Hanseniaspora vineae and the Concept of Friendly Yeasts to Increase Autochthonous Wine Flavor Diversity

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In this perspective, we will explain the concept of “friendly” yeasts for developing wine starters that do not suppress desirable native microbial flora at the initial steps of fermentation, as what usually happens with Saccharomyces strains. Some non-Saccharomyces strains might allow the development of yeast consortia with the native terroir microflora of grapes and its region. The positive contribution of non-Saccharomyces yeasts was underestimated for decades. Avoiding them as spoilage strains and off-flavor producers was the main objective in winemaking. It is understandable, as in our experience after more than 30 years of wine yeast selection, it was shown that no more than 10% of the isolated native strains were positive contributors of superior flavors. Some species that systematically gave desirable flavors during these screening processes were Hanseniaspora vineae and Metschnikowia fructicola. In contrast to the latter, H. vineae is an active fermentative species, and this fact helped to build an improved juice ecosystem, avoiding contaminations of aerobic bacteria and yeasts. Furthermore, this species has a complementary secondary metabolism with S. cerevisiae, increasing flavor complexity with benzenoid and phenylpropanoid synthetic pathways practically inexistent in conventional yeast starters. How does H. vineae share the fermentation niche with other yeast strains? It might be due to the friendly conditions it creates, such as ideal low temperatures and low nitrogen demand during fermentation, reduced synthesis of medium-chain fatty acids, and a rich acetylation capacity of aromatic higher alcohols, well-known inhibitors of many yeasts. We will discuss here how inoculation of H. vineae strains can give the winemaker an opportunity to develop ideal conditions for flavor expression of the microbial terroir without the risk of undesirable strains that can result from spontaneous yeast fermentations.

Keywords: microbial terroir, mixed cultures, yeast consortia, low-input winemaking, minimal intervention

INTRODUCTION

Traditionally, our ancestors produced wines by exploiting the indigenous yeast diversity present on grapes without knowledge of their winemaking capability. As wine quality was highly variable within and across different vintages, the challenge at that time was to obtain consistency and to search for vinification conditions that allowed greater standardization of their own wine styles.
After Pasteur in 1866 showed that many yeast species rather than simple chemical reactions were responsible for wine fermentation (Barnett, 2000), wine masters from the beginning of the 20th century began to search for the “pure ferment” concept (Hansen, 1895; Regenberg and Hansen, 2001), so as to increase the reliability of vintage quality. This period saw the improvement of wine quality until the 1980s which promoted the successful growth of the wine industry globally and increased the consumption of better quality wine. Thanks to a considerable expansion of knowledge on the roles of yeast and bacteria in winemaking over the past 60 years (Ribereau-Gayon et al., 1951; Rankine, 1967; Amerine and Kunkee, 1968), wines became more flavorsome, especially more fruity, and less faulty than in the past (Lambrechts and Pretorius, 2000; Fleet, 2003; Ugliano and Henschke, 2009; Steensels and Verstrepen, 2014; Pretorius, 2020). However, in the 1990s, some winemakers and consumers noticed that wines in the general wine markets were becoming more uniform in flavor in terms of lacking complexity and diversity. In particular, the flavor characteristics of wines from major producing countries, which have specific terroirs, were becoming less apparent (Bisson et al., 2002). The majority of the wine producers were using the same conventional fermentation technology based on Saccharomyces cerevisiae (Jolly et al., 2014), which essentially precluded the opportunity for their respective microbial terroir to participate in the vinification process. This led some artisans to suggest that wine styles needed to return to their “grape roots” and develop the concept of microbial terroir flavors based on “low-input” winemaking strategies (Ramey, 1995). The challenge was to return to traditional winemaking technologies but guided by a greater wealth of knowledge compared with earlier times (Carrau, 2006; van Wyk et al., 2020). It is now well established that increased yeast diversity can contribute to the diversity of the volatile chemical composition of wine (Romani et al., 2020), which might increase the sensory diversity of wine flavor, a still controversial concept of sensory complexity (Varela et al., 2009; Smith, 2012; Köster and Mojet, 2016; Borren and Tian, 2021). Moreover, increased diversity can also result in process stability and productivity in microbial communities (Lehman and Tilman, 2000; Briones and Raskin, 2003). Whilst complex ecological studies are needed to prove this phenomenon in several fermentation niches, there is an increasing numbers of studies showing that mixed cultures can more efficiently exhaust nutrients when compared with single cultures (Medina et al., 2012; Perez et al., 2020). This effect might further allow the reduction of microbial contamination during wine maturation. However, studies in real winemaking conditions at winery scale are very scarce (Romani et al., 2020), and when scaling up this technology of increased yeast diversity, the risks of appearance of undesirable flavors might result. At industrial sized fermentations, unpredictable interactions within a complex natural microflora, as the result of increasing reductive or oxidative conditions (Fariña et al., 2012; Varela et al., 2021) will demand careful daily process control by tasting. The attraction for food and wine is all about the flavor phenotype (Cordente et al., 2012; Carrau et al., 2015). These winemaking strategies are not romantic, and understanding deeply wine microbiology management is fundamental to obtain high-quality differentiated wines which reflect the region. Our proposal for developing friendly starters is based on careful strain selection by flavor and low nitrogen demand that was developed in order to obtain consistent screenings of superior native strains (Carrau et al., 2015). In our experience these aspects combined with an active but moderate fermentation capacity at temperatures below 20°C, resulted in wines of desirable flavors with certain yeast strains. By this selection strategy, we detected Hanseniaspora vineae at the initial steps of fermentation. In addition, we have characterized this yeast as having very different metabolic synthetic pathways compared with Saccharomyces that enrich wines with several grape flavor compounds related to the three aromatic amino acids such as benzenoids (Martin et al., 2016b; Valera et al., 2020a), other phenylpropanoids, and isoprenoids (Giorello et al., 2019; Del Fresno et al., 2020; Martin et al., 2021b). In this perspective, we will discuss the concept of “friendly” yeasts for developing wine starters and how the inoculation of H. vineae strains compared with conventional Saccharomyces strains gives the winemaker new tools to manage the microbial terroir flavor expression without the risk of unreliable spontaneous fermentations (Knight et al., 2020; Griggs et al., 2021).

**FRIENDLY YEASTS COOPERATE WITH THE MICROBIAL TERROIR**

It is well known that the majority of commercial yeasts added to a grape juice fermentation as starter culture rapidly control the process by reaching above 90% of the total yeast flora (Fleet, 1993; Medina et al., 2013; Carrau et al., 2020). This situation is common with Saccharomyces starters that are highly competitive compared with the native microflora, which we can define as “selfish” or “unfriendly” strains (Rendueles and Ghigo, 2012). This species has evolved a highly competitive strategy for removing key nutrients in a grape juice, such as amino acids and vitamins, within a few hours (Alonzo-del-Real et al., 2019). Furthermore, they actively use the glycolytic and

![FIGURE 1](https://example.com/figure1.png)
alcohol fermentation pathways to exclude the native microflora of grape must niches, not only by producing ethanol and CO₂ but also increasing temperature and producing compounds such as short- and medium-chain fatty acids, isoacids, or higher alcohols that can inhibit other yeast species (Goddard, 2008; Valera et al., 2019). These aroma compounds can be defined as “toxic” intercellular communication mechanisms, in contrast to the reduced production of these flavors by a friendly yeast strain. During the last decade or more, many non-Saccharomyces starters in mixed culture fermentation with a strain of Saccharomyces have been studied, usually inoculated in sequential mode, so as to enhance the opportunity of the non-Saccharomyces to influence the winemaking process compared with co-inoculations where its impact can be limited by exclusion or competition with Saccharomyces (Padilla et al., 2016; Aranda, 2019; Borren and Tian, 2021). In summary, the competitive advantage of Saccharomyces over the majority of non-Saccharomyces yeasts can reside in various stress mechanisms, including nitrogen depletion, sugar transporter adaptations to high osmotic pressure of grape juice (high sugar content) and to a more active proton-pump ATPase adapted to low pH and high ethanol (Ganucci et al., 2018; Palmgren and Morsonn, 2019). These key adaptations that are strain dependent can explain why Saccharomyces species and their related hybrids typically dominate the fermentation niche. Interestingly, in reference to secondary fermentation compounds, that are inhibitors of cell activity, we have noted that some characteristics of H. vineae metabolism can explain the resulting friendlier environment from its activity. Metabolic reactions such as its extreme capacity for higher alcohol acetylation represents a well-known mechanisms for detoxifying acetates and their corresponding alcohol in the fermentation medium (Peddie, 1990). Furthermore, we have determined that H. vineae produces significantly lower concentrations of fatty acids and has a slower rate of ethanol and CO₂ production compared with Saccharomyces strains (Valera et al., 2020b). These attributes are characteristic of H. vineae within the apeculate group of the Hanseniaspora genus (Martin et al., 2018; Giorello et al., 2019; Valera et al., 2020a,b). During Chardonnay winemaking with H. vineae strains (Medina et al., 2013), we noted that in control-inoculated wines with S. cerevisiae ALG804, the native grape microflora was invariably dominated by this strain within several days. In Figure 1, we show a yeast profile of conventional wine fermentation inoculated with 10 g/Hl of active dry yeast, which clearly reveals numerical dominance by this species. In contrast, Figure 2 shows that inoculation with H. vineae HV025 permitted the participation of eight different strains at day 10 of the fermentation process. Although we can see in Figure 2 that two commercial strains appeared that might come from previous winemaking vintages, it is interesting to note that there were also four native Saccharomyces strains according to our yeast commercial data bank by DNA microsatellite profiles (Jubany et al., 2008).

THE IMPACT OF NON-SACCHAROMYCES STRAINS ON SENSORY WINE DIFFERENTIATION

Few publications have focused on demonstrating increased sensory diversity associated with fermentation microflora using high-quality sensory techniques. Over the past two decades, many investigations have shown a great diversity of chemical compound analysis by GCMS of wines made with non-Saccharomyces and mixed species cultures compared with Saccharomyces yeasts. A large range of volatile compounds can be detected, but their impact on the sensory properties of wine is often lacking (Spence and Wang, 2019). The challenge today is to correlate wine chemical data with formal sensory analysis using for example a trained panel of judges or simple methods at the winery (Leschaeve and Noble, 2010). Wines made with different yeast strains can often be consistently differentiated by sensory profiling techniques or can be judged as being sensorially more complex than others (Padilla et al., 2016; Binati et al., 2019). Many reports showing detailed chemical analyses will need to be validated with sensory analysis as they are merely showing just the chemical diversity of yeast metabolism in different winemaking conditions. Many of the reports on experimental vinification were performed with either chemically defined or natural grape fermentation media with either undefined or an excess concentration of yeast assimilable nitrogen (Carrau et al., 2017; Perli et al., 2020). Although it is clear that many of the recent non-Saccharomyces vinification studies have demonstrated an impact on wine quality in different grape varieties or under mixed culture conditions, there is still very limited information on the behavior of non-Saccharomyces strains at winery scale (Jolly et al., 2014; Comitini et al., 2017; Morata et al., 2020). Since 2007, white wines have been produced with H. vineae on a commercial scale (Medina et al., 2007) and could be sensorially differentiated from wines produced by conventional Saccharomyces fermentations (Medina et al., 2013; Lleixà et al., 2016; Martin et al., 2018; Del Fresno et al., 2020, 2021). However, although red wines made under these treatments can be differentiated by chemical techniques, the

![FIGURE 2](https://example.com/figure2.png) **FIGURE 2** This shows an example of the yeast profile of a Chardonnay grape must inoculated with a single strain of HV025. This treatment shows an increase in yeast diversity during fermentation, as H. vineae allows the native or winery microflora to participate during fermentation. We apply the concept of “friendly starters” to a yeast inoculum that allows various yeast species and strains to share the fermentation environment.
sensory differentiation or quality conclusions of these processes in red wine vinification at winery scale were less clear. Our observation is that young red wines might be differentiated more easily than strong body red wines and even less following barrel maturation. Chemically, red wines made with *H. vineae* showed the presence of increased concentrations of benzenoids and acetate esters when compared with conventionally vinified red wines. In the meantime, metabolic footprinting techniques of these wines (Howell et al., 2006) allowed us to show that *H. vineae* had contributed to the aromatic chemical composition of wine on an industrial wine fermentation scale (Martin et al., 2021a).

INCREASED YEAST DIVERSITY INCREASES FLAVOR COMPLEXITY

It is widely known that protection of biodiversity should be a main biological focus for conserving agricultural ecosystems. As we previously mentioned, it was demonstrated that increased biodiversity in a given ecosystem niche increased community stability and productivity (Lehman and Tilman, 2000). The flavor complexity concept is still not clear and some authors believe it is more of an increase of flavor compound diversity from a chemical point of view. However, the relevant concept should include the sensory complexity when we talk about fermented beverages (Tempère et al., 2018). Increased flavor diversity has been demonstrated by the use of mixed culture inocula by which increased flavor complexity from a sensory point of view could be achieved, for example, in Chardonnay (Soden et al., 2000; Medina et al., 2013) or Sauvignon Blanc wines (Anfang et al., 2009; Knight et al., 2018). More recently, sensory studies of mixed culture fermentations in some other white and red varieties, have been reported (Varela, 2016; Padilla et al., 2017; Hranilovic et al., 2018; Benito et al., 2019; Castrillo et al., 2019; Binati et al., 2020; Del Fresno et al., 2020; Romani et al., 2020; Muñoz-Redondo et al., 2021). In our experience, a chemically defined grape must medium with low assimilable nitrogen has allowed the selection of non-*Saccharomyces* and *Saccharomyces* strains with a combination of both low nitrogen demand and intense desirable flavors production (Carrau et al., 2015). This procedure contributed to the selection of a new generation of native yeast strains adapted for low-input winemaking strategies, where the addition of DAP can be avoided. It is known that ammonium salts inhibit the synthesis of some of the main aroma compounds of particular interest for the varietal character of some grapes, such as phenylpropanoids or sulfur thiols (Martin et al., 2016a). This strategy is expected to identify yeasts that can share fermentation medium nutrients as they will have decreased nitrogen demand characteristics and will ensure the development of clean flavors. However, further studies regarding unpredictable yeast interactions at winery scale are being carried out to better understand the appearance of sluggish fermentations in some rich sugar white grape musts such as Petit Manseng and Chardonnay grapes which would yield ethanol concentrations exceeding above 13% alcohol by volume (Carrau et al., 2020).

YEAST DIVERSITY FROM NATIVE ENVIRONMENTS TO THE WINERY

The systematic use of *Saccharomyces* starter cultures in wine production has contributed to more uniform wine quality in the past 50 years. However, it is now considered by some that there is a limitation of flavor diversity due to this phenomenon affecting the development of new wine styles. This has led to a decrease in the flavor differentiation of wines from regional terroir sites which were previously described as having a “typical”

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**FIGURE 3** Yeast diversity along the different environments of a wine region ecosystem. Winemakers have the alternative to increase strain diversity when working with yeast strains that might share the fermentation niche with other native strains. The number of species data of the different niches was obtained from recent studies (Morrison-Whittle and Goddard, 2018), and fermentation data with *H. vineae* were adapted from Carrau et al. (2020).
flavor profile as associated with a given region (Liu et al., 2019, 2020). In Figure 3, we show the interesting process of how yeast diversity in a certain wine region is reduced from the vine to the final wine. This graph clearly shows that the addition of a pure culture inoculum of Saccharomyces at the initial step of fermentation in the winery restricts the rich microbial terroir flora to a single culture process.

Studies concerning yeast diversity in a wine region showed that the grape plants, including the fruit and soil niche ecosystems, lodge the higher number of yeast species compared with the native and winery environments (Morrison-Whittle and Goddard, 2018). Figure 3 shows the process from nature to the fermentation ecosystem revealing how yeast diversity undergoes a slight increase at the fruit ecosystem and, subsequently, a continuous decrease until the commencement of fermentation. Winemaker management of fermentation, that is interventions, can increase or even decrease yeast diversity to a single fermentation strain. In contrast to the rich biodiversity in a certain wine region, conventional inoculation of a pure Saccharomyces ferment finished with a single culture fermentation process.

DISCUSSION

The development of “friendly” yeasts can promote a sharing of the fermentation medium with some strains from natural environments so as to recover potential metabolic diversity along the process. The concept of introducing a starter with the characteristics described is based on the difficult task that would be incurred by a winery in selecting native yeasts with superior flavors from their vines, as it has been mentioned such strains might represent less than 10% of the total natural microflora. We have shown that a selection strategy for friendly yeast based on sensory analysis at low YAN concentrations and moderate fermentation capacity minimize the risk of sluggish processes or the production of undesirable flavors such as acetic acid, hydrogen sulfide, or acetaldehyde. H. vineae might also well fit the concept of friendly yeast owing to its low formation of higher alcohols and fatty acids compared with Saccharomyces, known inhibitors of many yeast strains. However, this approach to allowing the natural microflora to participate in the process should be controlled daily by tasting. However, in the wineries the fermentation increase microbial competition under such complex situations might be a small risk compared to the benefit of obtaining increased flavor complexity. The potential risks that could result from this approach are believed to be significantly smaller than the application of spontaneous fermentation processes. The future challenge for wine microbiologists and winemakers will be to understand fermentation from a holistic view point, so as to provide an alternative management of the process with the aim of minimizing the loss of strain diversity which would contribute to the loss of terroir characteristics. Strain interactions which increase strain diversity give infinite opportunities to explore flavor development. It is well known that in a complex consortia fermentation, there are many factors that can change the final flavor of a wine, such as size of the inocula, nutrient competition, metabolite–cell and cell–cell interactions, temperature, tank sizes, redox situations, and aeration. The development of friendly yeasts that share their environment with other strains is an interesting low-input winemaking strategy. This biotechnological approach might be considered a “romantic” way of producing particular or unique wines representing regionality. However, these strategies demand deep knowledge of microbiology and systematic tasting and sensory training at the winery to obtain quality and market differentiation.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article, further inquiries can be directed to the corresponding author.

AUTHOR CONTRIBUTIONS

FC wrote the article. PH wrote part of the article and reviewed the general presentation. Both authors contributed to the article and approved the submitted version.

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Carrau and Henschke

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