEYWORDS flavor compounds, genome, Illumina, metabolome, transcriptome, wine aroma

It is well known that yeast transforms grape sugars to ethanol and CO₂ as the main wine fermentation products; however, cell secondary metabolism generates the highest impact compounds that dramatically affect the final flavor of wine. Flavor traits matter most in fermented beverages and should be considered the key properties when developing yeast selection within food biotechnology industries (1, 2). In wine, non-*Saccharomyces* yeast strains that account for more than 99% of the grape native flora are still poorly explored (2), and their impact on flavor richness will require multidisciplinary studies from genetics to metabolomic analyses of yeast cells. The limited numbers of commercial yeast strains applied by today's winemakers are not contributing to flavor diversity, decreasing the possibilities to obtain more differentiated wine styles. Besides the grape selection and viticulture and vinification technologies used, which have been traditionally emphasized for quality wine production, yeast aspects should be taken into account. In a highly competitive market with more than one million brands of wines, differentiation and increased flavor diversity will be obtained with the application of increased yeast diversity during the fermentation process. Non-*Saccharomyces* species of yeast have been reported as beneficial for winemaking because they contribute to the sensory complexity of wines (3, 4). The main non-*Saccharomyces* genus associated with grapes is *Hanseniaspora*. Among the species comprising this genus, *Hanseniaspora vineae* is one of the most promising, with high oenological potential (5). Recently, *H. vineae* demonstrated the ability to provide increased levels of acetate esters and benzenoids and decreased levels of higher alcohols (except benzyl and 2-phenylethyl alcohols) in wines by pure fermentation or by cofermentation with *S. cerevisiae* (6–10). An aroma sensory analysis of wines, attributed to *H. vineae* winemaking, indicated a significant increase in fruit intensity, described as banana, pear, apple, citric fruits, and guava (9). These favorable characteristics for the winemaking industry have turned *H. vineae* into a species increasingly regarded as a means to improve aroma quality (5). Flavor diversity, including subtle characteristic differences in fine wines, has been described for other non-*Saccharomyces* species such as *Pichia*, *Metschnikowia*, and *Torulaspota* (2, 4, 11). Various genes have been identified as contributors to higher alcohol, acetate ester, and ethyl ester biosynthesis in *S. cerevisiae*; however, other species remain uncharacterized in this regard (12).

Higher alcohol formation via the Ehrlich pathway is subdivided into three steps: transamination, decarboxylation, and reduction (Fig. 1). In transamination, the key enzymes are the branched-chain amino acid transaminases (encoded by *BAT* genes) and the aromatic amino acid aminotransferases (encoded by *ARO8* and *ARO9* genes), which catalyze the transfer of amines between amino acids and their respective α -keto acid. In the second step, the α -keto acids are decarboxylated through pyruvate decarboxylases (encoded mainly by *PDC* and *ARO10* genes) to form the respective aldehydes. Finally, the reduction from aldehydes to alcohols is carried out by alcohol dehydrogenases (encoded by *ADH* genes) and aryl-alcohol dehydrogenases (encoded by *AAD* genes). The formation of the fruity- and flowery-like aroma acetate esters is dependent on acetate and alcohols, and they are due to two alcohol acetyltransferases (AATases) encoded by *ATF* genes in *S. cerevisiae*. The biosynthesis of ethyl esters is carried out by two acyl coenzyme A (acyl-CoA)/ethanol O-acyltransferases (encoded by the *EEB1* and *EHT1* genes) and involves ethanol and acyl-CoA units (derived from fatty acid synthesis). Ethyl esters as well as acetate esters contribute fruity-like aromas,

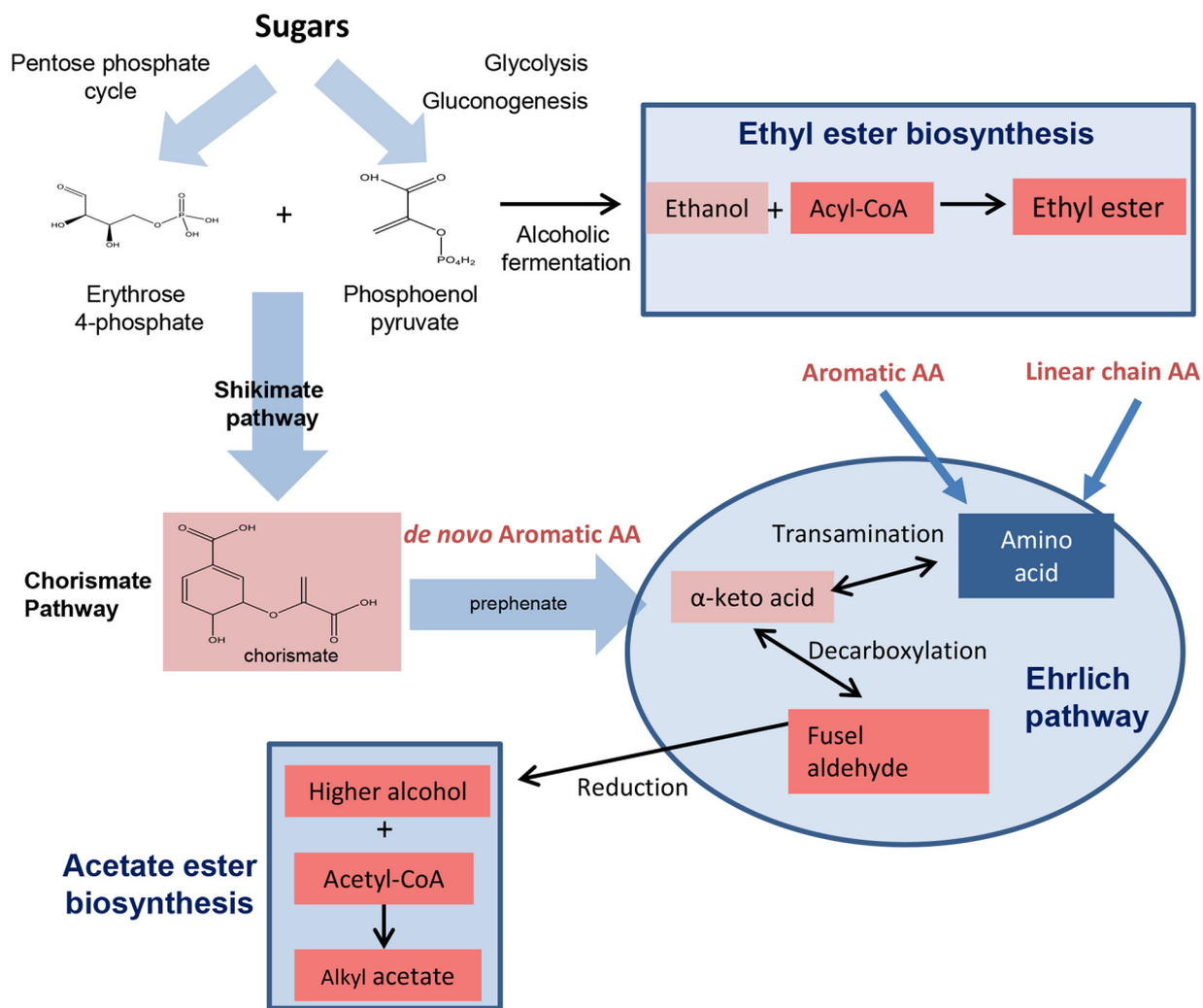


FIG 1 Metabolic pathways studied in this work involved in wine aroma formation. Ehrlich pathway for higher alcohol production, acetate ester biosynthesis, and ethyl ester biosynthesis from amino acids (AA) and sugars.

although their concentration levels in wine are significantly lower than those of the acetate esters (13–18).

In this work, genomic, transcriptomic, and metabolomic analyses of the novel and native yeast for winemaking (*H. vineae*) was conducted, and the results were compared with those of *S. cerevisiae* to understand the aroma compound differences produced. We identified several changes in the dosage of key genes involved in higher levels of alcohol, fatty acid, acetate ester, and ethyl ester biosynthesis in *H. vineae*. We analyzed the expression profiles of these genes through transcriptomics and by assessing the concentrations of several aroma compounds during three different phases of the *H. vineae* fermentation process. A comparative work that analyzed the genomic, transcriptomic, and metabolomic profiles of a member of the apiculata group of the Saccharomycodaceae yeast family is presented. An understanding of the alternative metabolic pathways of *H. vineae* compared to those of *S. cerevisiae* will contribute to an understanding of apiculata yeast biology, which is the main yeast group associated with fruits (16, 17).

RESULTS AND DISCUSSION

The yeasts analyzed in this work are shown in Table 1, and putative genes and codes related to aroma synthesis by *S. cerevisiae* are described in Data Set S1 in the supplemental material.

TABLE 1 Yeast strains analyzed in this work

Species	Strain	Ploidy	Source	BioSample ID from NCBI database ^a	Use
<i>H. vineae</i>	T02/19AF	Haploid	Uruguayan Tannat grape vines	SAMN02644989	Genomic transcriptomic and phenomic study
<i>H. vineae</i>	T02/05AF	Haploid	Uruguayan Tannat grape vines	SAMN04487210	Genomic study
<i>S. cerevisiae</i>	BY4742	Haploid	Laboratory strain, derived from S288c	SAMN03020230	FCM analysis
<i>S. cerevisiae</i>	BY4743	Diploid	Laboratory strain, derived from S288c	SAMN01822968	FCM analysis
<i>S. cerevisiae</i>	Montrachet 522	Diploid	Fortified wines, CA	SAMN03325349	Flavor compound analysis
<i>S. cerevisiae</i>	S288c	Haploid	Laboratory strain, CA	SAMD00065885	Genomic comparison
<i>S. cerevisiae</i>	AWRI1631	Haploid	Australian derivative of South African commercial wine strain N96	SAMN02953734	Genomic comparison
<i>S. cerevisiae</i>	AWRI796	Diploid	South African wine strain	SAMN04286136	Genomic comparison
<i>S. cerevisiae</i>	BC187	Diploid	Derivative of CA wine barrel isolate	SAMEA687137	Genomic comparison
<i>S. cerevisiae</i>	DBVPG6044	Diploid	West African isolate	SAMEA687132	Genomic comparison
<i>S. cerevisiae</i>	EC1118	Diploid	Commercial wine strain	SAMEA2272624	Genomic comparison
<i>S. cerevisiae</i>	L1528	Diploid	Chilean wine strain	SAMN03020223	Genomic comparison
<i>S. cerevisiae</i>	LalvinQA23	Diploid	Portuguese Vinho Verde white wine strain	SAMN02981266	Genomic comparison
<i>S. cerevisiae</i>	M22	Diploid	Italian vineyard isolate	SAMN00189351	Genomic comparison
<i>S. cerevisiae</i>	PW5	Diploid	Nigerian Raphia palm wine isolate	SAMN00199004	Genomic comparison
<i>S. cerevisiae</i>	RM11-1A	Haploid	Natural isolate collected from a vineyard, CA	SAMN02953602	Genomic comparison
<i>S. cerevisiae</i>	T73	Near-diploid	Spanish red wine strain	SAMN00198997	Genomic comparison
<i>S. cerevisiae</i>	Vin13	Diploid	South African white wine strain	SAMN02981268	Genomic comparison
<i>S. cerevisiae</i>	VL3	Diploid	French white wine strain	SAMN02981289	Genomic comparison
<i>S. cerevisiae</i>	YJM269	Diploid	Austrian wine from Blauer Portugieser grapes isolate	SAMN02981310	Genomic comparison

^aID, identifier; NCBI, National Center of Biotechnology Information.

Genome characterization of *H. vineae*. The two strains of *H. vineae* most used at the winemaking level by our group since 2009 were selected for genome sequencing: T02/19AF and T02/05AF. The sequencing of both strains was performed on an Illumina Genome Analyzer IIx platform.

The genome analysis revealed high similarity in both genomes with regard to size and the prediction of genes (Table 2; see also Table S1 and Fig. S1a). Therefore, only the data obtained from T02/19AF genomes are specified below in detail.

The sequencing run generated a mean of 13,302,566 paired-end reads (2×100 cycles). After filtering and removing redundant reads, a final set of 9,203,956 reads was used for the genome assembly. A total of 87 scaffolds with a median length of 76,832 bp were assembled through MaSuRCA software, yielding a genome (haploid) of 11.3 Mb, representing an average coverage of 163-fold, with an N_{50} of 261 kb and a GC content of 37% (Table 2; Table S1; Fig. S1a). Higher quality data and a more extensive analysis of the genome of *H. vineae* were obtained than in our previous report (19).

Genome size and ploidy level were also addressed by flow cytometry (FCM) analysis using linear plots of fluorescence intensity of cell populations stained with propidium iodide (PI). This technique discriminated two cellular subpopulations with different DNA contents, namely R1 and R2 (see Fig. S2). All tested samples presented a half-peak coefficient of variation of R1 of less than 10% (data not shown), indicating high-resolution DNA measurements. As the references for genomic DNA estimation, we used both *S. cerevisiae* haploid (BY4742) and diploid (BY4743) strains, containing genomes of 11.67 and 23.35 Mb, respectively (20). A concurrent FCM analysis of *S. cerevisiae* haploid and diploid strains revealed three distinct peaks (Fig. S2), corresponding to 1n, 2n, and

TABLE 2 Genome assembly report of the two strains of *H. vineae*

Strain	Genome size assembly (Mb)	Total no. of contigs	No. of ORFs ^a	No. of predicted proteins homologous to <i>S. cerevisiae</i>
<i>H. vineae</i> T02/05AF	11.37	741	4,741	3,862
<i>H. vineae</i> T02/19AF	11.33	305	4,708	3,861

^aORF, open reading frame.

TABLE 3 Comparison of genes involved in biosynthesis routes for key flavor compound production in *S. cerevisiae* and *H. vineae*

Biosynthesis route	Enzymatic activity	Genes identified (% amino acid identity with <i>S. cerevisiae</i> homologous protein) ^a
Higher alcohols	Aromatic amino acid transferases	3× <i>ARO8</i> (45.51, 59.84, 56.06), 4× <i>ARO9</i> (42.70, 35.27, 36.08, 36.91)
	Branched-chain amino acid transferases	<i>BAT1</i> (78.84), <i>BAT2</i>
	Decarboxylase	2× <i>ARO10</i> (34.10, 30.99), 2× <i>PDC1</i> (80.46, 50.66), <i>PDC5</i> , <i>PDC6</i> , <i>THI3</i>
	Alcohol dehydrogenase	2× <i>ADH1</i> (77.71, 78.74), <i>ADH2</i> , 2× <i>ADH3</i> (74.79, 74.80), <i>ADH4</i> , <i>ADH5</i> , 4× <i>ADH6</i> (44.74, 44.47, 44.74, 44.06), <i>ADH7</i> , <i>SFA1</i> (68.16), 4× <i>GRE2</i> (44.74, 50.73, 47.51, 43.02), <i>YPR1</i> , <i>PAD1</i> , <i>SPE1</i> , 3× <i>OYE2</i> (55.10, 58.06, 57.25), <i>HOM2</i> (78.24)
	Aryl alcohol dehydrogenase	<i>AAD3</i> , <i>AAD4</i> , <i>AAD6</i> , <i>AAD10</i> , <i>AAD14</i> , <i>AAD15</i> , <i>AAD16</i>
	Regulation	<i>ARO80</i> (34.80), <i>GAT2</i> , <i>GLN3</i> , <i>GZF3</i> , <i>DAL80</i>
Acetate esters	Alcohol acetyl transferases	<i>ATF1</i> , <i>ATF2</i> (26.58), 4× <i>SLI1</i> (22–24), g4599.t1
Ethyl esters	Ethanol <i>O</i> -acyltransferase and esterase	<i>EEB1</i> , <i>EHT1</i> (51.35), <i>MGL2</i> (30.06), <i>AAD</i> , <i>IAH1</i> (54.67)
Volatile organic acids	Aldehyde dehydrogenase	2× <i>ALD2</i> (40.55, 44.01), <i>ALD3</i> , <i>ALD4</i> , <i>ALD5</i> (53.45), <i>ALD6</i> (55.07)
Aromatic amino acid synthesis	Synthesis of chorismate, phenylalanine, tryptophan, and tyrosine	<i>ARO1</i> (66.79), <i>ARO2</i> (80.59), <i>ARO3</i> (77.03), <i>ARO4</i> (83.51), <i>TRP2</i> (70.84), <i>TRP3</i> (69.14), <i>ARO7</i> (67.97), <i>PHA2</i> (41.99), <i>TYR1</i> (62.37)
Benzyl alcohol/benzaldehyde synthesis	Mandelate pathway	2× <i>ARO10</i> (34.10, 30.99), 2× <i>PDC1</i> (80.46, 50.66), <i>SCS7</i> (66.50), <i>ALD6</i> (55.07), 2× <i>ALD2</i> (40.55, 44.01), <i>DLD1</i> (53.00), <i>DLD2</i> (70.00), <i>DLD3</i>

^aHomologous genes in *H. vineae* are not underlined, and the copy numbers (e.g., 2×) are indicated as prefixes for repeated genes. Predicted amino acid sequences from the genome of *H. vineae* were compared with protein homologs found in *S. cerevisiae*. Underlined genes represent absent homologous genes in *H. vineae*.

4n DNA contents, where the mean PI fluorescence intensity of each peak was directly correlated ($r^2 > 0.999$) to the amount of DNA of its corresponding cell subpopulation (Fig. S2). The genome size of each *H. vineae* strain was estimated in accordance with the R1 cell subpopulation (Fig. S2). The analysis by FCM revealed a diploid genome size of 16.71 ± 0.79 Mb (Table S1). Regarding gene copy number, we expected that the *H. vineae* genome would show a certain (but unknown) level of ploidy given its sporulation capacity (21). In any case, diploidy of both strains was confirmed. However, the slight difference in genome size obtained by FCM and our genome assembly-based calculations might be explained by the principles of the technique. *H. vineae* genome size was estimated using *S. cerevisiae* as the control strain. Because the cells themselves can act as a lens, changes in cell size or shape can affect the PI fluorescence detected by FCM (22), resulting in differences in genome size estimations obtained by FCM versus sequencing.

A total of 4,708 gene models were predicted using Augustus software, of which 3,855 had at least one Pfam domain of the Pfam platform databases. We identified 3,861 sequences homologous to *S. cerevisiae* genes and more than 4,141 sequences aligned to the National Center for Biotechnology Information (NCBI) nonredundant protein database (Table 2; Table S1). Due to the presence of a different number of homologous genes than reported for *S. cerevisiae* strain S288c in *H. vineae*, an Augustus prediction number (gXXXX.t1) is provided to clarify the putative gene, which was analyzed in each case.

We identified 243 of the 248 core eukaryotic genes (CEGs) and 445 of the 458 CEGs from the Augustus predictions, showing that our genome is ~98% complete. Interestingly, the protein identity between *H. vineae* and *S. cerevisiae* is only 52%, demonstrating a great divergence between these two species. Moreover, a high heterozygosity level was evidenced by single-nucleotide polymorphism (SNP) analysis using different *S. cerevisiae* strains (23). A total of 56,662 SNPs (1 heterozygous SNP per 200 bp) were found, of which, 30,740 SNPs (54%) were present in coding sequences (Fig. S1b). According to the high genetic similarity found between T02/19AF and T02/05AF, the nucleotide diversity between both *H. vineae* strains was 1 variant per 179 bp (63,021 SNPs), a similar rate to those found among different *S. cerevisiae* strains (24, 25).

The genes related to yeast aroma compound synthesis in *H. vineae* were compared with those reported for *S. cerevisiae* (see Tables 3 and S2). Absent homologies and repeated genes were found.

***H. vineae* diverged before the WGD clade of the *Saccharomyces* complex.** To determine the phylogenetic position of *H. vineae*, a phylogenetic tree was inferred by

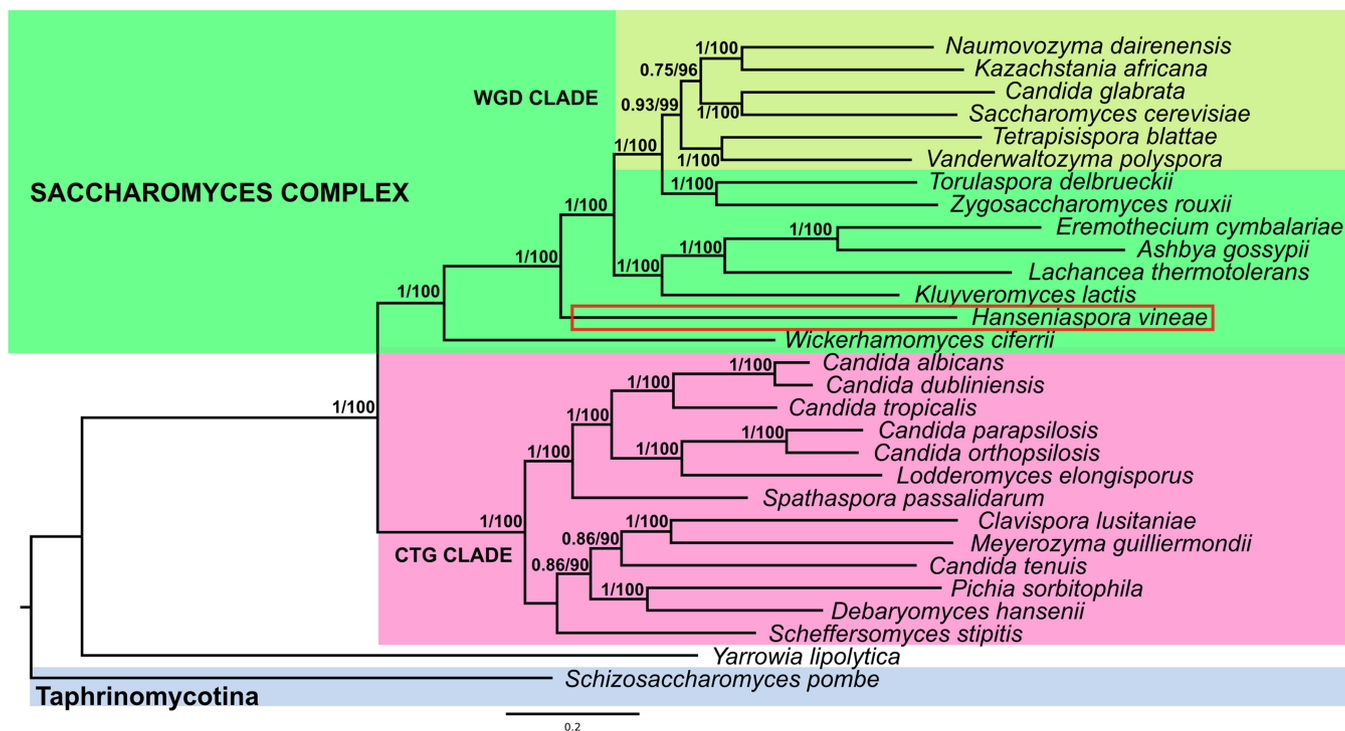
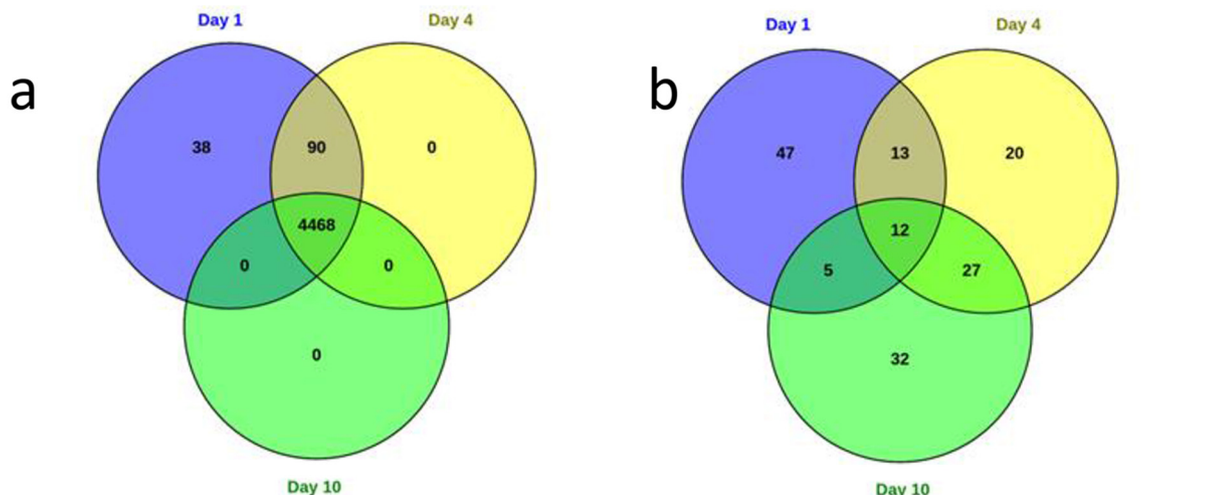


FIG 2 Maximum likelihood phylogeny of *Saccharomyces* complex species from concatenation of 227 genes. *H. vineae* is framed in red inside the *Saccharomyces* complex and outside the whole-genome duplication (WGD) clade. The clade CTG groups yeasts with alternative genetic codes. Numbers close to the node match bootstrap support (BS) for those values above 70 and internode certainty (IC). The scale bar represents units of amino acid substitutions per site. The tree has a midpoint root for easier visualization.

concatenating 227 genes from 29 species. The proteins were selected by an orthologous alignment of the predicted proteins from the *H. vineae* genome compared with those from yeast species obtained from databases. The maximum likelihood phylogeny classified *H. vineae* as part of the *Saccharomyces* complex but out of the whole-genome duplication (WGD) clade with very high support (Fig. 2). *H. vineae* was recovered as the sister taxa to two lineages, one composed of the WGD yeasts *Kazachstania africana*, *Naumovozyma dairenensis*, *Saccharomyces cerevisiae*, *Candida glabrata*, and *Tetrapisispora blattae* and the other composed of species diverged before the WGD, including *Ashbya (Eremothecium) gossypii*, *Eremothecium cymbalariae*, *Lachancea thermotolerans*, and *Kluyveromyces lactis*. Node support for this placement of *H. vineae* was very high (internode certainty [IC] = 1.0 and bootstrap support [BS] = 100).

A total of 372 orthologous groups were expanded in *S. cerevisiae* compared to *H. vineae*, which involved 427 genes. These genes have a 2:1 relationship between *S. cerevisiae* and *H. vineae*, supporting the theory that *H. vineae* diverged previously to the WGD and arose out of the fungal CTG clade formed by yeasts that present differences in their genetic codes (26). Although this phylogeny presents some differences to that previously reported for the *Saccharomyces* complex (27, 28), the phylogenetic position of *H. vineae* presents a high node support and is similar to that obtained by Kurtzman and Robnett (27). The phylogenies inferred by these authors were based on divergence in genes of the ribosomal DNA (rDNA) repeat (18S, 26S, ITS), single copy nuclear genes (translation elongation factor 1 α , actin-1, RNA polymerase II), and mitochondrially encoded genes (small-subunit rDNA, cytochrome oxidase II).

Overview of transcriptome dynamics during fermentation. To perform a comprehensive analysis, we obtained transcriptomic profiles of *H. vineae* strain T02/19AF along three different days of the fermentation process (see Fig. S3): exponential growth phase (day 1), end of exponential phase (day 4), and end of stationary phase (day 10). Fermentations were carried out in triplicates using 25-ml Erlenmeyer flasks with 125 ml



C Main biochemical cascades of the more expressed genes at each of the sampled days of fermentation

Day 1	Day 4	Day 10
-Biosynthesis of amino acids -Glycolysis/Gluconeogenesis -Methane metabolism -Pentose phosphate pathway	-Steroid biosynthesis -Arginine biosynthesis	-Protein processing in endoplasmic reticulum -GPI anchor biosynthesis
-Pyruvate metabolism -Synthesis of secondary metabolism -Oxidative phosphorylation -Citrate cycle TCA -Glycine, serine and threonine metabolism		

FIG 3 Overview of transcriptomic analysis. (a) Venn diagram showing the differentially expressed genes shared between each fermentation point. (b) Venn diagram showing the genes shared between each fermentation point for the top 100 most highly expressed genes. (c) Main biochemical cascades of the most expressed genes at each sampled day of fermentation. GPI, glycosylphosphatidylinositol.

of chemically defined grape (CDG) medium that presents a similar nutrient composition to grape juice. The medium was supplemented with 100 mg N/liter yeast available nitrogen (YAN), 200 g/liter of an equimolar mixture of glucose, and fructose, and the pH was adjusted to 3.5.

These data were analyzed to compare the expression of the key genes related to the flavor compounds present in *H. vineae*, and, moreover, to those extra copies identified exclusively in *H. vineae* and not in *S. cerevisiae*. Transcriptome sequencing of nine libraries was performed in three replicates for the three fermentation points with Illumina. Similar quantities of genes were expressed at the three fermentation points, although their expression levels differed considerably. More than 2,500 (~56%) genes were differentially expressed according to the false discovery rate (FDR) calculation (FDR < 0.05) between each pair of fermentation time points.

Transcriptome assembly enabled the identification of 15 more genes than those obtained by genome analysis, and almost all paralogous genes identified within the genome were confirmed. The transcriptome analyses for days 1, 4, and 10 presented 4,596, 4,558 and 4,468 expressed genes, respectively, of which 4,468 were in common (Fig. 3a and b). The most significant gene ontology (GO) terms associated with the genes shared between the three fermentation points were tRNA processing for biological processes, GTPase regulator activity for molecular function, and Golgi apparatus for cellular components.

High number of differentially expressed genes in *H. vineae* during fermentation. For the three fermentation points (1, 4, and 10 days), the differentially expressed genes were analyzed using edgeR software. Important changes in gene expression were detected between any pair of the three fermentation points, while the differences between replicates were minimal (see Fig. S4).

In *H. vineae*, the large number of differentially expressed genes (DEGs) identified along the fermentation process was remarkable. Of the 4,468 genes shared among the 3 days, more than 2,500 (~56%) were differentially expressed between each point (FDR < 0.05). However, microarray studies of various *S. cerevisiae* strains reported a smaller number of DEGs, ranging from 1,000 to 1,500 genes (29). The largest number of DEGs was identified between the first and last point; on the other hand, the fewest numbers were detected between day 4 and day 10. This situation is consistent with the fact that day 4 is the start of stationary phase and at day 10 the stationary phase is ending. As the fermentation process approaches stationary phase, fewer genes are expected to be differentially expressed.

Unique and expanded orthologous groups in *H. vineae* compared to those in *S. cerevisiae*. Using OrthoMCL software, 85 expanded orthologous groups were detected in *H. vineae* compared to data from the *S. cerevisiae* S288c sequence.

There was consistently higher expression at days 1 and 4 of genes related to growth biochemical cascades (such as amino acid biosynthesis, the pentose phosphate pathway, oxidative phosphorylation, and tricarboxylic acid [TCA]) and glycolysis (such as pyruvate metabolism and the synthesis of secondary compounds) (Fig. 3c; Table S3). However, at day 10, the protein turnover genes were expressed the most, as at the middle and end of fermentation, amino acids are generally exhausted from the medium. The expression of genes related to protein processing at the end of stationary phase might be related to autophagy processes. Autophagy in yeast is a response to nutrient limitation, and the endoplasmic reticulum and glycosylphosphatidylinositol (GPI) anchor mechanisms are activated under this stress situation for the recovery process of proteins (30, 31). Interestingly, methane metabolism genes were mainly expressed at exponential growth to early stationary phase (days 1 to 4), but this might be specific to *Hanseniaspora* yeasts, as they are a methylotrophic group that may be active when oxygen is present at the beginning of fermentation (32).

The most complete KEGG (Kyoto Encyclopedia of Genes and Genomes) Pathways were those related to tyrosine and phenylalanine metabolism, both aromatic amino acids that are related to phenylpropanoid synthesis (see Table S4a). The main genes that are exclusive to *H. vineae* belong to the following KEGG modules: β -lactam resistance and lysine biosynthesis, with five genes, and bacterial proteasome and benzoate degradation, with three genes (Table S4b). Five serine endopeptidases that might be involved in diverse functions were related to the β -lactam resistance module, while for lysine biosynthesis, two aldehyde dehydrogenases (g3618.t1 and g3619.t1), an unknown (4147.t1), one mlo2-like protein (g2280.t1), and ssm4 (g570.t1) proteins were found.

Genomics and yeast flavors. Several genetic and phenomic characteristics were analyzed to compare *H. vineae* and *S. cerevisiae* strains. The comparative genomics analysis included *H. vineae* and up to 14 wine industry strains of *S. cerevisiae* whose genomes were analyzed in previous studies (33, 34) (see Table S5). The aroma compound profiles determined by gas chromatography-mass spectrometry (GC-MS) of *H. vineae* were compared with those of *S. cerevisiae* strain M522 and are shown in Table 4. On the other hand, in Table 5., the aroma compounds produced by *H. vineae* at days 4 and 10 are shown, and the differential expression of genes involved in higher alcohol, acetate ester, and ethyl ester metabolism were evaluated (see Fig. S5). These results are discussed in the following sections.

Alcohols and 2-phenylethanol. The aroma compound analysis showed that overall, alcohol production was more than twice as high in *S. cerevisiae* M522 than in *H. vineae* (Fig. 4a; Table 5). In fact, other studies comparing *H. vineae* to the wine yeast *S. cerevisiae* EC1118 have found similar results (6). However, the proportion of 2-phenylethanol in *H. vineae* with respect to *S. cerevisiae* M522 is approximately equivalent (Fig. 4a) if 2-phenylethyl acetate is taken into account as a derived compound of 2-phenylethanol.

TABLE 4 Exometabolome of *H. vineae* flavor compounds produced at days 4 and 10

Compound	LRI ^a	Average \pm SD content (g/liter) ^b	
		Day 4	Day 10
Alcohols			
2-Methyl-2-butanol	975	66 \pm 5	42 \pm 2
1-Propanol	996	116 \pm 5	40 \pm 5
2-Methyl-1-propanol	1,067	3,620 \pm 268	2,990 \pm 290
1-Butanol	1,128	149 \pm 51	122 \pm 10
3-Methyl-1-butanol	1,187	42,525 \pm 1,288	36,859 \pm 1,693
2,3-Butanediol	1,526	1,310 \pm 74	1,450 \pm 252
3-Ethoxy-1-propanol	1,389	13 \pm 13	177 \pm 8
2-Ethyl-1-hexanol	1,453	ND ^c	39 \pm 2
Methionol	1,716	1,605 \pm 60	1,925 \pm 60
3-Acethoxy-1-propanol	1,756	1,335 \pm 109	1,520 \pm 50
Benzyl alcohol	1,822	280 \pm 9	407 \pm 33
2-Phenylethanol	1,906	6,657 \pm 317	7,587 \pm 361
Tyrosol	3,012	33 \pm 33	2,213 \pm 638
Esters			
3-Methylbutyl acetate	1,126	91 \pm 21	112 \pm 33
Ethyl lactate	1,341	ND	62 \pm 3
Ethyl 2-hydroxyhexanoate	1,650	ND	20 \pm 10
Benzyl acetate	1,690	ND	10 \pm 1
2-Phenylethyl acetate	1,813	5,862 \pm 627	10,260 \pm 995
Ethyl 4-hydroxy-butoanoate	1,819	ND	1,344 \pm 47
Diethyl 2 hydroxy glutarate	2,202	ND	10 \pm 2
Fatty acids			
2-Methylpropanoic acid	1,588	2,366 \pm 158	3,024 \pm 138
Butanoic acid	1,625	57 \pm 12	97 \pm 6
3-Methylbutanoic acid	1,650	71 \pm 11	128 \pm 5
Hexanoic acid	1,843	50 \pm 4	110 \pm 4
Octanoic acid	2,070	44 \pm 12	164 \pm 17
Decanoic acid	2,243	15 \pm 15	308 \pm 67
Other compounds			
2,3-Butanedione	935	407 \pm 53	58 \pm 9
2,3-Pentanedione	1,046	76 \pm 25	15 \pm 3
3-Hydroxy-2-butanone	1,270	12,691 \pm 348	9,669 \pm 275
3-Hydroxy-2-pentanone	1,330	1,353 \pm 45	1,121 \pm 184
γ -Butyrolactone	1,620	64 \pm 32	116 \pm 7
N-Formyl tyramine	2,890	727 \pm 145	8,788 \pm 451

^aLinear retention index based on a series of *n*-hydrocarbons reported according to elution order on Carbowax 20M.

^bMeans and standard deviations from triplicate fermentations at 20°C in chemically defined grape synthetic medium.

^cND, not detected.

The three steps of higher alcohol biosynthesis (transamination, decarboxylation, and reduction) (Fig. 4b) were analyzed attending to transcriptomics and phenomic results.

(i) Transamination. In *S. cerevisiae*, the most important gene involved in transamination leading to the production of higher alcohols is *BAT2* (35), which encodes the branched-chain amino acid aminotransferase. *BAT2* is absent in the *H. vineae* genome. This might explain the reduced presence of overall branched-chain higher alcohols in *H. vineae* fermentations compared to that in *S. cerevisiae* M522. In this scenario, the *BAT1* gene in *H. vineae* would perform the two reactions of the reversible transamination step. *BAT1* showed higher expression levels on day 1 and a decay in expression on days 4 and 10, while overall alcohol levels remained constant (Fig. 4d and S5). Therefore, the production of alcohols occurs early in fermentation, preceded by the expression of this gene.

On the other hand, the amounts of 2-phenylethanol/2-phenylethyl acetate remain constant between days 4 and 10, while the expression of the *ARO8* and *ARO9* genes reaches a peak by day 4 (Fig. 4c and d; Fig. S5). *S. cerevisiae* industrial strains present only one copy of these *ARO* genes (Table S5); however, *H. vineae* presents three copies

TABLE 5 Exometabolome of *H. vineae* and *S. cerevisiae* flavor compounds at the end of the fermentation

Compound	LRI ^a	Average \pm SD content (g/liter) ^b		
		<i>H. vineae</i>		<i>S. cerevisiae</i>
		T02/05AF	T02/19AF	M522
Alcohols				
2-Methyl-2-butanol	975	168 \pm 78	159 \pm 1	ND ^c
1-Propanol	996	ND	2 \pm 2	42 \pm 1
2-Methyl-1-propanol	1,067	631 \pm 490	750 \pm 22	3,488 \pm 4
1-Butanol	1,128	31 \pm 14	33 \pm 5	91 \pm 2
3-Methyl-1-butanol	1,187	25,028 \pm 3,699	28,326 \pm 954	54,953 \pm 41
2,3-Butanediol	1,526	422 \pm 68	1,076 \pm 65	ND
3-Ethoxy-1-propanol	1,389	75 \pm 17	135 \pm 6	175 \pm 1
2-Ethyl-1-hexanol	1,453	29 \pm 4	26 \pm 6	312 \pm 1
Methionol	1,716	2,032 \pm 230	2,601 \pm 170	4,980 \pm 6
Benzyl alcohol	1,822	141 \pm 25	179 \pm 8	ND
2-Phenylethanol	1,906	8,029 \pm 2,067	9,879 \pm 120	18,387 \pm 2
Tyrosol	3,012	814 \pm 188	1,006 \pm 11	7,683 \pm 4
Esters				
3-Methylbutyl acetate	1,126	33 \pm 19	20 \pm 4	54 \pm 1
Ethyl lactate	1,341	66 \pm 5	81 \pm 17	116 \pm 1
Benzyl acetate	1,690	6 \pm 0	4 \pm 0	ND
2-Phenylethyl acetate	1,813	10,054 \pm 929	9,205 \pm 1,435	1,185 \pm 6
Fatty acids				
2-Methylpropanoic acid	1,588	301 \pm 21	668 \pm 52	168 \pm 1
Butanoic acid	1,625	59 \pm 6	55 \pm 6	133 \pm 1
3-Methylbutanoic acid	1,650	67 \pm 10	146 \pm 3	448 \pm 1
Hexanoic acid	1,843	82 \pm 19	67 \pm 4	461 \pm 1
Octanoic acid	2,070	127 \pm 37	89 \pm 14	875 \pm 2
Decanoic acid	2,243	170 \pm 111	81 \pm 26	96 \pm 2
Other compounds				
3-Hydroxy-2-butanone	1,270	4,328 \pm 1,858	5,165 \pm 742	303 \pm 20
γ -Butyrolactone	1,620	90 \pm 22	153 \pm 14	338 \pm 2

^aLRI, linear retention index based on a series of *n*-hydrocarbons reported according to their elution order on Carbowax 20M.

^bMeans and standard deviations from triplicate fermentations at 20°C in chemically defined grape (CDG) synthetic medium.

^cND, not detected.

of *ARO8* and four of *ARO9* that are all very similarly expressed during fermentation (Fig. 4d; Fig. S5). *ARO8* and *ARO9* encode aromatic amino acid transaminases, which act as broad-substrate-specificity amino acid transaminases in the Ehrlich pathway (15) and they are involved in the anabolism and catabolism of the aromatic amino acids phenylalanine and tyrosine. These data are in agreement with the KEGG pathways overrepresented in *H. vineae* as shown in Table S4a. Therefore, the overexpression of these two expanded genes might explain the larger proportion of 2-phenylethanol in two ways: first, for their incremented specificity for aromatic amino acids present in the medium, and second, for an increased synthesis of phenylalanine that is known as 2-phenylethanol precursor (36).

(ii) Decarboxylation. Five genes are involved in the decarboxylation step in *S. cerevisiae* (*PDC1*, *PDC5*, *PDC6*, *ARO10*, and *THI3*) (15), of which, *H. vineae* has two copies of *ARO10* and two of *PDC1*. The most highly expressed paralogous copy of *PDC1* had an expression pattern similar to that of *BAT1* on day 1, prior to alcohol production (Fig. 4c and d; Fig. S5a).

It is possible that *ARO10* duplication (Table 3) enables an efficient decarboxylation of aromatic α -keto acids derived from the enhanced transamination step. In fact, this is supported by the expression profiles (Fig. 4d; Fig. S5a) of both *ARO10* genes that are very similar to the expression profiles found for *ARO8* and *ARO9* copies. It should be noted that the cofermentation of *H. vineae* with *S. cerevisiae* resulted in an increased

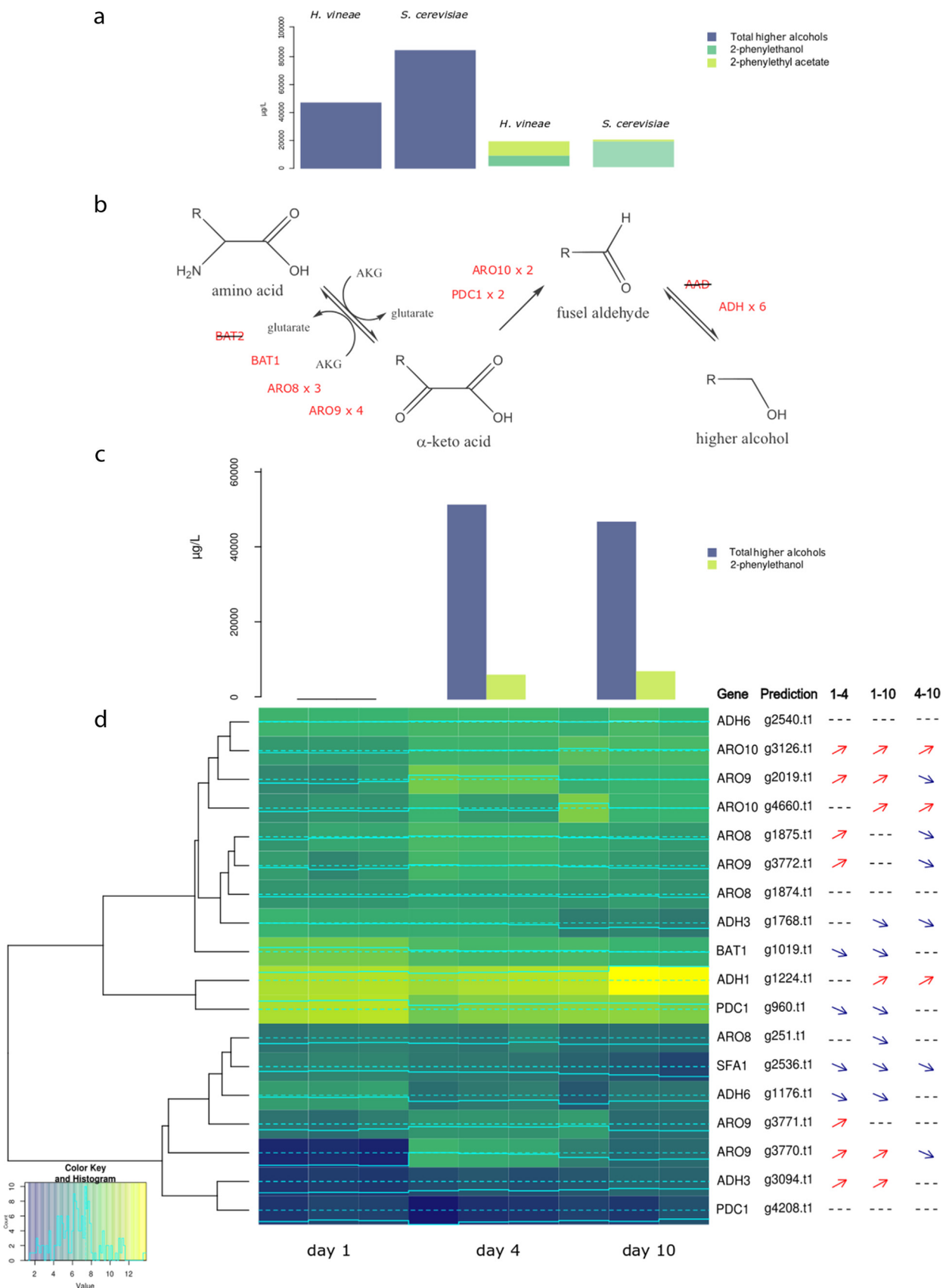


FIG 4 Higher alcohols and 2-phenylethanol production and putatively related genes. (a) Comparison of total higher alcohols, 2-phenylethanol, and 2-phenylethyl acetate produced in *H. vineae* and *S. cerevisiae* at day 10 of fermentation. (b) Three steps of metabolic pathway of higher alcohols (Continued on next page)

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intensity of citrusy aromas of which 2-phenylethanol (and therefore the *ARO* gene duplications) might be responsible (9). All 14 *S. cerevisiae* industrial strains only showed one copy of *ARO10* (Table S5). Further, *ARO10* has been shown to be related to the production of benzyl alcohol in a putative metabolic pathway of mandelate (7). Therefore, this decarboxylase activity might be involved in the enhanced (more than two orders of magnitude) synthesis of benzylic alcohol in *H. vineae* compared to that in *S. cerevisiae* (Table 5).

(iii) Reduction to higher alcohol. Surprisingly, *H. vineae* did not contain homologous sequences or any transcriptional evidence of the seven aryl alcohol dehydrogenases (*AAD* genes) present in the *S. cerevisiae* S288c sequenced genome (Table 3). This activity catalyzes the chemical reaction between aromatic aldehydes and alcohols. Given the overproduction of benzyl and 2-phenylethyl alcohol (precursor of the 2-phenylethyl acetate) in *H. vineae* compared to that in *S. cerevisiae* M522 (Table 5), at least one aryl alcohol dehydrogenase protein would be expected. However, it should be noted that the final step of the Ehrlich pathway (higher alcohol formation) can be catalyzed by any one of the ethanol dehydrogenases (*Adh1*, *Adh2*, *Adh3*, *Adh4*, and *Adh5*) or by *Sfa1* (a formaldehyde dehydrogenase) in *S. cerevisiae* (37). In *S. cerevisiae*, the alcohol dehydrogenases are present in two multigenic families, with four genes each (according to Ensembl): *ADH6*, *ADH7*, *YAL061W*, and *YAL060W* in one family and *ADH1*, *ADH2*, *ADH3*, and *ADH5* in the other. *H. vineae* presents two copies of *ADH1* and *ADH3* and four of *ADH6*, totaling eight genes, as in *S. cerevisiae* (Table 3). The *ADH4* gene does not belong to either of these multigenic families and is absent in *H. vineae*. In *H. vineae*, not all paralogous copies of *ADH* genes showed significant transcriptional activity (many paralogous copies were assembled but they were filtered out before differential expression analysis). Interestingly, two of the four paralogous copies of *ADH6* found in *H. vineae* were not expressed under these conditions (Fig. S5a).

In regard to the expression levels, four of the other five alcohol dehydrogenase genes, as well as *SFA1*, were significantly more expressed on days 1 and 4, while just one copy of *ADH1* and *ADH3* were more expressed on days 4 and 10 (Fig. 4d; Fig. S5a). One of the *ADH6* copies showed a significant decline in expression levels between days 1 and 4, which is consistent with that previously reported for *S. cerevisiae* (29). In contrast, one *ADH3* gene copy showed a 2-fold increase in expression by day 4 relative to day 1, similar to that for the *AAD10* and *AAD14* genes during *S. cerevisiae* wine fermentation (29). As a result, we suggest that the *ADH* genes that may be replacing the *AAD* genes might be those that show the same expression profile found in *S. cerevisiae*. Further biochemical studies will be necessary to confirm this suggestion.

Acetate esters. *H. vineae* and *S. cerevisiae* M522 also showed notable differences in overall acetate production, whereby *H. vineae* produced concentrations one order of magnitude higher than *S. cerevisiae* (Fig. 4a). As mentioned, *H. vineae* also showed a larger turnover from 2-phenylethanol to 2-phenylethyl acetate than *S. cerevisiae*. For example, 2-phenylethyl acetate only constituted a small fraction in *S. cerevisiae* of the total 2-phenylethanol produced compared to that in *H. vineae* (Fig. 4a; Table 5).

With regard to the genes involved in acetate ester formation, the *H. vineae* genome presented a highly divergent putative ortholog of the *S. cerevisiae* *ATF2* gene, and it did not present any sequences homologous to *ATF1*. However, there were also five predictions containing the AATase Pfam domain. The four *SLI1* *N*-acetyltransferase homologues are repeated in tandem in the *H. vineae* genome (one of them is out of the transcriptomic analysis according to threshold evaluation) (Fig. 5d). Three of these genes that were highly expressed in *H. vineae* (Fig. 5d) have weak similarity (22% to

FIG 4 Legend (Continued)

biosynthesis with putative enzymes involved in *H. vineae*. (c) Production of total higher alcohols and 2-phenylacetate by *H. vineae* at 1, 4, and 10 days of fermentation. (d) Expression heatmap of genes putatively involved in higher alcohols and 2-phenylethanol production from *H. vineae* at 1, 4, and 10 days of fermentation. Lighter colors indicate higher expression values, and data are shown for triplicates. Significant changes in expression of each gene are indicated with arrows to the right of the heatmap as analyzed using the package edgeR (FDR < 0.05). 1–4, differential expression between days 1 and 4; 1–10, differential expression between days 1 and 10; 4–10, differential expression between days 4 and 10.

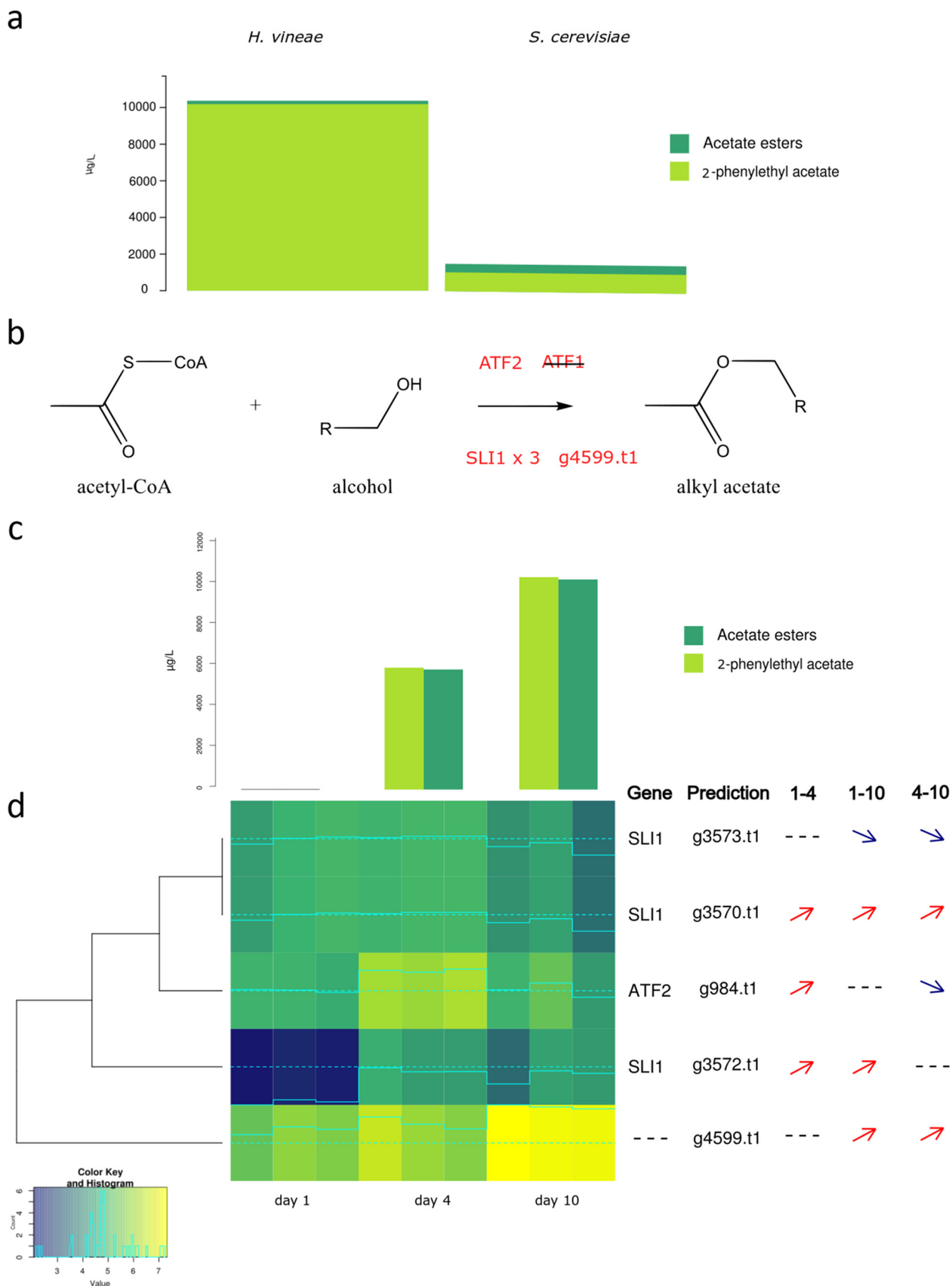


FIG 5 Acetate ester production and putatively related genes. (a) Comparison of total acetate esters and 2-phenylethyl acetate produced in *H. vineae* and *S. cerevisiae* at day 10 of fermentation. (b) Metabolic pathway of acetate esters biosynthesis with putative enzymes involved in *H. vineae*. (c) (Continued on next page)

24% at the amino acid level) with *S. cerevisiae* *SLI1*, which is a unique copy gene encoding *N*-acetyltransferase activity. It is known that *SLI1* has wide specificity for aromatic amines, similar to the *ATF* genes (38). The other *H. vineae* AATase predicted (g4599.t1) has no homology with any *S. cerevisiae* gene previously reported; however, it is the most highly expressed gene at the end of stationary phase (Fig. 5d). The *ATF2* gene and the most highly expressed *SLI1* gene copy were both highly expressed on day 4, which explains the notable 2-fold increment of acetate esters between days 4 and 10 (Fig. 5c; Fig. S5b). Curiously, only *S. cerevisiae* strain M522 did not present the *ATF1* gene, while none of the 14 industrial *S. cerevisiae* strains showed more than one gene similar to *SLI1* (Table S5).

Therefore, the presence of six sequences with AATase domains (one *ATF2*, four for *SLI1*, and g4599.t1) might explain why *H. vineae* produces significantly more acetate esters than *S. cerevisiae*. The higher turnover of 2-phenylethanol to its corresponding acetate esters in *H. vineae* compared to that in *S. cerevisiae* clearly suggests that some of the *H. vineae* AATases (e.g., *SLI1* paralogs) might be specific for this aromatic alcohol. The increased level of acetate esters in *H. vineae* explains the more intense fruity aroma resulting from the fermentation of *H. vineae* in Chardonnay (9) and Macabeo (8) wines. In fact, other apiculate yeasts from the *Hanseniaspora* genus are higher acetate esters producers than *S. cerevisiae* (39). However, high production of 2-phenylacetate is a particular characteristic of *H. vineae* compared to other species of this genus (40). Other *Hanseniaspora* species commonly produce increased levels of ethyl acetate. It is noteworthy that with regard to information about sequenced genomes of other *Hanseniaspora* species available in databases (41, 42), most do not present *SLI1* homologous sequences. The exception is *Hanseniaspora osmophila* with two putative *SLI1* copies. The detection of six AATases in *H. vineae* provides a relevant higher number of proteins for acetate ester biosynthesis than from the three copies of *S. cerevisiae*. These variations might contribute to improved functional designs for 2-phenylethanol acetylation and the synthesis of other phenylpropanoid aroma compounds, which are scarce pathways in *S. cerevisiae* strains.

Ethyl esters. *EEB1* and *EHT1* code for ethanol *O*-acyltransferases responsible for medium-chain fatty acid ethyl ester biosynthesis in *S. cerevisiae* (43). A decrease in the production of ethyl esters was observed in *H. vineae* compared to that in *S. cerevisiae* M522 (Fig. 6; Table 5). Furthermore, the absence of one of the main genes involved in ethyl ester production (*EEB1*) in *H. vineae* is consistent with this result (Fig. 6b). Only three strains of *S. cerevisiae* did not present this gene (Table S5). *EHT1* is present in the *H. vineae* genome and it is highly and significantly expressed on days 1 and 4 relative to day 10 (Fig. 6c; Fig. S4). This might be consistent with the fact that esterified fatty acids were quantified on day 10 but were not detected on day 4 (Table 5).

Even so, an important interstrain difference in the expression of acyltransferases was found during the fermentation process in *S. cerevisiae*. In general, the expression of *EHT1* in *S. cerevisiae* increased somewhat as fermentation progressed (30), which differs from our findings in *H. vineae*. Regarding our data, ethyl ester compounds were detectable on day 10 of the fermentation process. Here, it should be noted that our results are consistent with those obtained with this species in wines of Chardonnay (9) and Macabeo (8) fermentations, in which they exhibited decreased levels of ethyl esters compared to those of acetate esters.

Conclusion. The use of non-*Saccharomyces* yeasts in winemaking is limited due to the insufficient characterization of many species that naturally participate in these processes. *H. vineae* has proved to contribute with flavor diversity in winemaking

FIG 5 Legend (Continued)

Production of total acetate esters and 2-phenylethyl acetate by *H. vineae* at 1, 4, and 10 days of fermentation. (d) Expression heatmap of genes putatively involved in total acetate esters and 2-phenylethyl acetate production from *H. vineae* at 1, 4, and 10 days of fermentation. Lighter colors indicate higher expression values, and data are shown for triplicates. Significant changes in expression of each gene are indicated with arrows to the right of the heatmap as analyzed using the package edgeR (FDR < 0.05). 1–4, differential expression between days 1 and 4; 1–10, differential expression between days 1 and 10; 4–10, differential expression between days 4 and 10.

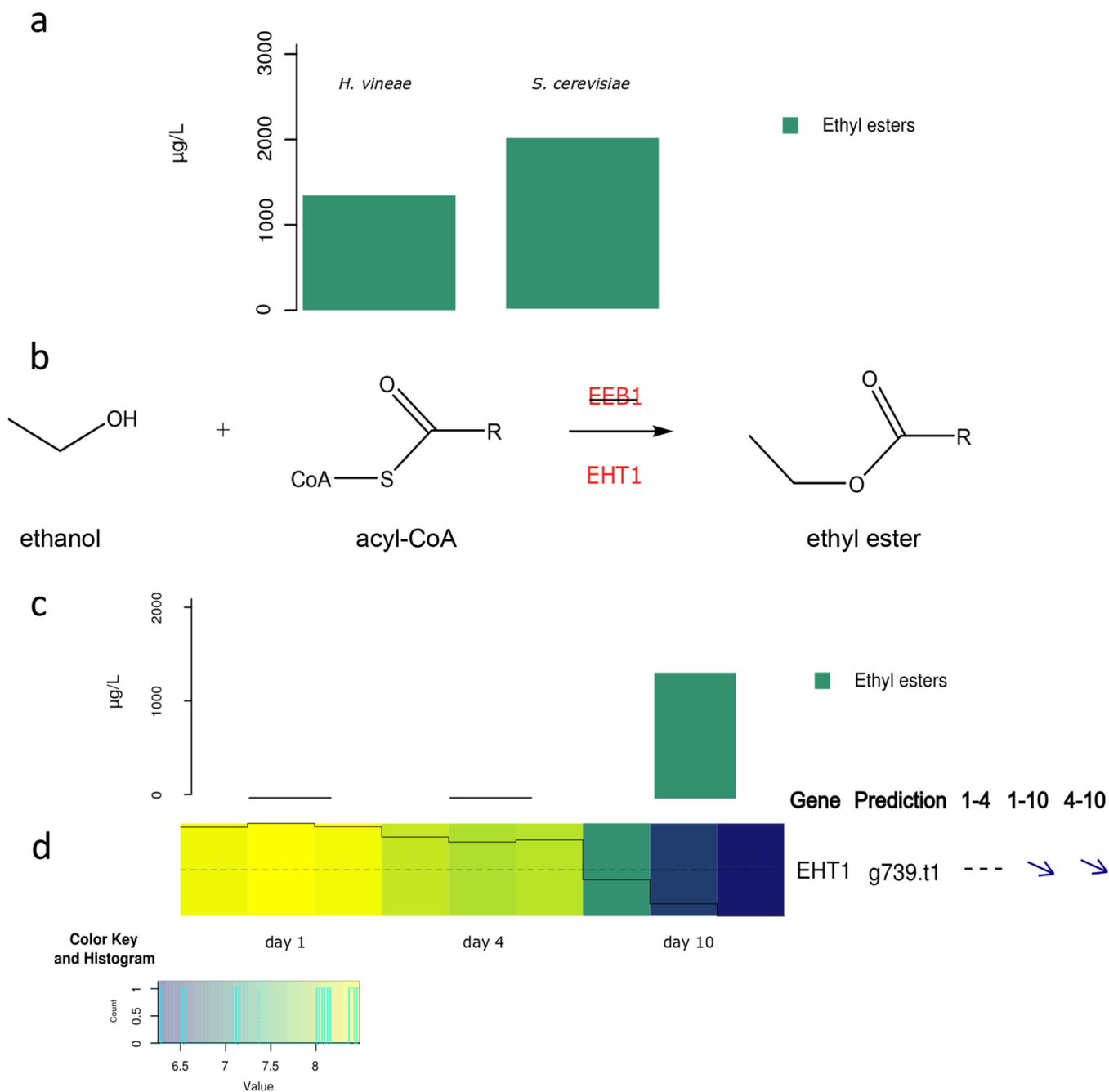


FIG 6 Ethyl esters production and putatively related genes. (a) Comparison of ethyl esters produced in *H. vineae* and *S. cerevisiae* at day 10 of fermentation. (b) Metabolic pathway of acetate ester biosynthesis with putative enzymes involved in *H. vineae*. (c) Production of ethyl esters by *H. vineae* at 1, 4, and 10 days of fermentation. (d) Expression heatmap of genes putatively involved in ethyl ester production from *H. vineae* at 1, 4, and 10 days of fermentation. Lighter colors indicate higher expression values, and data shown are of triplicates. Significant changes in expression of each gene are indicated with arrows to the right of the heatmap as analyzed using the package edgeR (FDR < 0.05). 1–4, differential expression between days 1 and 4; 1–10, differential expression between days 1 and 10; 4–10, differential expression between days 4 and 10.

conditions. Here, we present a deep genomic, transcriptomic, and metabolomic analyses and their comparisons with *Saccharomyces* strain data. On the basis of our results with a synthetic chemically defined grape juice medium, this work represents a relevant contribution to understand the biology and phylogenic relationship of the main yeast genus associated with grapes. The larger production of acetate esters, the increased ratio of 2-phenylethyl acetate to 2-phenylethanol, and the reduced amount of ethyl esters found in *H. vineae* may be due to the high presence of putative alcohol

acetyltransferase proteins and the absence of *EEB1*. These results are in agreement with previous reports studied in real winemaking conditions. As was shown, *H. vineae* produced a large amount of phenylpropanoids compared to that by *S. cerevisiae* and other yeasts, which might be explained by gene duplications and highly expressed *ARO* genes. This work established that *H. vineae* may be a potential model eukaryotic species to study benzenoid synthesis pathways, an alternative to the phenylalanine ammonia lyase (PAL) pathway commonly found in plants and Basidiomycetes. These phenolic volatile compounds have several known key functions in plants, such as cell-cell communication, antimicrobial activity, or phytohormone production, that make them highly attractive to the yeast biotechnology industry.

MATERIALS AND METHODS

Yeasts. Table 1 shows all the yeast strains utilized in this work.

Genomic characterization of *H. vineae*. (i) **DNA and RNA isolation from *H. vineae* strains.** *H. vineae* T02/19AF and T02/05AF strains were isolated from the Uruguayan Tannat vineyards. These strains were identified as *H. vineae* by sequencing the ribosomal D1/D2 region, and the strains were differentiated using the tandem repeats tRNA PCR technique (44). Genomic DNA was obtained from *H. vineae* cultures grown in yeast extract-peptone (YP) medium (1% yeast extract and 2% peptone, supplemented with 2% glucose) at 30°C, using the Wizard Genomic DNA purification kit (Promega, Madison, WI, USA), according to the manufacturer's instructions. Total RNA was obtained from the *H. vineae* T02/19AF strain grown under static batch fermentation conditions using the RiboPure RNA purification kit yeast (Life Technologies, Grand Island, USA). The poly(A) mRNA fraction was then isolated using the Oligotex mRNA Minikit (Qiagen, Hilden, Germany) and converted to indexed transcriptome sequencing (RNA-seq) libraries with the ScriptSeq v2 RNA-seq library preparation kit (Epicentre Biotechnologies, Madison, WI, USA).

(ii) **Genome length and ploidy estimation by flow cytometry.** *H. vineae* strains were grown in YP medium supplemented with 2% glucose, and 1×10^7 cells were pelleted at $3,000 \times g$ for 3 min and washed with ice-cold phosphate-buffered saline (PBS; 138 mM NaCl, 3 mM KCl, 8.1 mM Na_2HPO_4 , and 1.5 mM KH_2PO_4). To fix cells, 1 ml of 70% cold ethanol was slowly added, and the samples were stored at 4°C overnight. After removing the ethanol by centrifugation, the cell pellet was washed with PBS and resuspended in 700 μl of the same buffer. Each sample was sequentially treated with 250 μl of 1 mg/ml RNase A (Applichem, USA) (1 h at 50°C), 50 μl of proteinase K (20 mg/ml; Sigma-Aldrich, USA) (1 h at 50°C), and incubated overnight at 4°C in the dark with 50 μl of propidium iodide (PI, 1 mg/ml; Life Technologies, USA). The analysis of DNA content by FCM requires staining yeasts with PI, a fluorochrome that binds to DNA.

FCM analyses were performed using a CyAn ADP LX, 7-color flow cytometer (Beckman Coulter, USA). The blue laser (488 nm) was selected to excite the PI fluorophore. The fluorescence area signal was detected with a 575/25-nm (FL2) emission filter and plotted on a linear scale. Data acquisition and analysis were achieved using Summit v4.3 software (DakoCytomation, UK), and 10,000 events per sample were collected. The gating strategy comprised a forward scatter (FSC) versus side scatter (SSC) cell region that excluded cellular debris and irrelevant small particles. This region was applied to a PI histogram so that only gated events were displayed. *S. cerevisiae* strains BY4742 and BY4743 (Table 1) were used as the controls. The mean fluorescence intensity of stained cells as measured by FCM was taken as indicative of the total DNA content, and a direct correlation between fluorescence intensity measurements and the amount of DNA in each control strain was established. All cultures generated bimodal fluorescence profiles composed of two peaks: one corresponding to a population of a majority of cells in G phase (lower intensity peak) and the other (higher intensity peak) attributed to cells in S phase undergoing DNA synthesis. The genome size of each *H. vineae* strain was estimated in accordance with the mean fluorescence of the peak subpopulation that showed lower intensity values. Three independent biological experiments were performed, and samples were analyzed in triplicates for each experiment.

(iii) **Genome assembly and gene annotation.** Genomes were sequenced using an Illumina Genome Analyzer IIx platform in paired-end mode. A shotgun genomic library was generated on the basis of standard methods.

The reads were filtered and trimmed with the QC Toolkit (45). The first 15 bases at the 5' end and the last bases of the 3' end with a Phred quality smaller than 30 were trimmed. The reads with average Phred quality scores smaller than 20 were filtered.

Digital normalization to the paired reads was applied to systematize the coverage, from uneven $200 \times$ to $30 \times$ across the genome, to gain computation efficiency and to eliminate most of the erroneous kmer (46, 47). The *de novo* genome assembly was performed using MaSuRCA (48) (insert length, 900). To reduce heterozygosity redundancy and find any potential gene tandem repeats, HaploMerger (49) was applied using default parameters.

Gene prediction was carried out using Augustus (50) trained with *S. cerevisiae* gene models. Peptide predictions were then annotated through BLASTp (cutoff for E value, $1e-10$) against *S. cerevisiae* proteins obtained from the *Saccharomyces* genome database (20). The Pfam protein families database (51) was used to predict possible protein domains. To evaluate genome completeness, core eukaryotic genes (CEGs) (52) were sought with BLASTp (cutoff for E value, $1e-10$). Gene ontology analysis was carried out using topGO (53).

TABLE 6 Transcriptomic assembly reference metrics for *H. vineae* T02/19AF

Parameter	Transcriptomic reference
Total length (bp)	9,362,444
Total contig number	4,725
Contig length (bp)	
Maximum	17,336
Minimum	226
Mean	1,982
Median	1,683
No. of genes annotated	4,725

(iv) SNP identification. Genomic short reads sequences were mapped to the assembled genome of T02/19AF using Bowtie2 in paired-end mode with default conditions (54) and processed using SAMtools (55) and Picard (<http://broadinstitute.github.io/picard/>). Through the GATK pipeline (56, 57), SNPs were identified using Unified Genotyper applying hard filter (QD < 2.0, FS > 60.0, MQ < 40.0, HaplotypeScore > 13.0, MappingQualityRankSum < -12.5, ReadPosRankSum < -8.0). Base pair coverage was calculated using BEDTools (58). The reads of *H. vineae* T02/05AF were aligned to those of T02/19AF to estimate the nucleotide divergence between these two strains.

Analysis of 14 *S. cerevisiae* industrial wine strains. For several genes with known functions in the biosynthesis of acetate esters, ethyl esters, and higher alcohols, we determined which ones were present, duplicated, or absent in the *H. vineae* genome compared to *S. cerevisiae* S288c and an additional 14 *S. cerevisiae* wine strains. These strains were selectively chosen because they are used in wine fermentation and commercial winemaking studies (Table 1).

Ortholog cluster analysis. The proteomes of 31 fungal species were downloaded from OrthoDB (59). This web service has the orthologous relationships among a broad group of predefined species. For orthologous identification, we first used pairwise BLASTp against *H. vineae* and selected the reciprocal best hit. Then, we compared our orthologous group with those present on the OrthoDB database, and if they contained at least one gene not belonging to the corresponding OrthoDB group, they were filtered out. The protein alignment was conducted with MUSCLE v3.8.31 (60). We used PAL2NAL (61) for aligning the nucleotides on the basis of the protein alignment and Gblocks v0.91b (62) to eliminate poorly aligned positions. We finally obtained 227 proteins for 29 species (two species had to be discarded because we could not find the correspondence between their protein and nucleotide sequences) to recover the phylogenetic position of *H. vineae*.

To establish orthologous clusters between *S. cerevisiae* S288c and *H. vineae* T02/19AF, the predicted proteins were analyzed with the OrthoMCL web server (63). Orthologous clusters were classified as expanded in *H. vineae* if the number of *H. vineae* genes in one OrthoMCL group was larger than the number of *S. cerevisiae* genes present in that group. To identify the pathways involved in each group, *S. cerevisiae* genes were used as input on the DAVID functional annotation pipeline (64). Those orthologous cluster groups exclusive to *H. vineae* (not containing any *S. cerevisiae* sequences) were analyzed using the EC enzymes and KEGG modules of the corresponding orthologous group (65) using custom Python scripts.

Phylogenetic analysis. A supermatrix tree was constructed using a set of 227 genes from 29 species, including *H. vineae*. First, FASconCAT (62) was used to concatenate the supermatrix of 214,302 bases. The problematic aligned regions were previously removed with Gblocks v0.91b (66). For this supermatrix, the best partition scheme was chosen through PartitionFinder (67). The phylogenetic inference under maximum likelihood was performed with RAxML employing a GTRCAT substitution model for each of the 32 partitions suggested by PartitionFinder and using 200 starting trees. Node support was summarized in RAxML. Bootstrap support (BS) was calculated using extended majority-rule consensus for the bootstrapped trees set. Support is also shown as internode certainty (IC) values, a recently developed metric that considers the frequency of the bipartition defined by the internode in a given set of trees jointly with that of the most prevalent conflicting bipartition in the same tree set (68).

Transcriptome analyses. Nine transcriptomes, three replicates from three different fermentation stages, at days 1, 4, and 10, were analyzed. Paired-end transcriptome sequencing was performed using Illumina MiSeq. High-quality raw sequencing reads were directly assembled using Trinity (47). They yielded a total of 7.8 Gb of data and 52 million 75-bp paired reads. The transcriptomic reference constructed resulted in 4,725 contigs with an average and median length of 1,982 and 1,683 bp, respectively (Table 6).

A transcriptomic reference was constructed using the transcriptome of each sample, and an assembly was constructed by joining all of the reads for the subsequent gene expression analysis. For the construction of the transcriptomic reference, we selected the best reciprocal hit between the contigs among the 10 assembled transcriptomes and the subject sequences (19). The subject sequences were constructed using *H. vineae* T02/19AF protein predictions and *S. cerevisiae* proteins from the OMA browser (69). The alignments were carried out using reciprocal BLASTx (E value cutoff, $1e-10$).

The reads were aligned against the transcriptomic reference implementing RSEM (default settings) (70). The obtained expected counts for each gene were then used for the differential gene expression analysis carried out with edgeR (71). Genes with cpm of <5 in 2 samples or more per each fermentation

point were removed from the differential expression analysis. Genes with an FDR of <0.05 were considered differentially expressed.

Aroma compound analysis in a synthetic medium. (i) Fermentation conditions. Chemically defined grape (CDG) fermentation medium (simulating the nutrient components of grape juice but devoid of grape precursors) was prepared with the same composition to study the *de novo* formation of aroma compounds and for the transcriptome analysis with a previously described process (72) with some variations. The modifications were as follows: the total nitrogen content was adjusted to a basic amount of 50 mg of nitrogen (N)/liter with each amino acid and ammonium component added in the same proportions as indicated previously (72). The final CDG medium used for inoculum preparation and fermentations was made by increasing the basic concentration by supplementing with diammonium phosphate (DAP) up to a yeast available nitrogen (YAN) concentration of 100 mg N/liter. This YAN amount was not a limiting concentration for the complete fermentation of sugars by the yeast strains used. The final pH of the medium was adjusted to 3.5 with HCl. Equimolar concentrations of glucose and fructose were added to reach a total of 200 g/liter, and the mixed vitamins and salts were as described previously (72). Tween 80 was excluded from the medium, because it was not found to be necessary for complete fermentation and it had a negative impact on the sensory characteristics of the resultant wines. Ergosterol was added as the only supplemented lipid at a final concentration of 10 mg/liter.

Inocula were prepared in 10 ml of the same CDG medium by incubating for 12 h in a rotary shaker at 150 rpm and 25°C. Fermentations were carried out in 125 ml of medium contained in 250-ml Erlenmeyer flasks closed with cotton plugs to simulate microaerobic conditions (73). The inoculum size was 1×10^5 cells/ml in the final medium for all strains. Static batch fermentations were conducted at 20°C in triplicates, simulating winemaking conditions. Wine samples for GC analysis were taken at days 4 and 10 during fermentation and at the end of the process. The samples were filtered through 0.45- μ m pore membranes; SO₂ was added as 50 mg/liter of sodium metabisulfite.

(ii) Aroma volatile compounds. The extraction of aroma compounds was performed using adsorption and separate elution from an Isolute ENV1 cartridge (IST Ltd., Mid Glamorgan, UK) packed with 1 g of a highly cross-linked styrene-divinylbenzene (SDVB) polymer. The treatment of samples and GC-MS analysis were performed as described previously (4) in a Shimadzu-QP 2010 ULTRA (Tokyo, Japan) mass spectrometer equipped with a Stabilwax (30 m by 0.25 mm inside diameter [i.d.], 0.25- μ m film thickness; Restek) capillary column.

(iii) Identification and quantification. The components of wine aromas were identified by comparing their linear retention indices with pure standards (Aldrich, Milwaukee, WI). A comparison of mass spectral fragmentation patterns with those stored in databases was also performed. GC-flame ionization detection (GC-FID) and GC-MS instrumental procedures using an internal standard (1-heptanol) were applied for quantitative purposes, as described previously (4). All fermentations and chemical analysis were performed in triplicates. Analyses of variance (ANOVAs) were conducted to determine the differences in aroma compound concentrations among the strains with Statistica 7.0 (StatSoft Inc., Tulsa, OK, USA).

Accession number(s). This whole-genome shotgun project has been deposited in DDBJ/EMBL/GenBank under the accession number [JFAV00000000](https://doi.org/10.1101/2019.08.28.281111). The version described in this paper is [JFAV03000000](https://doi.org/10.1101/2019.08.28.281111).

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at <https://doi.org/10.1128/AEM.01959-18>.

SUPPLEMENTAL FILE 1, PDF file, 0.9 MB.

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