



AperTO - Archivio Istituzionale Open Access dell'Università di Torino

# **Bioprotection strategies in winemaking**

This is the author's manuscript
Original Citation:
Availability:
This version is available http://hdl.handle.net/2318/1851245 since 2022-06-22T15:24:09Z
Published version:
DOI:10.1016/j.ijfoodmicro.2022.109532
Terms of use:
Open Access
Anyone can freely access the full text of works made available as "Open Access". Works made available under a Creative Commons license can be used according to the terms and conditions of said license. Use of all other works requires consent of the right holder (author or publisher) if not exempted from copyright protection by the applicable law.

(Article begins on next page)

1	Bioprotection strategies in winemaking
2	
3	Paola Di Gianvito, Vasileios Englezos, Kalliopi Rantsiou, Luca Cocolin
4	
5	Università degli Studi di Torino, Dipartimento di Scienze Agrarie, Forestali e Alimentari, Largo
6	Braccini 2, 10095 Grugliasco, Italy.
7	
8	*Corresponding author: Luca Cocolin, Fax: +39-011-6708553, e-mail: lucasimone.cocolin@unito.it.
9	
10	
11	
12	
13	
14	
15	
16	
17	
18	
19	
20	
21	
22	
23	
24	
25	
26	

#### 27 Abstract

Worldwide the interest for biological control of food spoilage microorganisms has significantly increased over the last decade. Wine makes no exception to this trend, as consumer demands for wines free of preservatives that are considered negative for human health, increase. Biological control during wine fermentation aims at producing high quality wines, while minimizing, or even eliminating, the use of chemical additives. Its success lies in the inoculation of microorganisms to prevent, inhibit or kill undesired microbes, therefore maintaining wine spoilage at the lowest level. The food industry already makes use of this practice, with dedicated commercial microbes already on the market. In winemaking, there are commercial microbes currently under investigation, particularly with the aim to reduce or replace the use of sulfur dioxide. In this review, the potential of wine yeasts and lactic acid bacteria as bioprotection agents and their mechanisms of action during wine fermentation are presented.

41 Keywords: Bioprotection mechanisms; Wine fermentation; Antagonism strategies; Competition
42 strategies; Yeasts; Lactic Acid bacteria

55 There is a growing concern about the negative effects on human health of chemical additives 56 used in foods and drinks to improve their stability and shelf life. For this reason, several research 57 lines are focusing on the development of alternative strategies to avoid spoilage. Bioprotection (or 58 biopreservation or biological control) fits well in this new concept of spoilage prevention, and several 59 microorganisms or their antimicrobial products have already been identified as bioprotection agents 60 (Dokka et al., 2018; Mewa-Ngongang et al., 2019; Mukherjee et al., 2020; Pu et al., 2014; Singh et 61 al., 2018;). This practice is widely used, mainly in agriculture and food industries for the protection 62 of fruits from postharvest spoilage microorganisms, and the extension of the shelf life of food products (Ferraz et al., 2019; Keswani et al., 2019). For example, a commercial product based on 63 64 Aureobasidium pullulans was developed to protect grapes, strawberry and tomato against 65 Penicillium, Botrytis and Monilinia, while a product based on Metschnikowia fructicola is used to 66 protect strawberry, blueberry, grape, stone fruit and pome against Botrytis, Penicillium, Rhizopus, 67 Aspergillus and Monilinia (Zhang et al., 2020). To date, there is a patent (WO 2015/110484) related 68 to the exploitation of a Lactiplantibacillus plantarum strain as bioprotective agent against undesired 69 microorganisms in wine (Krieger-Weber et al., 2020). Different genera of bacteria (e.g. Lactiplantibacillus, Bacillus, Pseudomonas, Streptomyces), fungi (Trichoderma) and yeasts 70 71 (belonging mainly to the genera, Candida, Cryptococcus and Aureobasidium) are commercially 72 available for these purposes (Droby et al., 2016; López-Seijas et al., 2020; Mukherjee et al., 2020).

The bioprotection strategy consists of the inoculation of living microorganisms (Bio Protective Cultures, BPCs), or the addition of their metabolites (Bio Protective Metabolites, BPMs), in purified form, during the production of the food or there after. These microorganisms prevent food microbial spoilage through different bioprotection mechanisms that can be divided in passive (competition for space, nutrient and oxygen) and active antagonistic strategies (production of antimicrobial molecules) (Pandin et al., 2017). The addition of BPCs during the early stages of the production process could also enhance the characteristics of fermented foods, such as flavour, texture
and nutritional value (Gaggia et al., 2011).

81 In the wine industry, the most common chemical additive is sulfur dioxide (SO<sub>2</sub>). This 82 compound is considered an essential tool for winemakers, due to its low-cost and its antioxidant and 83 antimicrobial properties against a wide spectrum of microorganisms (Roullier-Gall et al., 2017). 84 Therefore, its presence during both the fermentation and the storage of the wine is desired to avoid 85 spoilage. In spite of the advantages, high doses of SO<sub>2</sub> may cause consumer health problems, 86 particularly in sensitive individuals (Guerrero and Cantos-Villar, 2015). For this reason, the World 87 Health organization (WHO) encourages alternative methods to reduce, or even eliminate, its use in 88 wine production. Consequently, several alternative technologies able to control microorganisms have 89 been investigated. These include ultrasound, ultraviolet radiation, pulsed electric field, electrolyzed 90 water, high hydrostatic pressure treatments, or the addition of lysozyme, sorbic acid, dimethyl 91 dicarbonate and chitosan (Guerrero and Cantos-Villar, 2015). However, a definitive substitute for 92 SO<sub>2</sub> has not yet been found, especially for wines undergoing long-term storage (Giacosa et al., 2019). 93 The application of bioprotection strategies in viticulture has increased significantly in the last 94 years, with studies investigating its applicability against grapevine trunk diseases (Mondello and 95 Songy, 2018), the fungal pathogen Botrytis cinerea causing bunch (Carbó et al., 2018; Garrido et al., 96 2017; Jacometti et al., 2010) and sour rot (Carbó et al., 2018) in the vineyard (Nardi et al., 2020). 97 However, little is known about the application of bioprotection strategies during or after the 98 completion of the fermentation process (Simonin et al., 2018). Bioprotective strategies found in wine 99 yeasts and Lactic Acid Bacteria (LAB) are represented in Fig. 1. In this review, the current knowledge 100 about bioprotection opportunities and mechanisms exerted by wine yeasts and LAB, during

102

101

#### 103 2. Bioprotection mechanisms

fermentation to control spoilage microorganisms, will be discussed.

105 Wine microorganisms form complex ecological interactive webs, which result in the 106 dominance of a specific species or strains able to determine the final quality of wine. In inoculated 107 wine fermentations, the use of a starter culture aims to establish dominance of the inoculated culture 108 and determine wine characteristics. Conversely, the BPCs aim to prevent the growth of the spoilage 109 microorganisms, without affecting the performances of useful microorganisms responsible for the 110 alcoholic/malolactic fermentation (Nardi, 2020). Furthermore, BPCs may also be directly responsible 111 for wine fermentation (Table 1). Understanding the antagonistic mechanisms that confer 112 bioprotection activity to the inoculated microorganisms is fundamental for their successful 113 application as BPCs. These competition strategies can be either passive or active (Bauer et al., 2018) 114 as shown in Fig. 2. The difference is that passive competition strategies are exerted by all 115 microorganisms, while active competition strategies are performed only by some microorganisms 116 that, consequently, result stronger than the other and able to dominate the fermentation process.

117

# 118 2.1. Passive competition strategies

119

Passive competition strategies comprise all the mechanisms that are beneficial including strategies to take away resources from competitors (exploitative competition), or to achieve a superior positioning within the niche (Parijs and Steenackers, 2018). During wine fermentation, the factors that mainly modulate the fitness advantage of some species above others are the ability to rapidly uptake nutrients (carbohydrates, nitrogen, vitamins, sterols and microelements) and oxygen or to colonize the space (Curiel et al., 2017; Petigonnett et al., 2019; Prior et al., 2019; Rollero et al. 2018; Siedler et al., 2019; Taillandier et al. 2014).

127

128 2.1.2. Nutrients competition

130 In a specific environment all microorganisms compete with each other for nutrients necessary 131 for their growth (Liu et al., 2015). Passive competitive microorganisms are able to consume the substrates faster and in a more efficient way than the competitors, producing higher levels of biomass 132 133 (Boynton et al., 2019). Proteomic and transcriptomic studies have revealed that, during mixed wine 134 fermentation, an increased resource uptake (glucose, nitrogen, oxygen, sterols, copper, iron and 135 vitamins) occurring in several yeast strains, compared to the respective control pure culture 136 fermentations. These yeasts belong to Saccharomyces cerevisiae (Alonso-del-Real et al., 2019; 137 Barbosa et al., 2015; Curiel et al., 2017; Kosel et al., 2017; Peng et al., 2019; Rossouw et al., 2012; 138 Shekhawat et al., 2019; Tronchoni et al., 2017;), Torulaspora delbrueckii (Prior et al., 2019; Su et al., 139 2020), even in presence of different nitrogen sources (Roca-Mesa et al., 2020), and Lachancea 140 thermotolerans (Shekhawat et al. 2019). Furthermore, nutrient competition was identified as the 141 major antagonistic strategy exhibited by Lpb. plantarum (Sielder et al., 2020). A metagenomic 142 analysis revealed that during wine fermentation, this species has the largest number of 143 phosphotransferase systems (the major mechanism used by bacteria for carbohydrates uptake) among 144 the bacteria observed in wine (Melkonian et al., 2019). The ability of the inoculated microorganisms 145 to subtract high levels of nutrients from the medium, could increase the microbial stability in the 146 wines at the end of the fermentation process by inhibiting the growth of unwanted microorganisms 147 and reduce or even avoid the use of SO<sub>2</sub>. Since, residual nitrogen levels could favor the growth of 148 undesirable microorganisms in unwanted steps of wine production chain and could act as precursors 149 for the formation of biogenic ammines (Restuccia et al., 2018). However, particular attention should 150 be paid in order to find the most appropriate strain selection that fits with the fermentation conditions, 151 since low nitrogen concentration may lead to stuck or sluggish fermentations (Bisson et al., 1999).

152 Competition for nutrients can also be achieved by the secretion of extracellular molecules 153 such as digestive enzymes, or siderophores (nutrient exclusion) (Ghoul and Mitri, 2016). In this 154 context, antibacterial and antifungal activity of *Metschnikovia pulcherrima* and *Metschnikovia* 155 *fructicola* was mainly attributed to the iron depletion caused by the insoluble maroon-red

156 pulcherrimin pigment (Lachance, 2016; Sipiczki, 2006; 2020). In particular, this antimicrobial feature 157 is the result of the reduction of the iron in the medium, due to the precipitation of iron (III) ions, 158 caused the precursor of the pulcherrimin pigment called pulcherriminic acid. As a result, low iron 159 levels in the medium could limit the growth of the microorganisms that require this compound for 160 their growth. This mechanism was found to have an inhibitory effect against the following 161 microorganisms: Candida tropicalis, Candida albicans, Brettanomyces/Dekkera, Hanseniaspora and 162 Pichia genera, B. cinerea and Penicillium (Morata et al., 2019a). While, contradictory results were 163 observed against S. cerevisiae (Kantor et al., 2015, Melvyda et al., 2020, Oro et al., 2019). This 164 behaviour let us to hypothesize a strain specific inhibition. Therefore, careful selection of S. cerevisiae 165 strains should be performed to find the most suitable partner of *M. pulcherimma* in mixed culture 166 fermentations to reduce ethanol content and enhance the aroma volatile fraction of the wines (Jolly 167 et al., 2014). However, there was not a clear correlation between the antagonistic activity and the iron 168 deprivation by pulcherrimin (Gore-Lloyd et al., 2019; Saravanakumar et al., 2008). In a recent study, 169 a genome comparison between a *M. pulcherrima* wild type and three pigmentless mutants, allowed 170 the identification of a point mutation in the MPUL0C08850 gene, which encodes for an ortholog of 171 S. cerevisiae SNF2 gene. The resultant truncated form of this transcriptional regulator caused a down-172 regulation of the transcription of PUL genes (pulcherrimin biosynthesis and utilisation) explaining 173 why the colourless strains showed an antimicrobial behaviour even if it was less than the wild type 174 (Gore-Lloyd et al., 2019). These authors speculated that the antagonistic activity of *M. pulcherrima* 175 was due to the combination of the effect of pulcherrimin and diverse proteins regulated by SNF2 such 176 as genes encoding different types of transporters or secreted proteases (Gore-Lloyd et al., 2019).

A similar system was also identified in many Lactobacilli, including *Lpb. plantarum*. In particular, Sielder et al. (2020) demonstrated that the Lactobacilli showed an overexpression of a manganese transporter (*mntH*1 gene) during milk fermentation. The depletion of this microelement in the media resulted in the reduction of the growth of several yeasts (also present in wine) such as *Debariomyces hansenii* and *T. delbrueckii*. These authors found that for *T. delbrueckii* the addition 182 of manganese restored the growth only partially and hypothesized that other mechanisms183 (antimicrobial compounds production) were involved in the antagonistic interactions.

184

185 *2.1.3. Space competition* 

186

187 Another passive competition phenotype is the ability to colonize the space where competitors are 188 established (Ghool and Mitri, 2016) or grow faster than other microorganisms. With this purpose, 189 microorganisms are able of change their lifestyle forming multicellular structures like biofilms 190 (Váchová and Palková, 2018). These structures may exhibit different phenotypic characteristics, 191 compared to free-floating cells. Usually, after the adhesion of single cells to the host surface, the 192 biofilm formation determines the modification of the cell wall properties due to the secretion of 193 extracellular matrix and often the formation of hyphae and pseudohyphae (Freimoser et al., 2019). 194 The biofilm formation is considered a desired characteristic in biocontrol yeasts and has been widely 195 investigated. For example, M. pulcherrima and Pichia kluvveri are able to form biofilms on grape 196 berries surface (Cordero-Bueso et al., 2017; Parafati et al., 2015; Pawlikowska et al., 2019). This 197 ability confers a great competitive advantage to these yeasts to colonize both intact and damaged 198 grapes as well as establishing themselves in grape juice and starting the fermentation process even in 199 harsh conditions such as those found in desiccated grapes (Cordero-Bueso et al., 2017). 200 Microorganisms with ability to form biofilms are generally characterised by a higher competitiveness 201 against pathogens due to the higher environment persistence and higher biocontrol activity (Freimoser 202 et al., 2019). Furthermore, it has been demonstrated that Oenococcus oeni is able to produce 203 exopolysaccharides and form biofilms on cellar surfaces (Dimopoulou et al., 2015). This organization 204 is formed by stress-tolerant cells able for efficient malolactic fermentation under winemaking 205 conditions (Bastard et al., 2016).

206

207 2.1.4. Competition for oxygen

209 The presence of oxygen during the fermentation process is considered as one of the key factors that 210 regulate the presence and population of yeast species. This molecule could positively affect ethanol 211 tolerance and the fermentation performance of yeast cells, through its key role in the production of 212 sterols and unsaturated fatty acids (Albergaria and Arneborg, 2016). Grape must contains relatively 213 low dissolved oxygen levels and becomes rapidly anaerobic due to the fast consumption by the 214 microbial consortium and the grape-derived polyphenol oxidase (Romano et al., 2019). Therefore, 215 the concentration of oxygen may have an important role in the survival and dominance of a specific 216 yeast during the fermentation process. Indeed, the low levels could promote the growth of species 217 that are able to grow in anaerobic conditions, like S. cerevisiae (Hansen et al., 2001). While, on the 218 other hand low oxygen levels can influence the growth and death rate of several non-Saccharomyces 219 yeast species like T. delbrueckii, L. thermotolerans, Starmerella bacillaris and Hanseniaspora vinae 220 and reduce the competitiveness against S. cerevisiae (Englezos et al., 2018a, Hansen et al., 2001, Yan et al., 2020). Oxygen addition is also believed to enhance the impact of non-Saccharomyces yeasts 221 222 on the chemical and aroma composition of the wines produced by mixed culture fermentations 223 (Englezos et al., 2018a, Hansen et al., 2001, Yan et al., 2020). While the limited addition of oxygen 224 at the beginning of mixed fermentations with M. pulcherrima or Starm. bacillaris and S. cerevisiae 225 could help to reduce the ethanol content in wines due to the higher sugar consumption of the first 226 yeast species (Varela et al., 2017). However, a fine tuning of oxygen addition is required, since 227 excessive oxygenation might lead to stuck fermentation due to increased antagonism between non-228 Saccharomyces and S. cerevisiae yeasts, as previously observed for H. vinae and S. cerevisiae (Yan 229 et al., 2020). Interestingly, the use of selected yeasts species or strains within species could potentially 230 protect the wine against oxidation due to the rapid consumption of oxygen, as previously 231 demonstrated for T. delbrueckii (Simonin et al., 2018). The effect of oxygen on yeast interactions 232 during mixed fermentations and as consequence on wine characteristics needs further investigation.

236 Generally, active competition includes the strategies exploited to confer a disadvantage to the 237 unwanted species through a direct and active interference (Ghoul and Mitri, 2016) due to the 238 production of antimicrobial compounds or through a contact dependent inhibition (Parijs and 239 Steenackers, 2018). During fermentation, yeasts release ethanol that is the most studied antimicrobial 240 compound. Its toxicity on cells is well understood (Gao and Fleet, 1988). Yeasts are also able to 241 release SO<sub>2</sub> as result of their metabolism (Andorrà et al., 2018). Some S. cerevisiae strains can 242 produce more than 100 mg/L of this compound (Rauhut, 2017), while information regarding its 243 production by non-Saccharomyces yeasts are scarce, probably because these yeasts are generally 244 more sensitive to SO<sub>2</sub> than S. cerevisiae (Jolly et al., 2014). However, some spoilage yeasts namely 245 Zygosaccharomyces bailii and Brettanomyces spp. are more tolerant to these conditions than S. 246 *cerevisiae* and therefore are still detected in wines with relatively high SO<sub>2</sub> levels.

247 Several studies highlighted that an array of other antimicrobial compounds are produced in 248 wine environment by yeasts (Albergaria and Arneborg, 2016; Balmaseda et al., 2018; Liu et al., 2017) 249 and bacteria (Bartle et al, 2019; Siedler et al., 2019) of oenological interest. These molecules differ 250 and possess diverse inhibition mechanisms. Antimicrobial compounds are actually subjected to an 251 extensive research with the purpose to exploit their action during wine fermentation. However, it is 252 important to consider that these compounds are often produced after the logarithmic stage and 253 therefore their competitive ability is strictly correlated with the growth performance of the inoculated 254 strain (Mazzucco et al., 2019; Singh, 2018; Villaba et al., 2016). However, the use of these 255 compounds for commercial purpose cannot be authorised without the approval by the International 256 Office of Vine and Wine (OIV) and/or the national regulators (Mehlomakulu et al., 2015). On the 257 other hand, the yeasts producing these compounds can be used if their origin is oenological. In the 258 next sections, the active antagonistic strategies used by yeasts and LAB that confer a possible bioprotection in wine will be discussed. These molecules can be grouped according to the chemical 259

groups they possess. In particular, nitrogen antimicrobial compounds are antimicrobial peptides,
killer toxins, bacteriocins and lytic enzymes as listed in the next section.

262

# 263 2.2.2. Antimicrobial peptides and proteins

264

#### 265 2.2.1.1 Antimicrobial peptides

266

Antimicrobial peptides (AMPs) are low molecular weight oligopeptides with a varying number (from five to over a hundred) of amino acids and high sequence variability. Furthermore, the majority of AMPs present charged and hydrophobic amino acids at physiological pH, defining the amphipathic nature of these molecules (Zhang et al., 2019). AMPs are evolutionary conserved in the genome and have a broad antimicrobial spectrum of action from viruses to parasites (Bahar and Ren, 2013). Up to date, more than 5,000 AMPs have been discovered from prokaryotes (e.g., bacteria) and eukaryotes (e.g., protozoa, fungi, plants, insects, and animals) or synthesized (Zhao et al., 2013).

274 Several studies reported that S. cerevisiae is able to secrete a proteinaceous antimicrobial 275 compound able to inhibit O. oeni growth and as a consequence malolactic fermentation (Comitini et 276 al., 2005; Osborne and Edwards, 2007; Mendoza et al., 2010; Nehme et al., 2010; Rizk et al., 2016). Furthermore, the AMP "saccharomycin" produced by S. cerevisiae was found to act as a natural 277 278 biocide (2-10kDa) against several wine-related non-S. cerevisiae yeasts such as Hanseniaspora 279 uvarum (Pérez-Nevado et al., 2006), Brettanomyces/D. bruxellensis and LABs (Branco et al., 2014). 280 These authors demonstrated that saccharomycin is comprises by two anionic (isoelectric point of 281 4.37) peptides (AMP1 and AMP2/3) able to alter intracellular pH, membrane permeability and 282 cultivability of non-Saccharomyces strains. These two peptides are fragments of the glycolytic 283 enzyme glyceraldehyde-3-phosphate dehydrogenase (GAPDH); which is considered an energy 284 metabolism-related enzyme, able to generate NADH in glycolysis (Branco et al., 2017).

285 The production of these molecules is widespread among wine related yeasts and bacteria and 286 the use of yeasts able to release AMPs was recently proposed for the biocontrol against undesired 287 microorganisms. However, this strategy has been poorly applied to wine-related microorganisms 288 (Peña and Ganga, 2019). A recent study reported the production of an AMP by the non-289 Saccharomyces strain Candida intermedia LAMAP1790. The application of this wine strain affected 290 the growth of several strains of Brettanomyces bruxellensis, without influencing S. cerevisiae 291 performance during fermentation (Peña and Ganga, 2019). These findings opens new prospects 292 regarding the use of these AMPs as alternatives to SO<sub>2</sub> in wine production industrial process 293 (Albergaria and Arneborg, 2016). Since, some spoilage microorganisms like Pichia spp., Dekkera 294 spp. and Z. bailii could tolerate medium-high SO<sub>2</sub> levels and the AMPs might have a key role in 295 controlling unwanted microorganisms without affecting the beneficial microorganisms.

296

297 2.2.1.2 Killer toxins

298

299 Killer toxins play an important role in the defence mechanisms of the yeast starter culture 300 against undesired microorganisms. This character consists in the production of proteins or 301 glycoproteins (molecular weight between 30 and 70 kDa) that are lethal for sensitive cells 302 taxonomically related or not. It was demonstrated that the action of these compounds is mediated by 303 the presence of two kinds of specific receptors in the sensitive microorganism cell wall (Liu et al., 304 2015). Even if this characteristic was first observed in S. cerevisiae, it is well distributed in several 305 yeast genera like Candida, Ogataea, Pichia, Williopsis, Tetrapisispora, Schwanniomyces, 306 Debaryomyces, Ustilago, Cryptococcus, Metschnikowia, Kluyveromyces and Zygosaccharomyces 307 (de Ullivarri et al., 2014; Liu et al., 2015). Killer toxins are encoded chromosomally, or their 308 production has been related with the presence of dsRNA viruses, or linear dsDNA plasmids (Belda 309 et al., 2017), however this is the case of some of the non-Saccharomyces species listed above. Killer 310 yeasts have been used to control pathogenic fungi in plants (Schmitt & Breinig, 2002) and to develop antimycotics for the treatment of human and animal infections (Liu et al., 2015). Furthermore, they
were purposed as an alternative strategy of biocontrol to combat contaminating wild yeasts in food
industries and in particular in winemaking (Çorbacı, and Uçar, 2018; de Ullivarri et al., 2014;
Mazzucco, et al., 2019; Santos et al., 2011; Villaba et al., 2016;).

315 The possibility of using S. cerevisiae as BPCs (active antagonistic phenotype) in wine was 316 first introduced in the 1960s, when it was found that S. cerevisiae strains are able to release killer 317 toxins and inhibit other S. cerevisiae strains as well as spoilage yeast species using reduced SO<sub>2</sub> 318 concentrations (de Ullivarri et al., 2014). These authors proposed the use of a sequential inoculation 319 in wine of two S. cerevisiae killer strains (Cf8 and M12) to improve the biocontrol against undesired 320 wine yeasts Dekkera anomala and Z. bailii. This new strategy showed a reduction in the growth of 321 the undesired yeasts to about 78% and 50%, respectively. Subsequently, Oro et al. (2016) found that 322 Kwkt and Pikt, two killer toxins produced by Kluyveromyces wickerhamii and Wickerhamomyces 323 anomalus, respectively, showed an antimicrobial activity against B. bruxellensis. These authors 324 demonstrated that this spoilage wine yeast, after a treatment of 0.7 mg/l of SO<sub>2</sub> at pH 3.2, entered in 325 a viable but non culturable state. Divergently, after treatment with Kwkt or Pikt, B. bruxellensis D46 326 strain completely lost its viability. Furthermore, Mehlomakulu et al. (2017) exposed this spoilage 327 yeast to the killer toxin CpKT1 produced by Candida pyralidae and demonstrated that the loss of 328 viability was due to the induction of cell membrane and cell wall damages. Mazzucco et al. (2019) 329 also observed an inhibition against *B. bruxellensis*. These authors, demonstrated that the killer toxin 330 SeKT produced by Saccharomyces eubayanus in wine, could be used for the biocontrol of four 331 common spoilage wine yeasts, namely B. bruxellensis, Pichia membranifaciens, Pichia 332 guilliermondii and Pichia manshurica.

Villaba and collegues (2016) investigated the ability of *T. delbrueckii* to release a killer toxin (TdKT). In particular, the NPCC 1033 strain was tested for its killer activity against some undesired wine yeasts, namely *H. uvarum*, *B. bruxellensis*, *P. guilliermondii* and *P. membranifaciens* (Lopes and Sangorrín, 2010). It was also verified that the TdKT toxin was active in oenological conditions 337 (at different pH and temperatures values, ethanol, glucose and SO<sub>2</sub> concentrations simulating several 338 wine environmental conditions). It was also found that some L. thermotolerans strains secreted killer 339 toxins against S. cerevisiae (Aponte and Blaiotta 2016), while Nally et al. (2018) demonstrated that 340 two strains were able to inhibit Aspergillus without affecting S. cerevisiae in a mixed fermentation. 341 Some *P. kluyveri* strains were found able to release killer toxins in the medium inhibiting the growth 342 of some food and beverage spoilage yeast genera like Dekkera, Kluyveromyces, Pichia, 343 Saccharomyces and Zygosaccharomyces (with the highest activity against D. bruxellensis) (Labbani 344 et al., 2015; Zagorc et al., 2001). Finally, the use of killer strains in sparkling wine production was 345 proposed in order to accelerate the maturation time (Todd et al., 2000, Velázquez et al., 2016, 2019). 346 Since, the production of this category of wines using the traditional Methode Champenoise is 347 characterized by a secondary fermentation in a bottle of a base wine, followed by a long period of 348 aging in which the yeast cells are in contact with the fermented wine. During this period, which may 349 last several months, yeast cells undergone autolysis and release cell components into the wine with 350 positive repercussions on the aroma, mouthfeel and foaming properties of wines (Di Gianvito et al., 351 2019). Such investigations, suggested that killer T. delbrueckii strains and sensitive yeasts in mixed 352 cultures could accelerate the onset of yeast autolysis, improving the organoleptic quality and foam properties of sparkling wines, without affecting fermentation kinetics (Velázquez et al., 2016, 2019). 353 354 Another study demonstrated that the use of a combination of a killer S. cerevisiae strain and a sensitive 355 Saccharomyces bayanus induced the autolysis of the sensitive yeast in a shorter time, compared to 356 the control wine (without the addition of these yeasts), under oenological conditions (Lombardi et al., 357 2016). In particular, in a pilot scale production, the selected strains influenced the concentrations of 358 free amino acids, total proteins and polysaccharides reaching, after 3 months of aging, the levels 359 showed by control wines after nine months of aging. These findings, raise the possibility of using a 360 biotechnological approach based on specific combinations of killer/sensitive strains or different types 361 of killer strains to reduce the time needed to release of cell components in the medium and

362 consequently the time for sparkling wine production with economic benefits for the producers363 (Mannazzu et al., 2019).

364

# 365 2.2.1.3 Bacteriocins and bacteriocin-like inhibitory substance (BLIS)

366

367 Many wine related LAB are known for their ability to produce proteinaceous antimicrobial 368 molecules called bacteriocins, active against other bacteria that are closely related to the producing 369 organism growing in the same medium (Ndlovu et al., 2015). Several studies highlighted the ability 370 of wine LAB to produce bacteriocins, however, bacteriocin producing activity has not yet been 371 demonstrated under wine conditions (Krieger-Weber et al., 2020) and their efficacy in wine is only 372 demonstrated in combination with SO<sub>2</sub> (Royo-Bezares et al., 2007). In fact, the production and application of bacteriocins in food is influenced by the physical and chemical conditions of food and 373 374 the presence of competitive microorganisms (Singh et al., 2018).

375 Knoll et al. (2008) found that both indigenous and commercial O. oeni strains showed an 376 antagonistic activity against several wine-related LAB strains in vitro. These authors identified some 377 putative bacteriocin-encoding genes, but they did not demonstrate the release of these bacteriocins in 378 wine. Recently, Lasik-Kurdyś and Sip (2019) reported the ability of one O. oeni strain to synthetize 379 a peptidic BLIS active against Leuconostoc mesenteroides and Pediococcus pentosaceus. In 380 particular, a synergistic effect between the presence of *O. oeni* (competition for nutrients and space) 381 the release of BLIS and other inhibition compounds (organic acids and H<sub>2</sub>O<sub>2</sub>) was highlighted. It was 382 further found that the removal of viable cells, as well as the neutralization of the supernatant or the 383 treatment with the catalase enzyme, significantly decreased or removed the inhibition activity against 384 some LAB indicator strains. Although several studies have indicated the use of bacteriocins in food 385 and beverages production, the application of these molecules in wine has a great potential, but it's 386 not yet approved. Interestingly, studies have proposed the use of bacteriocins to control biofilm 387 formation on stainless steel tanks and surfaces (Nel et al., 2002), suggesting that their application 388 could be a valid response to the chemical cleaning detergents with a negative impact to the 389 environment.

390

391 2.2.1.4 Lytic enzymes

392

393 The production of cell wall lytic enzymes by wine related yeasts and bacteria is a less known 394 form of antagonism. However, there are growing evidences that suggest the involvement of cell-wall 395 degrading enzymes in the mechanism of action of BPCs. Lytic enzymes such as  $\beta$ -glucanase 396 determine the leakage of cell contents by cleaving the  $\beta$ -glucan polysaccharide of fungi and bacteria, 397 such as Penicillium expansum, Fusarium oxysporum and B. cinerea (Edison et al., 2018; Wisniewski 398 et al., 1991). Cordero-Bueso et al., (2017), observed the release of lytic enzymes by P. kluyveri that 399 could result in the degradation of the fungal cell wall ( $\beta$ -1, 3-glucanase, proteolytic and pectinolytic 400 activities). Moreover, on grapes A. pullulans releases extracellular  $\beta$ -1, 3-glucanase which could play 401 a role in the biocontrol activity towards grape pathogens (Bozoudi and Tsaltas, 2018). Lastly, several 402 strains of the *M. pulcherrima* clade secrete extracellular chitinase and  $\beta$ -1, 3-glucanases able to 403 degrade the cell wall of B. cinerea, however their impact is strain and matrix-dependent (Sipiczki, 404 2020).

405

406 2.2.2. Volatile Organic Compounds (VOCs)

407

Among mechanisms of antagonism, VOCs are considered superior over the nonvolatile compounds ones because direct contact and closeness among microorganisms are not required. In fact, these small organic molecules (<C<sub>20</sub>) can diffuse to greater distances in heterogeneous environment, since are characterized by a low molecular mass (100 to 500 Da), a lipophilic moiety, low solubility in the water, low boiling point and a high vapor pressure that permit a readily volatilization at ambient temperatures (Zhang et al., 2021). VOCs are molecules potentially produced 414 by all living microorganisms and can be ascribed to several molecular classes, including 415 hydrocarbons, alcohols, thioalcohols, aldehydes, ketones, thioesters, cyclohexanes, heterocyclic 416 compounds, phenols, and benzene derivatives and can be produced by the primary metabolism or via 417 specialized secondary metabolic pathways.

It is widely known that wines produced by mixed fermentations have a complex aroma profile, but the role of these volatile molecules in microorganisms' interactions is still poorly understood. This is the case of the higher alcohols tryptophol, phenylethanol, and tyrosol which are produced at relative high levels by *S. cerevisiae*. Recently, interactions studies highlighted their role as antagonistic VOCs. In fact, through *in vitro* experiments, it was demonstrated that these molecules caused the reduction of the growth of non-*Saccharomyces* like *H uvarum*, *Starm. bacillaris*, *T. delbrueckii* and *M. pulcherrima*, even at low concentration (Gonzalez et al., 2018).

425 In another study, Farbo et al. (2018) attributed the bioprotective activity of L. thermotolerans 426 against toxinogenic fungi (Aspergillus spp.) to the production of VOCs, particularly 2-phenylethanol. 427 These authors demonstrated that this volatile compound was able to downregulate key genes 428 implicated in Ochratoxin A biosynthesis in a species-dependent way. The implication of this higher 429 alcohol in the antagonistic role during alcoholic fermentation was also suggested by the 430 transcriptomic profile of the yeast during fermentation. In fact, it was highlighted that in a mixed 431 fermentation with S. cerevisiae, L. thermotolerans showed an up regulation of all genes involved in 432 the conversion of phenylalanine to phenylethanol (Shekhawat et al., 2019).

Furthermore, through *in vitro* experiments, it was demonstrated that VOCs such as isoamyl acetate, isoamyl alcohol, 2-phenyl ethylacetate and 2-phenyl ethanol were responsible of the bioprotective activity of *P. kluyveri* on grapes (Mewa-Ngongang et al., 2019b). These authors have also observed that this yeast inhibited the growth of wine spoilage yeasts (*D. bruxellensis* and *D. anomala*) and fruit fungi (*B. cinerea, Colletotrichum acutatum* and *Rhizopus stolonifer* (Mewa-Ngongang et al., 2019a, b). A biocontrol ability on grape berries was also observed for another non-*Saccharomyces* yeast, namely *Starm. bacillaris*. This species is able to survive until the last stages of 440 the fermentation process, contributing significantly to wine quality (Englezos et al., 2017). Its 441 antagonistic activity was investigated against fungal infections such as Alternaria alternata (Prendes 442 et al., 2018) and B. cinerea (Lemos Junior et al., 2016). In the latter study, through in vivo tests, it 443 was found that a co-inoculation of 14 Starm. bacillaris strains in artificially wounded grape berries 444 lead to a decrease of *B. cinerea* infection. A recent study attributed this bioprotection ability to the 445 production of VOCs and in particular to benzyl alcohol (Lemos Junior et al., 2020). In line with this 446 finding, several authors observed an increased amount of this alcohol in mixed fermentations with S. 447 cerevisiae (Englezos et al., 2019a; Binati et al., 2020). Antimicrobial activity against B. cinerea was 448 also observed for *M. pulcherrima* even if in a strain dependent way, due to the production of ethyl 449 alcohol and ethyl acetate (Parafati et al., 2015; Contarino et al., 2019). The inhibitory effect of yeastderived VOCs against postharvest food pathogens was greatly investigated and proposed as an 450 451 effective biocontrol strategy to inhibit the growth of B. cinerea, Colletotrichum acutatum, P. 452 expansum, Penicillium digitatum and Penicillium italicum (Contarino et al., 2019). VOCs could be 453 considered as ideal antimicrobials, since contact between biocontrol agent and food pathogen is not 454 required, to inhibit the growth of the last. Such approach could be useful on postharvest table grapes 455 to improve shelf life and therefore reduce food loss during storage. Further studies are necessary to 456 understand the type and the concentration of VOCs that could be considered crucial in the biocontrol 457 mechanism.

458

### 459 2.2.3. Organic acids

460

461 Organic acids production by the BPCs could play an important role on their dominance over the 462 indigenous microorganisms, increasing the microbial stability of the wine. Bagheri et al. (2018) found 463 that in a consortium of six non-*Saccharomyces* and a *S. cerevisiae* strains, a higher concentration of 464 *Starm. bacillaris* resulted in the disappearance of *Candida parapsilosis* and *M. pulcherrima*. This 465 antagonistic ability was attributed to the capacity to produce high levels of organic acids, increasing 466 the total acidity and reducing pH of the wines. In wine conditions, L. thermotolerans bioprotective 467 ability was investigated in combination with a Lpb. plantarum strain (Rubio-Bretón et al., 2018). The results showed the potential of this inoculation strategy as alternatives to SO<sub>2</sub> addition. The final 468 469 wines had a higher titratable acidity and esters concentration than control wines inoculated with S. 470 cerevisiae. The antimicrobial activity was attributed to the production of high lactic acid amounts by 471 both microorganisms and consequently to the decrease of the pH (Gobbi et al., 2013; Rubio-Bretón 472 et al., 2018; Vilela, 2018). This peculiar feature is of great interest in wine starter cultures, since 473 current climate change in many wine-growing regions may cause a shift in grapes composition, 474 increasing sugar content and decreasing must acidity (Mira de Orduña, 2010). In a recent study, 475 Tremonte et al., (2017) demonstrated that the highest antimicrobial activity of 106 Lpb. plantarum 476 strains was exerted when growing cells were present. It was also highlighted that the use of a cell free 477 supernatant reduced the bioprotective ability of all strains and in particular that the main antagonistic 478 activity was due to the production of organic acids. Interestingly, in this study a relationship between 479 the antagonistic activity and the origin of the strains was underlined, with the wine strains showing 480 the strongest growth inhibition capacity. Generally, biocontrol activity, by the decrease of pH due to 481 the organic acid production could have an impact on wine color stability, due to the enhance of SO<sub>2</sub> 482 properties and the shift of the equilibrium of anthocyanins from the colourless to colored form 483 (Giacosa et al., 2019).

484

# 485 2.2.4. Cell-to-cell contact

486

Among the interaction mechanisms, different researches underlined the ability of yeasts to form cellular aggregates called flocs in different environments, wine included. These flocs can be formed by cells of the same strain, by different strains of the same species or by microorganisms of different species (Rossouw et al., 2015). This physical association increases the metabolic exchange among cells and, consequently, determines the ecological interactions like inhibition or stimulation. It was demonstrated that in a wine environment this cell-to-cell contact mechanism is involved in antagonistic interactions among microorganisms. This phenomenon also called co-aggregation was firstly observed in *S. cerevisiae* against several wine yeasts namely *L. thermotolerans, T. delbrueckii*, *H. uvarum* and *Starm. bacillaris* (Alonso-del-Real et al., 2019; Englezos et al., 2019b; Kemsawasd et al., 2015; Nissen and Arneborg, 2003; Petigonnet et al., 2019; Pietrafesa et al., 2020; Renault et al., 2013; Rossouw et al., 2018; Shekhawat et al., 2019). It was also observed between different strains of *S. cerevisiae* and is involved in dominance (Pérez-Torrado et al., 2018).

499 Co-aggregation is manly studied using a double compartment bioreactor where two species 500 are separated by a membrane that permits the exchange of metabolites (Nissen and Arneborg, 2003), 501 however this system limits the nature and extent of the interactions investigated because it does not 502 permit the immediate and complete transfer of all relevant metabolites and macromolecules (Rossouw 503 et al., 2018).

504 Recently, in S. cerevisiae the genetic basis of this ability was mainly attributed to the 505 expression of FLO family genes, which encode for adhesive proteins on cell wall surface (Rossouw 506 et al., 2018). In fact, transcriptomic analyses highlighted that, during wine and sparkling wine 507 fermentation, S. cerevisiae is achieved in a general cell wall remodelling with the up-regulation of 508 genes involved in asexual cell aggregation (FLO genes) (Shekhawat et al., 2019; Di Gianvito et al., 509 2018). Genetic analyses suggested that non-Saccharomyces yeasts like H. uvarum contain adhesion-510 related domains in their genome (Pu et al., 2014). However, the antagonistic role in these 511 microorganisms is not yet investigated.

512

### 513 **3.** Conclusions and future perspectives

514

515 Success in biological control of unwanted microorganisms requires consideration of the wine 516 fermentation ecology as a balanced ecosystem, where the resident microbiota contribute to enhance 517 wine quality, while maintaining spoilage microorganisms at the lowest levels. Implementing 518 fermentation conditions that favour the population of microorganisms with antimicrobial activity is 519 essential to this purpose. However, the use of chemical additives such as SO<sub>2</sub> may still be needed 520 when the concentration of microbial spoilage metabolites is high, especially in wines with low or 521 moderate ethanol levels. Despite the limitations observed in the different stages, the use of viable 522 cells of antagonist microorganisms well adapted in specific conditions or their derived antimicrobial 523 metabolites are of relevance in winemaking industry, since they could help to reduce or even replace chemical additives. In the same context, the selection of yeasts and bacteria of oenological interest 524 525 with the double role of starter culture and BPCs represents an opportunity for winemakers to enhance wine quality and reduce the production costs, as the demand for sustainable wines will continue to 526 527 grow. A better knowledge of the fermentation conditions and oenological practices which modulate microorganism's performance, will allow a greater management of specific unwanted 528 529 microorganisms during the alcoholic and/or malolactic fermentation. Therefore, studies aiming to 530 further investigate the interaction mechanisms among the starter culture and BPCs and the spoilage 531 microorganisms will be of great value in the near future for this field of research.

532

#### 533 Funding

534

535 This research did not receive any specific grant from funding agencies in the public, commercial, or 536 not-for-profit sectors.

537

### 538 **Declaration of competing interest**

539

540 The authors declare no conflict of interest.

- 542 **References**
- 543

544	Albergaria, H., Arneborg, N., 2016. Dominance of Saccharomyces cerevisiae in alcoholic
545	fermentation processes: role of physiological fitness and microbial interactions. Appl. Microbiol.
546	Biotechnol. 100(5), 2035-2046. http://dx.doi.org/10.1007/s00253-015-7255-0.
547	
548	Alonso-del-Real, J., Pérez-Torrado, R., Querol, A., Barrio, E., 2019. Dominance of wine
549	Saccharomyces cerevisiae strains over Saccharomyces kudriavzevii in industrial fermentation
550	competitions is related to an acceleration of nutrient uptake and utilization. Environ. Microbiol. 21(5),
551	1627-1644. http://dx.doi.org/10.1111/1462-2920.14536.
552	
553	Andorrà, I., Martín, L., Nart, E., Puxeu, M., Hidalgo, C., Ferrer-Gallego, R., 2018. Effect of grape
554	juice composition and nutrient supplementation on the production of sulur dioxide and carboxylic
555	compounds by Saccharomyces cerevisiae. Aust. J. Grape Wine Res. 24(2), 260-266.
556	http://dx.doi.org/10.1111/ajgw.12325.
557	
558	Aponte, M., Blaiotta, G., 2016. Potential role of yeast strains isolated from grapes in the production
559	of Taurasi DOCG. Front. Microbiol. 7, 809. https://doi.org/10.3389/fmicb.2016.00809.
560	
561	Bagheri, B., Zambelli, P., Vigentini, I., Bauer, F. F., Setati, M. E., 2018. Investigating the effect of
562	selected non-Saccharomyces species on wine ecosystem function and major volatiles. Front. Bioeng.

565 Bahar, A. A., Ren, D., 2013. Antimicrobial peptides. Pharmaceuticals. 6(12), 1543-1575.
566 https://doi.org/10.3390/ph6121543.

Biotechnol. 6, 169. https://doi.org/10.3389/fbioe.2018.00169.

568	Balmaseda, A., Bordons, A., Reguant, C., Bautista-Gallego, J., 2018. Non-Saccharomyces in wine:
569	effect upon Oenococcus oeni and malolactic fermentation. Front. Microbiol. 9, 534.
570	https://doi.org/10.3389/fmicb.2018.00534.
571	
572	Barbosa, C., Mendes-Faia, A., Lage, P., Mira, N. P., Mendes-Ferreira, A., 2015. Genomic expression
573	program of Saccharomyces cerevisiae along a mixed-culture wine fermentation with Hanseniaspora
574	guilliermondii. Microb. Cell Factories. 14(1), 124. https://doi.org/10.1186/s12934-015-0318-1.
575	
576	Bartle, L., Sumby, K., Sundstrom, J., Jiranek, V., 2019. The microbial challenge of winemaking:
577	yeast-bacteria compatibility. FEMS Yeast Res. 19(4), foz040. https://doi.org/10.1093/femsyr/foz040.
578	
579	Bastard, A., Coelho, C., Briandet, R., Canette, A., Gougeon, R., Alexandre, H., Guzzo, J., Weidmann,
580	S., 2016. Effect of biofilm formation by Oenococcus oeni on malolactic fermentation and the release
581	of aromatic compounds in wine. Front. Microbiol. 7, 613. https://doi.org/10.3389/fmicb.2016.00613.
582	
583	Bauer, M. A., Kainz, K., Carmona-Gutierrez, D., Madeo, F., 2018. Microbial wars: competition in
584	ecological niches and within the microbiome. Microb. Cell. 5(5), 215.
585	https://doi.org/10.15698/mic2018.05.628.
586	
587	Belda, I., Ruiz, J., Alonso, A., Marquina, D., Santos, A., 2017. The biology of Pichia
588	membranifaciens killer toxins. Toxins. 9(4), 112. https://doi.org/10.3390/toxins9040112.
589	
590	Benito, S., 2018a. The impact of Torulaspora delbrueckii yeast in winemaking. Appl. Microbiol.
591	Biotechnol. 102(7), 3081-3094. https://doi.org/10.1007/s00253-018-8849-0.
592	

Benito, S., 2018b. The impacts of *Lachancea thermotolerans* yeast strains on winemaking Appl.
Microbiol. Biotechnol. 102 (16), 6775-6790. https://doi.org/10.1007/s00253-018-9117-z.

595

Binati, R. L., Junior, W. J. L., Luzzini, G., Slaghenaufi, D., Ugliano, M., Torriani, S., 2020.
Contribution of non-*Saccharomyces* yeasts to wine volatile and sensory diversity: A study on *Lachancea thermotolerans, Metschnikowia* spp. and *Starmerella bacillaris* strains isolated in Italy.
Int. J. Food Microbiol. 318, 108470. <u>https://doi.org/10.1016/j.ijfoodmicro.2019.108470</u>.
Bisson, L. F., 1999. Stuck and sluggish fermentations. Am. J. Enol. Vitic. 50, 107-119.

602

- Boynton, P. J., Peterson, C. N., Pringle, A., 2019. Superior dispersal ability can lead to persistent
  ecological dominance throughout succession. Appl. Environ. Microbiol. 85(6), e02421-18.
  https://doi.org/10.1128/AEM .02421-18.
- 606
- Bozoudi, D., Tsaltas, D., 2018. The multiple and versatile roles of *Aureobasidium pullulans* in the
  vitivinicultural sector. Fermentation. 4(4), 85. https://doi.org/10.3390/fermentation4040085.

- Branco, P., Francisco, D., Chambon, C., Hébraud, M., Arneborg, N., Almeida, M. G., Caldeira, J.,
  Albergaria, H., 2014. Identification of novel GAPDH-derived antimicrobial peptides secreted by *Saccharomyces cerevisiae* and involved in wine microbial interactions. Appl. Microbiol. Biotechnol.
  98, 843-853. https://doi.org/10.1007/s00253-013-5411-y.
- 614
- Branco, P., Francisco, D., Monteiro, M., Almeida, M. G., Caldeira, J., Arneborg, N., Prista, C.,
  Albergaria, H., 2017. Antimicrobial properties and death-inducing mechanisms of saccharomycin, a
  biocide secreted by *Saccharomyces cerevisiae*. Appl. Microbiol. Biotechnol. 101, 159-171.
- 618 https://doi.org/10.1007/s00253-016-7755-6.

620	Brizuela, N., Tymczyszyn, E. E., Semorile, L. C., La Hens, D. V., Delfederico, L., Hollmann, A.,
621	Bravo-Ferrada, B., 2019. Lactobacillus plantarum as a malolactic starter culture in winemaking: A
622	new (old) player?. Electron. J. Biotechnol. 38, 10-18. https://doi.org/10.1016/j.ejbt.2018.12.002.
623	
624	Carbó, A., Torres, R., Teixidó, N., Usall, J., Magan, N., Medina, A., 2018. Predicted ecological niches
625	and environmental resilience of different formulations of the biocontrol yeast Candida sake CPA-1
626	using the Bioscreen C. BioControl. 63(6), 855-866. https://doi.org/10.1007/s10526-018-09910-4.
627	
628	Comitini, F., Ferretti, R., Clementi, F., Mannuzzu, I., Ciani, M., 2005. Interactions between
629	Saccharomyces cerevisiae and malolactic bacteria: preliminary characterization of a yeast
630	proteinaceous compound(s) active against Oenococcus oeni. J. Appl. Microbiol. 99, 105-111.
631	doi:10.1111/j.1365-2672.2005.02579.x.
632	
633	Contarino, R., Brighina, S., Fallico, B., Cirvilleri, G., Parafati, L., Restuccia, C., 2019. Volatile
634	organic compounds (VOCs) produced by biocontrol yeasts. Food Microbiol. 82, 70-74.
635	https://doi.org/10.1016/j.fm.2019.01.008.
636	
637	Çorbacı, C., Uçar, F. B., 2018. Purification, characterization and in vivo biocontrol efficiency of killer
638	toxins from Debaryomyces hansenii strains. Int. J. Biol. Macromol. 119, 1077-1082.
639	https://doi.org/10.1016/j.ijbiomac.2018.07.121.
640	
641	Cordero-Bueso, G., Mangieri, N., Maghradze, D., Foschino, R., Valdetara, F., Cantoral, J. M.,
642	Vigentini, I., 2017. Wild grape-associated yeasts as promising biocontrol agents against Vitis vinifera

fungal pathogens. Front. Microbiol. 8, 2025. https://doi.org/10.3389/fmicb.2017.02025.

645	Curiel, J. A., Morales, P., Gonzalez, R., Tronchoni, J., 2017. Different non-Saccharomyces yeast
646	species stimulate nutrient consumption in S. cerevisiae mixed cultures. Front. Microbiol. 8, 2121.
647	https://doi.org/10.3389/fmicb.2017.02121.
648	
649	de Ullivarri, M. F., Mendoza, L. M., Raya, R. R., 2014. Killer activity of Saccharomyces cerevisiae
650	strains: partial characterization and strategies to improve the biocontrol efficacy in winemaking.
651	Antonie Leeuwenhoek. 106(5), 865-878. https://doi.org/10.1007/s10482-014-0256-7.
652	
653	Di Gianvito, P., Arfelli, G., Suzzi, G., Tofalo, R., 2019. New trends in sparkling wine production:
654	yeast rational selection. In Alcoholic Beverages, Woodhead Publishing., pp. 347-386.
655	
656	Di Gianvito, P., Tesnière, C., Suzzi, G., Blondin, B., Tofalo, R., 2018. Different genetic responses to
657	oenological conditions between a flocculent wine yeast and its FLO5 deleted strain: insights from the
658	transcriptome. Food Res. Int. 114, 178-186. https://doi.org/10.1016/j.foodres.2018.07.061.
659	
660	Dimopoulou, M., Bardeau, T., Ramonet, P.Y., Miot-Certier, C., Claisse, O., Doco, T., Petrel, M.,
661	Lucas, P., Dols-Lafargue, M., 2015. Exopolysaccharides produced by Oenococcus oeni: from
662	genomic and phenotypic analysis to technological valorization. Food Microbiol. 53, 10-17.
663	https://doi.org/10.1016/j.fm.2015.07.011.
664	
665	Dimopoulou, M., Raffenne, J., Claisse, O., Miot-Sertier, C., Iturmendi, N., Moine, V., Coulon, J.,
666	Dols-Lafargue, M., 2018. Oenococcus oeni exopolysaccharide biosynthesis, a tool to improve
667	malolactic starter performance. Front. Microbiol. 9, 1276. https://doi.org/10.3389/fmicb.2018.01276.
668	
669	Dokka, A., kotilinga Reddy, Y., Spoorthy, G. S., Reddy, I. S., 2018. Protective cultures-a review. Int.
670	J. Curr. Microbiol. App. Sci. 7(6), 228-238. https://doi.org/10.20546/ijcmas.2018.706.028.

- Droby, S., Wisniewski, M., Teixidó, N., Spadaro, D., Jijakli, M. H., 2016. The science, development,
  and commercialization of postharvest biocontrol products. Postharvest Biol. Technol. 122, 22-29.
  https://doi.org/10.1016/j.postharvbio.2016.04.006.
- 675
- Edison, L. K., Shiburaj, S., Pradeep, N. S., 2018. Microbial beta glucanase in agriculture. In: Kumar,
  P., Patra, K. J., Chandra, P. (Ed.), Advances in Microbial Biotechnology. Apple Academic Press. P.,
  New York, pp. 53-72.
- 679
- Englezos, V., Cravero, F., Torchio, F., Rantsiou, K., Ortiz-Julien, A., Lambri, M., Gerbi, V., Rolle,
  L., Cocolin, L., 2018a. Oxygen availability and strain combination modulate yeast growth dynamics
  in mixed culture fermentations of grape must with *Starmerella bacillaris* and *Saccharomyces cerevisiae*. Food Microbiol. 69, 179-188. https://doi.org/10.1016/j.fm.2017.08.007.
- 684
- Englezos, V., Giacosa, S., Rantsiou, K., Rolle, L., Cocolin, L., 2017. *Starmerella bacillaris* in
  winemaking: opportunities and risks. Curr. Opin. Food Sci. 17, 30-35.
  https://doi.org/10.1016/j.cofs.2017.08.007.
- 688
- Englezos, V., Pollon, M., Rantsiou, K., Ortiz-Julien, A., Botto, R., Rio Segade, S., Giacosa S., Rolle
  L., Cocolin, L., 2019a. *Saccharomyces cerevisiae-Starmerella bacillaris* strains interaction
  modulates chemical and volatile profile in red wine mixed fermentations. Food Res. Int. 122, 392401. https://doi.org/10.1016/j.foodres.2019.03.072.
- 693
- Englezos, V., Rantsiou, K., Giacosa, S., Rio Segade, S., Rolle, L., Cocolin, L., 2019b. Cell-to-cell
  contact mechanism modulates *Starmerella bacillaris* death in mixed culture fermentations with

696Saccharomycescerevisiae.Int.J.FoodMicrobiol.289,106-114.697https://doi.org/10.1016/j.ijfoodmicro.2018.09.009.

698

Farbo, M. G., Urgeghe, P. P., Fiori, S., Marcello, A., Oggiano, S., Balmas, V., Hassan, Z. U. I., Jaoua,
S., Migheli, Q., 2018. Effect of yeast volatile organic compounds on ochratoxin A-producing *Aspergillus carbonarius* and *Aspergillus ochraceus*. Int. J. Food Microbiol. 284, 1-10.
https://doi.org/10.1016/j.ijfoodmicro.2018.06.023.

703

Ferraz, P., Cássio, F., Lucas, C., 2019. Potential of yeasts as biocontrol agents of the phytopathogen
causing cacao Witches' Broom Disease: is microbial warfare a solution?. Front. Microbiol. 10, 1766.
https://doi.org/10.3389/fmicb.2019.01766.

707

- 708 Fiori, S., Urgeghe, P. P., Hammami, W., Razzu, S., Jaoua, S., Migheli, Q., 2014. Biocontrol activity 709 of four non-and low-fermenting yeast strains against Aspergillus carbonarius and their ability to 710 remove ochratoxin A from grape juice. Int. J. Food Microbiol. 189, 45-50. 711 https://doi.org/10.1016/j.ijfoodmicro.2014.07.020.
- 712
- Gaggia, F., Di Gioia, D., Baffoni, L., Biavati, B., 2011. The role of protective and probiotic cultures
  in food and feed and their impact in food safety. Trends Food Sci. Technol. 22, S58-S66.
  <u>https://doi.org/10.1016/j.tifs.2011.03.003</u>.
- 716
- Gao, C., Fleet, G. H., 1988. The effects of temperature and pH on the ethanol tolerance of the wine
  yeasts, *Saccharomyces cerevisiae*, *Candida stellate* and *Kloeckera apiculata*. J. Appl. Bacteriol. 65,
  405-410. https://doi.org/10.1111/j.1365-2672.1988.tb01909.x.

- Garrido, C. C., Usall, J., Torres, R., Teixidó, N., 2017. Effective control of *Botrytis* bunch rot in
  commercial vineyards by large-scale application of *Candida sake* CPA-1. BioControl. 62, 161-173.
  https://doi.org/10.1007/s10526-017-9789-9.
- 724
- Ghoul, M., Mitri, S., 2016. The ecology and evolution of microbial competition. Trends Microbiol.
  24(10), 833-845. https://doi.org/10.1016/j.tim.2016.06.011.
- 727
- Giacosa, S., Río Segade, S., Cagnasso, E., Caudana, A., Rolle, L., Gerbi, V., 2019. SO<sub>2</sub> in wines:
  rational use and possible alternatives. In: Morata, A. (Ed.), Red Wine Technology. Academic Press,
  Cambridge, MA, USA, pp. 309-321.
- 731
- Gobbi, M., Comitini, F., Domizio, P., Romani, C., Lencioni, L., Mannazzu, I., Ciani, M., 2013. *Lachancea thermotolerans* and *Saccharomyces cerevisiae* in simultaneous and sequential cofermentation: a strategy to enhance acidity and improve the overall quality of wine. Food Microbiol.
  33(2), 271-281. https://doi.org/10.1016/j.fm.2012.10.004.
- 736
- González, B., Vázquez, J., Morcillo-Parra, M. Á., Mas, A., Torija, M. J., Beltran, G., 2018. The
  production of aromatic alcohols in non-*Saccharomyces* wine yeast is modulated by nutrient
  availability. Food Microbiol. 74, 64-74. https://doi.org/10.1016/j.fm.2018.03.003.
- 740
- Gore-Lloyd, D., Sumann, I., Brachmann, A. O., Schneeberger, K., Ortiz-Merino, R. A., MorenoBeltrán, M., Schläfli, M., Kirner, P., Santos Kron, A., Rueda-Mejia, M. P., Somerville, V., Wolfe, K.
  H., Piel, J., Ahrens, C. H., Henk, D., Freimoser, F. M., 2019. Snf2 controls pulcherriminic acid
  biosynthesis and antifungal activity of the biocontrol yeast *Metschnikowia pulcherrima*. Molecular
  Microbiol. 112(1), 317-332. https://doi.org/10.1111/mmi.14272.
- 746

- Guerrero, R. F., Cantos-Villar, E., 2015. Demonstrating the efficiency of sulphur dioxide
  replacements in wine: A parameter review. Trends Food Sci Technol. 42(1), 27-43.
  https://doi.org/10.1016/j.tifs.2014.11.004.
- 750
- 751 Jacometti, M. A., Wratten, S. D., Walter, M., 2010. Alternatives to synthetic fungicides for Botrytis 752 cinerea management in vineyards. Aust. J. Grape Wine Res. 154-172. 16(1), 753 https://doi.org/10.1111/j.1755-0238.2009.0067.x.
- 754
- Jolly, N. P., Varela, C., Pretorius, I. S., 2014. Not your ordinary yeast: non-*Saccharomyces* yeasts in
  wine production uncovered. FEMS Yeast Res. 14(2), 215-237. <u>https://doi.org/10.1111/1567-</u>
  <u>1364.12111</u>.
- 758
- Kántor, A., Hutková, J., Petrová, J., Hleba, L., Kac<sup>\*</sup>ániová, M., 2015. Antimicrobial activity of
  pulcherrimin pigment produced by *Metschnikowia pulcherrima* against various yeast species. J.
  Microbiol. Biotechnol. Food Sci. 5, 282-285.
- 762
- Kemsawasd, V., Branco, P., Almeida, M. G., Caldeira, J., Albergaria, H., Arneborg, N., 2015. Cellto-cell contact and antimicrobial peptides play a combined role in the death of *Lachanchea thermotolerans* during mixed-culture alcoholic fermentation with *Saccharomyces cerevisiae*. FEMS
  Microbiol. Lett. 362(14), fnv103. https://doi.org/10.1093/femsle/fnv103.
- 767
- Keswani, C., Singh, H. B., Hermosa, R., García-Estrada, C., Caradus, J., He, Y. W., MezaacheAichour, S., Glare, T. R., Borriss, R., Vinale, F., Sansinenea, E., 2019. Antimicrobial secondary
  metabolites from agriculturally important fungi as next biocontrol agents. Appl. Microbiol.
  Biotechnol. 103(23-24), 9287-9303. https://doi.org/10.1007/s00253-019-10209-2103.
- 772

773	Knoll	, C., Divol, B., Du Toit, M., 2	2008. Geneti	c screening	of lactic acid bact	eria of oenol	ogical origin
774	for	bacteriocin-encoding	genes.	Food	Microbiol.	25(8),	983-991.
775	https:/	//doi.org/10.1016/j.fm.2008.	.06.010.				

Kosel, J., Čadež, N., Schuller, D., Carreto, L., Franco-Duarte, R., Raspor, P., 2017. The influence of *Dekkera bruxellensis* on the transcriptome of *Saccharomyces cerevisiae* and on the aromatic profile
of synthetic wine must. FEMS Yeast Res. 17(4), fox018. https://doi.org/10.1093/femsyr/fox018.

Krieger-Weber, S., Heras, J. M., Suarez, C., 2020. *Lactobacillus plantarum*, a new biological tool to
control malolactic fermentation: a review and an outlook. Beverages. 6(2), 23.
https://doi.org/10.3390/beverages6020023.

784

Kuchen, B., Maturano, Y. P., Mestre, M. V., Combina, M., Toro, M. E., Vazquez, F., 2019. Selection
of native non-*Saccharomyces* yeasts with biocontrol activity against spoilage yeasts in order to
produce healthy regional wines. Fermentation, 5(3), 60.
https://doi.org/10.3390/fermentation5030060.

789

Labbani, F. Z. K., Turchetti, B., Bennamoun, L., Dakhmouche, S., Roberti, R., Corazzi, L., Meraihi,
Z., Buzzini, P., 2015. A novel killer protein from *Pichia kluyveri* isolated from an Algerian soil:
purification and characterization of its *in vitro* activity against food and beverage spoilage yeasts.
Antonie Leeuwenhoek. 107(4), 961-970. https://doi.org/10.1007/s10482-015-0388-4.

794

Lachance, M. A., 2016. *Metschnikowia*: half tetrads, a regicide and the fountain of youth. Yeast.
33(11), 563-574. https://doi.org/ 10.1002/yea.3208.

798	Lasik-Kurdyś, M., Sip, A., 2019. Evaluation of the antimicrobial activity of bacteriocin-like
799	inhibitory substances of enological importance produced by Oenococcus oeni isolated from wine.
800	Eur. Food Res. Technol. 245(2), 375-382. https://doi.org/10.1007/s00217-018-3169-2.

Lemos Junior, W. J., Fernandes, W. J., Bovo, B., Nadai, C., Crosato, G., Carlot, M., Favaron, F.,
Giacomini A., Corich, V., 2016. Biocontrol ability and action mechanism of *Starmerella bacillaris*(synonym *Candida zemplinina*) isolated from wine musts against gray mold disease agent *Botrytis cinerea* on grape and their effects on alcoholic fermentation. Front Microbiol. 7, 1249.
https://doi.org/10.3389/fmicb.2016.01499.

807

808 Lemos Junior, W. J., Binati, R. L., Felis, G. E., Slaghenaufi, D., Ugliano, M., Torriani, S., 2020.

809 Volatile organic compounds from *Starmerella bacillaris* to control gray mold on apples and modulate

810 cider aroma profile. Food Microbiol. 89, 103446. https://doi.org/10.1016/j.fm.2020.103446.

811

Liu, G. L., Chi, Z., Wang, G. Y., Wang, Z. P., Li, Y., Chi, Z. M., 2015. Yeast killer toxins, molecular
mechanisms of their action and their applications. Crit. Rev. Biotechnol. 35(2), 222-234.
https://doi.org/10.3109/07388551.2013.833582.

815

Liu, Y., Rousseaux, S., Tourdot-Maréchal, R., Sadoudi, M., Gougeon, R., Schmitt-Kopplin, P.,
Alexandre, H., 2017. Wine microbiome: a dynamic world of microbial interactions. Crit. Rev. Food
Sci. Nutr. 57(4), 856-873. https://doi.org/10.1080/10408398.2014.983591.

819

Lombardi, S. J., De Leonardis, A., Lustrato, G., Testa, B., Iorizzo, M., 2016. Yeast autolysis in
sparkling wine aging: use of killer and sensitive *Saccharomyces cerevisiae* strains in co-culture.
Recent Pat. Biotechnol. 9, 223-230.

824	Lopes, C.A., Sangorrín, M.P., 2010. Optimization of killer assays for yeast selection protocols. Rev.
825	Argent. Microbiol. 42, 298-306. https://doi.org/10.1590/S0325-75412010000400011.
826	
827	López-Seijas, J., García-Fraga, B., da Silva, A. F., Sieiro, C., 2020. Wine Lactic Acid Bacteria with
828	antimicrobial activity as potential biocontrol agents against Fusarium oxysporum f. sp. lycopersici.
829	Agronomy. 10(1), 31. https://doi.org/10.3390/agronomy10010031.
830	
831	Mazzucco, M. B., Ganga, M. A., Sangorrín, M. P., 2019. Production of a novel killer toxin from
832	Saccharomyces eubayanus using agro-industrial waste and its application against wine spoilage
833	yeasts. Antonie Leeuwenhoek. 112(7), 965-973. https://doi.org/10.1007/s10482-019-01231-5.
834	
835	Mannazzu, I., Domizio, P., Carboni, G., Zara, S., Zara, G., Comitini, F., Budroni, M., Ciani, M., 2019.
836	Yeast killer toxins: from ecological significance to application, Crit. Rev. Biotechnol. 35 (5), 603-
837	617. https://doi.org/10.1080/07388551.2019.1601679.
838	
839	Mehlomakulu, N. N., Prior, K. J., Setati, M. E., Divol, B., 2017. Candida pyralidae killer toxin
840	disrupts the cell wall of Brettanomyces bruxellensis in red grape juice. J. Appl. Microbiol. 122(3),
841	747-758. https://doi.org/10.1111/jam.13383.
842	
843	Mehlomakulu, N. N., Setati, M. E., Divol, B., 2015. Non-Saccharomyces killer toxins: possible
844	biocontrol agents against Brettanomyces in wine?. S. Afr. J. Enol. Vitic. 36(1), 94-104.
845	
846	Melkonian, C., Gottstein, W., Blasche, S., Kim, Y., Abel-Kistrup, M., Swiegers, H., Saerens, S.,
847	Edwards, N., Patil, K. R., Teusink, B., Molenaar, D., 2019. Finding functional differences between
848	species in a microbial community: case studies in wine fermentation and Kefir culture. Front.
849	Microbiol. 10, 1347. https://doi.org/10.3389/fmicb.2019.01347.
	33

851	Melvydas, V., Svediene, J., Skridlaite, G., Vaiciuniene, J., Garjonyte, R., 2020. In vitro inhibition of
852	Saccharomyces cerevisiae growth by Metschnikowia spp. triggered by fast removal of iron via two
853	ways. Braz. J. Microbiol., 51(4), 1953-1964. https://doi.org/10.1007/s42770-020-00357-3.
854	
855	Mendoza, L. M., de Nadra, M. C. M., Farías, M. E., 2010. Antagonistic interaction between yeasts
856	and lactic acid bacteria of oenological relevance: partial characterization of inhibitory compounds
857	produced by yeasts. Food Res. Int. 43(8), 1990-1998. https://doi.org/10.1016/j.foodres.2010.05.017.
858	
859	Mewa-Ngongang, M., du Plessis, H. W., Ntwampe, S. K. O., Chidi, B. S., Hutchinson, U. F., Mekuto,
860	L., Jolly, N. P., 2019a. The use of Candida pyralidae and Pichia kluyveri to control spoilage
861	microorganisms of raw fruits used for beverage production. Foods. 8(10), 454.
862	https://doi.org/10.3390/foods8100454.
863	
864	Mewa-Ngongang, M., du Plessis, H. W., Ntwampe, S. K., Chidi, B. S., Hutchinson, U. F., Mekuto,
865	L., Jolly, N. P., 2019b. Grape pomace extracts as fermentation medium for the production of potential
866	biopreservation compounds. Foods. 8(2), 51. https://doi.org/10.3390/foods8020051.
867	
868	Mira de Orduña R., 2010. Climate change associated effects on grape and wine quality and
869	production. Food Res. Int. 43:1844-1855. doi:10. 1016/j.foodres.2010.05.001.
870	
871	Mondello, V., Songy, A., Battiston, E., Pinto, C., Coppin, C., Trotel-Aziz, P., Clemént, C., Mugnai,
872	L., Fontaine, F., 2018. Grapevine trunk diseases: a review of fifteen years of trials for their control
873	with chemicals and biocontrol agents. Plant Dis. 102(7), 1189-1217. https://doi.org/10.1094/PDIS-
874	08-17-1181-FE.

- Morata, A., Loira, I., Tesfaye, W., Bañuelos, M. A., González, C., Suárez Lepe, J. A., 2018. *Lachancea thermotolerans* applications in wine technology. Fermentation. 4(3), 53.
  https://doi.org/10.3390/fermentation4030053.
- 879
- Morata, A., Loira, I., Escott, C., del Fresno, J. M., Bañuelos, M. A., Suárez-Lepe, J. A., 2019a.
  Applications of *Metschnikowia pulcherrima* in Wine Biotechnology. Fermentation. 5(3), 63.
  https://doi.org/10.3390/fermentation5030063.
- 883
- Morata, A., 2019b. Enological Repercussions of Non-*Saccharomyces* Species in Wine
  Biotechnology. Fermentation. 5, 72. https://doi.org/10.3390/fermentation5030072.
- 886
- Mukherjee, A., Verma, J. P., Gaurav, A. K., Chouhan, G. K., Patel, J. S., Hesham, A. E. L., 2020.
  Yeast a potential bio-agent: future for plant growth and postharvest disease management for
  sustainable agriculture. Appl. Microbiol. Biotechnol. 104, 1497-1510.
  https://doi.org/10.1007/s00253-019-10321-3.
- 891
- 892
- Nally, M. C., Ponsone, M. L., Pesce, V. M., Toro, M. E., Vazquez, F., Chulze, S., 2018. Evaluation
  of behaviour of *Lachancea thermotolerans* biocontrol agents on grape fermentations. Lett. Appl.
  Microbiol. 67(1), 89-96. https://doi.org/10.1111/lam.13001.
- 896
- Nardi, T., 2020. Microbial resources as a tool for enhancing sustainability in winemaking.
  Microorganisms. 8(4), 507. https://doi.org/10.3390/microorganisms8040507.
- 899

- Ndlovu, B., Schoeman, H., Franz, C. M. A. P., Du Toit, M., 2015. Screening, identification and
  characterization of bacteriocins produced by wine-isolated LAB strains. J. Appl. Microbiol. 118(4),
  1007-1022. https://doi.org/10.1111/jam.12752.
- 903
- Nehme, N., Mathieu, F., Taillandier, P., 2010. Impact of the co-culture of *Saccharomyces cerevisiae Oenococcus oeni* on malolactic fermentation and partial characterization of a yeast-derived
  inhibitory peptidic fraction. Food Microbiol. 27, 150-57. https://doi.org/10.1016/j.fm.2009.09.008.
- 908 Nel, H. A., Bauer, R., Wolfaardt, G. M., Dicks, L. M. T., 2002. Effect of bacteriocins pediocin PD-
- 909 1, plantaricin 423, and nisin on biofilms of *Oenococcus oeni* on a stainless steel surface. Am J Enol
  910 Vitic. 53, 191-196.
- 911
- Nissen, P., Arneborg, N., 2003. Characterization of early deaths of non-*Saccharomyces* yeasts in
  mixed cultures with *Saccharomyces cerevisiae*. Arch. Microbiol. 180(4), 257-263.
  https://doi.org/10.1007/s00203-003-0585-9.
- 915
- Oro, L., Ciani, M., Comitini, F., 2014. Antimicrobial activity of *Metschnikowia pulcherrima* on wine
  yeasts. J. Appl. Microbiol. 116, 1209-1217. https://doi.org/10.1111/jam.12446.
- 918
- Oro, L., Ciani, M., Bizzaro, D., Comitini, F., 2016. Evaluation of damage induced by Kwkt and Pikt
  zymocins against *Brettanomyces/Dekkera* spoilage yeast, as compared to sulphur dioxide. J. Appl.
  Microbiol. 121(1), 207-214. https://doi.org/10.1111/jam.13121.
- 922
- 923 Osborne, J. P., Edwards, C. G., 2007. Inhibition of malolactic fermentation by a peptide produced by
- 924 Saccharomyces cerevisiae during alcoholic fermentation. Int. J. Food Microbiol. 118, 27-34.
- 925 https://doi.org/10.1016/j. ijfoodmicro.2007.05.007.

927	Pandin, C., Le Coq, D., Canette, A., Aymerich, S., Briandet, R., 2017. Should the biofilm mode of
928	life be taken into consideration for microbial biocontrol agents?. Microb. Biotechnol. 10(4), 719-734.
929	https://doi.org/10.1111/1751-7915.12693.
930	
931	Parafati, L., Vitale, A., Restuccia, C., Cirvilleri, G., 2015. Biocontrol ability and action mechanism
932	of food-isolated yeast strains against Botrytis cinerea causing post-harvest bunch rot of table grape.
933	Food Microbiol. 47, 85-92. https://doi.org/10.1016/j.fm.2014.11.013.
934	
935	Parapouli, M., Vasileiadis, A., Afendra, A. S., Hatziloukas, E., 2020. Saccharomyces cerevisiae and
936	its industrial applications. AIMS microbial. 6(1), 1-31. https://doi.org/10.3934/microbiol.2020001.
937	
938	Parijs, I., Steenackers, H. P., 2018. Competitive inter-species interactions underlie the increased
939	antimicrobial tolerance in multispecies brewery biofilms. ISME J. 12(8), 2061-2075.
940	https://doi.org/10.1038/s41396-018-0146-5.
941	
942	Pawlikowska, E., James, S. A., Breierova, E., Antolak, H., Kregiel, D., 2019. Biocontrol capability
943	of local Metschnikowia sp. isolates. Antonie Leeuwenhoek. 112(10), 1425-1445.
944	https://doi.org/10.1007/s10482-019-01272-w.
945	
946	Pawlowska, A. M., Zannini, E., Coffey, A., Arendt, E. K., 2012. "Green preservatives": combating
947	fungi in the food and feed industry by applying antifungal lactic acid bacteria. In Jeyakumar H. (Ed.),
948	Advances in food and nutrition research. Academic Press, New York, Vol. 66, pp. 217-238.
949	

950	Peña, R., Gan	iga, M.	A., 2019	. No	ovel antimicrobi	ial pep	tides produced l	oy Candida	intermedia
951	LAMAP1790	active	against	the	wine-spoilage	yeast	Brettanomyces	bruxellensis	. Antonie
952	Leeuwenhoek.	112(2),	297-304.	https	s://doi.org/10.10	007/s10	482-018-1159-9.		

Peng, C., Andersen, B., Arshid, S., Larsen, M. R., Albergaria, H., Lametsch, R., Arneborg, N., 2019.
Proteomics insights into the responses of *Saccharomyces cerevisiae* during mixed-culture alcoholic
fermentation with *Lachancea thermotolerans*. FEMS Microbiol. Ecol. 95(9), fiz126.
https://doi.org/10.1093/femsec/fiz126.

958

959 Pérez-Nevado, F., Albergaria, H., Hogg, T., Gírio, F., 2006. Cellular death of two non960 Saccharomyces wine-related yeasts during mixed fermentations with Saccharomyces cerevisiae. Int.
961 J. Food Microbiol. 108, 336-345. https://doi.org/10.1016/j.ijfoodmicro.2005.12.012.

962

963 Pérez-Torrado, R., Rantsiou, K., Perrone, B., Navarro-Tapia, E., Querol, A., Cocolin, L., 2017.
964 Ecological interactions among *Saccharomyces cerevisiae* strains: insight into the dominance
965 phenomenon. Sci. Rep. 7, 43603. https://doi.org/10.1038/srep43603.

966

- 967 Petitgonnet, C., Klein, G. L., Roullier-Gall, C., Schmitt-Kopplin, P., Quintanilla-Casas, B., Vichi, S.,
  968 Julien-Davi, D., Alexandre, H., 2019. Influence of cell-cell contact between *L. thermotolerans* and *S.*969 *cerevisiae* on yeast interactions and the exo-metabolome. Food Microbiol. 83, 122-133.
  970 https://doi.org/10.1016/j.fm.2019.05.005.
- 971
- Pietrafesa, A., Capece, A., Pietrafesa, R., Bely, M., Romano, P., 2020. *Saccharomyces cerevisiae* and *Hanseniaspora uvarum* mixed starter cultures: influence of microbial/physical interactions on wine
  characteristics. Yeast. 37(11), 609-621. https://doi.org/10.1002/yea.3506.

- 976 Porter, T. J., Divol, B., Setati, M. E., 2019. *Lachancea* yeast species: Origin, biochemical
  977 characteristics and oenological significance. Food Res. Int. 119, 378-389.
  978 https://doi.org/10.1016/j.foodres.2019.02.003.
- 979
- Prendes, L. P., Merín, M. G., Fontana, A. R., Bottini, R. A., Ramirez, M. L., de Ambrosini, V. I. M.,
  2018. Isolation, identification and selection of antagonistic yeast against *Alternaria alternata*infection and tenuazonic acid production in wine grapes from Argentina. Int. J. Food Microbiol. 266,
  14-20. https://doi.org/10.1016/j.ijfoodmicro.2017.10.033.
- 984
- Prior, K. J., Bauer, F. F., Divol, B., 2019. The utilisation of nitrogenous compound by commercial
  non-*Saccharomyces* yeasts associated with wine. Food Microbiol. 79, 75-84.
  https://doi.org/10.1016/j.fm.2018.12.002.
- 988
- Pu, L., Jingfan, F., Kai, C., Chao-an, L., Yunjiang, C., 2014. Phenylethanol promotes adhesion and
  biofilm formation of the antagonistic yeast *Kloeckera apiculata* for the control of blue mold on citrus.
  FEMS Yeast Res. 14(4), 536-546. https://doi.org/10.1111/1567-1364.12139.
- 992
- Rauhut, D., 2017. Usage and formation of sulphur compounds. In Biology of Microorganisms on
  Grapes, in Must and in Wine, Springer, Cham., pp. 255-291..
- 995
- Renault, P. E., Albertin, W., Bely, M., 2013. An innovative tool reveals interaction mechanisms
  among yeast populations under oenological conditions. Appl. Microbiol. Biotechnol. 97, 4105-4119.
  https://doi.org/10.1007/s00253-012-4660-5.
- 999

- 1000 Restuccia, D., Loizzo, M. R, Spizzirri, U. G., 2018. Accumulation of biogenic amines in wine: role
  1001 of alcoholic and malolactic fermentation. Ferment. 4(1), 6.
  1002 https://doi.org/10.3390/fermentation4010006.
- 1003
- Rizk, Z., El Rayess, Y., Ghanem, C., Mathieu, F., Taillandier, P., Nehme, N., 2016. Impact of
  inhibitory peptides released by *Saccharomyces cerevisiae* BDX on the malolactic fermentation
  performed by *Oenococcus oeni* Vitilactic F. Int. J. Food Microbiol. 233, 90-96.
  https://doi.org/10.1016/j.ijfoodmicro.2016.06.018.
- 1008
- Roca-Mesa, H., Sendra, S., Mas, A., Beltran, G., Torija, M. J., 2020. Nitrogen preferences during
  alcoholic fermentation of different non-*Saccharomyces* yeasts of oenological interest.
  Microorganisms, 8(2), 157. https://doi.org/10.3390/microorganisms8020157.
- 1012
- 1013 Rojo-Bezares, B., Saenz, Y., Zarazaga, M., Torres, C., Ruiz-Larrea, F., 2007. Antimicrobial activity
  1014 of nisin against *Oenococcus oeni* and other wine bacteria. Int. J. Food Microbiol. 116(1):32-36.
  1015 https://doi.org/10.1016/j.ijfoodmicro.2006.12.020.
- 1016
- Rollero, S., Bloem, A., Ortiz-Julien, A., Camarasa, C., Divol, B., 2018. Altered fermentation
  performances, growth, and metabolic footprints reveal competition for nutrients between yeast
  species inoculated in synthetic grape juice-like medium. Front. Microbiol. 9, 196.
  https://doi.org/10.3389/fmicb.2018.00196.
- 1021
- 1022 Rossouw, D., Bagheri, B., Setati, M. E., Bauer, F. F., 2015. Co-flocculation of yeast species, a new
  1023 mechanism to govern population dynamics in microbial ecosystems. PloS one, 10(8).
  1024 https://doi.org/10.1371/journal.pone.0136249.
- 1025

- Rossouw, D., Du Toit, M., Bauer, F. F., 2012. The impact of co-inoculation with *Oenococcus oeni*on the trancriptome of *Saccharomyces cerevisiae* and on the flavour-active metabolite profiles during
  fermentation in synthetic must. Food Microbiol. 29(1), 121-131.
  https://doi.org/10.1016/j.fm.2011.09.006.
- 1030
- 1031 Rossouw, D., Meiring, S. P., Bauer, F. F., 2018. Modifying *Saccharomyces cerevisiae* adhesion
  1032 properties regulates yeast ecosystem dynamics. mSphere. *3*(5), e00383-18.
  1033 https://doi.org/10.1128/mSphere.00383-18.
- 1034
- Roullier-Gall, C., Hemmler, D., Gonsior, M., Li, Y., Nikolantonaki, M., Aron, A., Coelho, C.,
  Gougeon, R. D., Schmitt-Kopplin, P., 2017. Sulfites and the wine metabolome. Food Chem. 237,
  106-113. https://doi.org/10.1016/j.foodchem.2017.05.039.
- 1038
- Rubio-Bretón, P., Gonzalo-Diago, A., Iribarren, M., Garde-Cerdán, T., Pérez-Álvarez, E. P., 2018. 1039 1040 Bioprotection as a tool to free additives winemaking: Effect on sensorial, anthocyanic and aromatic 1041 profile of young red wines. LWT-Food Sci. Technol. 98. 458-464. 1042 https://doi.org/10.1016/j.lwt.2018.08.050.
- 1043
- Santos, A., Navascués, E., Bravo, E., Marquina, D., 2011. Ustilago maydis killer toxin as a new tool
  for the biocontrol of the wine spoilage yeast *Brettanomyces bruxellensis*. Int. J. Food Microbiol.
- 1046 145(1), 147-154. https://doi.org/10.1016/j.ijfoodmicro.2010.12.005.
- 1047
- 1048 Saravanakumar, D., Spadaro, D., Garibaldi, A., Gullino, M. L., 2009. Detection of enzymatic activity
- 1049 and partial sequence of a chitinase gene in *Metschnikowia pulcherrima* strain MACH1 used as post-
- 1050 harvest biocontrol agent. Eur. J. Plant Pathol. 123(2), 183-193. https://doi.org/10.1007/s10658-008-
- 1051 9355-5.

- Saravanakumar, D., Ciavorella, A., Spadaro, D., Garibaldi, A., Gullino, M. L., 2008. *Metschnikowia pulcherrima* strain MACH1 outcompetes *Botrytis cinerea*, *Alternaria alternata* and *Penicillium expansum* in apples through iron depletion. Postharvest Biol. Technol. 49, 121-128.
  https://doi.org/10.1016/j.postharvbio.2007.11.006.
- 1057
- Schmitt, M. J., Breinig, F., 2002. The viral killer system in yeast: from molecular biology to
  application. FEMS Microbiol. Rev. 26(3), 257-276. https://doi.org/10.1111/j.15746976.2002.tb00614.x.
- 1061
- Shekhawat, K., Patterton, H., Bauer, F. F., Setati, M. E., 2019. RNA-seq based transcriptional analysis
  of *Saccharomyces cerevisiae* and *Lachancea thermotolerans* in mixed-culture fermentations under
  anaerobic conditions. BMC Genomics. 20(1), 145. https://doi.org/10.1186/s12864-019-5511-x.
- 1065
- Siedler, S., Balti, R., Neves, A. R., 2019. Bioprotective mechanisms of lactic acid bacteria against
  fungal spoilage of food. Curr. Opin. Biotechnol. 56, 138-146.
  https://doi.org/10.1016/j.copbio.2018.11.015.
- 1069
- Siedler, S., Rau, M. H., Bidstrup, S., Vento, J. M., Aunsbjerg, S. D., Bosma, E. F., McNair, L. M.,
  Beisel, C. L., Neves, A. R., 2020. Competitive exclusion is a major bioprotective mechanism of
  lactobacilli against fungal spoilage in fermented milk products. Appl. Environ. Microbiol. 86(7),
  e02312-19. https://doi.org/10.1128/AEM.02312-19
- 1074
- 1075 Simonin, S., Alexandre, H., Nikolantonaki, M., Coelho, C., Tourdot-Maréchal, R., 2018. Inoculation
- 1076 of *Torulaspora delbrueckii* as a bio-protection agent in winemaking. Food Res. Int. 107, 451-461.
- 1077 https://doi.org/10.1016/j.foodres.2018.02.034

- Singh, V. P., 2018. Recent approaches in food bio-preservation-a review. Open. Vet. J. 8(1), 104111. https://doi.org/10.4314/ovj.v8i1.16.
- 1081
- Sipiczki, M., 2006. *Metschnikowia* strains isolated from botrytized grapes antagonize fungal and
  bacterial growth by iron depletion. Appl. Environ. Microbiol. 72(10), 6716-6724.
  https://doi.org/10.1128/AEM.01275-06.
- 1085
- 1086 Sipiczki, M., 2016. Overwintering of vineyard yeasts: survival of interacting yeast communities in
- 1087 grapes mummified on vines. Front. Microbiol. 7, 212. https://doi.org/10.3389/fmicb.2016.00212
- 1088
- Sipiczki, M., 2020. *Metschnikowia pulcherrima* and related pulcherrimin-producing yeasts: fuzzy
  species boundaries and complex antimicrobial antagonism. Microorganisms. 8(7), 1029.
  https://doi.org/10.3390/microorganisms8071029.
- 1092
- Su, Y., Seguinot, P., Sanchez, I., Ortiz-Julien, A., Heras, J. M., Querol, A., Camarasa, C., Guillamón,
  J. M., 2020. Nitrogen sources preferences of non-*Saccharomyces* yeasts to sustain growth and
  fermentation under winemaking conditions. Food Microbiol. 85, 103287.
  https://doi.org/10.1016/j.fm.2019.103287.
- 1097
- Taillandier, P., Lai, Q. P., Julien-Ortiz, A., Brandam, C., 2014. Interactions between *Torulaspora delbrueckii* and *Saccharomyces cerevisiae* in wine fermentation: influence of inoculation and nitrogen content. World J. Microbiol. Biotechnol. 30, 1959-1967. https://doi.org/10.1007/s11274014-1618-z.
- 1102

1103	Todd, B. E. N., Fleet, G. H., Henscke, P. A., 2000. Promotion of autolysis through the interaction of
1104	killer and sensitive yeasts: potential application in sparkling wine production. Am. J. Enol. Vitic. 51,
1105	65-72.
1106	
1107	Tremonte, P., Pannella, G., Succi, M., Tipaldi, L., Sturchio, M., Coppola, R., Luongo, D., Sorrentino,

E., 2017. Antimicrobial activity of Lactobacillus plantarum strains isolated from different

- 1109 environments: a preliminary study. Int. Food Res. J. 24(2), 852-859.
- 1110

- 1111 Tronchoni, J., Curiel, J. A., Morales, P., Torres-Pérez, R., Gonzalez, R., 2017. Early transcriptional 1112 response to biotic stress in mixed starter fermentations involving Saccharomyces cerevisiae and 1113 Torulaspora delbrueckii. Int. J. Food Microbiol. 241, 60-68. https://doi.org/ 1114 10.1016/j.ijfoodmicro.2016.10.017.
- 1115
- 1116 Váchová, L., Palková, Z., 2018. How structured yeast multicellular communities live, age and die?.
  1117 FEMS Yeast Res. 18(4), foy033. https://doi.org/10.1093/femsyr/foy033.
- 1118
- 1119 Varela, C., Borneman, A. R., 2017. Yeasts found in vineyards and wineries. Yeast 34(3), 111-128.
  1120 https://doi.org/10.1002/yea.3219.
- 1121
- 1122 Velázquez, R., Zamora, E., Álvarez, M., Álvarez, M. L., Ramírez, M., 2016. Using mixed inocula of
  1123 Saccharomyces cerevisiae killer strains to improve the quality of traditional sparkling-wine. Food
- 1124 Microbiol. 59, 150-160. https://doi.org/10.1016/j.fm.2016.06.006.
- 1125
- 1126 Velázquez, R., Zamora, E., Álvarez, M. L., Ramírez, M., 2019. Using Torulaspora delbrueckii killer
- 1127 yeasts in the elaboration of base wine and traditional sparkling wine. Int. J. Food Microbiol. 289, 134-
- 1128 144. https://doi.org/10.1016/j.ijfoodmicro.2018.09.010.

1130

1131

1132	
1133	Vilela, A., 2019. Use of nonconventional yeasts for modulating wine acidity. Fermentation. 5(1), 27.
1134	https://doi.org/10.3390/fermentation5010027.
1135	
1136	Villalba, M. L., Sáez, J. S., Del Monaco, S., Lopes, C. A., Sangorrín, M. P., 2016. TdKT, a new killer
1137	toxin produced by Torulaspora delbrueckii effective against wine spoilage yeasts. Int. J. Food
1138	Microbiol. 217, 94-100. https://doi.org/10.1016/j.ijfoodmicro.2015.10.006.
1139	
1140	Wang, C., Mas, A., Esteve-Zarzoso, B., 2015. Interaction between Hanseniaspora uvarum and
1141	Saccharomyces cerevisiae during alcoholic fermentation. Int. J. Food Microbiol. 206, 67-74.
1142	https://doi.org/10.1016/j.ijfoodmicro.2015.04.022.
1143	
1144	Wang, C., Mas, A., Esteve-Zarzoso, B., 2016. The interaction between Saccharomyces cerevisiae
1145	and non-Saccharomyces yeast during alcoholic fermentation is species and strain specific. Front.
1146	Microbiol. 7, 502. https://doi.org/10.3389/fmicb.2016.00502.
1147	
1148	Wisniewski, M., Biles, C., Droby, S., McLaughlin, R., Wilson, C., Chalutz, E., 1991. Mode of action
1149	of the postharvest biocontrol yeast Pichia guillermondii. I. Characterization of attachment to Botrytis
1150	cinerea. Physiol. Mol. Plant Pathol. 39(4), 245-258. https://doi.org/10.1016/0885-5765(91)90033-E.
1151	
1152	Zagorc, T., Maráz, A., Cadez, N., Jemec, K. P., Péter, G., Resnik, M., Nemaničd, J., Raspor, P., 2001.
1153	Indigenous wine killer yeasts and their application as a starter culture in wine fermentation. Food
1154	Microbiol. 18(4), 441-451. https://doi.org/10.1006/fmic.2001.0422.
	45

Vilela, A., 2018. Lachancea thermotolerans, the non-Saccharomyces yeast that reduces the volatile

acidity of wines. Fermentation. 4(3), 56. https://doi.org/10.3390/fermentation4030056.

spp. and its effect on the metabolic profiles of fermentation communities. Appl. Environ. Microbiol.
87(9), e02992-20. https://doi.org/10.1128/AEM.02992-20.
Zhang, C., Yang, M., Ericsson, A. C., 2019. Antimicrobial peptides: potential application in liver
cancer. Front. Microbiol. 10, 1257. https://doi.org/10.3389/fmicb.2019.01257.

Zhang, H., Du, H., Xu, Y., 2021. Volatile organic compound-mediated antifungal activity of Pichia

- 1163 Zhao, X., Wu, H., Lu, H., Li, G., Huang, Q., 2013. LAMP: a database linking antimicrobial peptides.
- 1164 PloS one. 8(6), e66557. https://doi.org/10.1371/journal.pone.0066557.

- ....

- 1181 Figure captions
- 1182 Fig. 1 Antagonistic behaviour demonstrated in wine microorganisms. Antagonistic behaviour
- 1183 was investigated *in vitro* (A), on grapes (B) and in wine (C).
- 1184 Fig. 2 Antagonistic strategies of wine microorganisms
- 1185
- 1186

# 1187 **Table 1**

# 1188 Role of starter cultures in wine and their bioprotective strategies

-

Yeasts	Starter ability	References	Bioprotec	References		
			Passive strategies	Active strategies		
Saccharomyces	- Alcoholic fermentation conduction	Parapouli et al.,	- Fast nutrient uptake	- Toxic compounds production	Alonso-del-Real et al., 2019	
cerevisiae	- Positive impact on the organoleptic	2020	- Multicellular consortia	- Ethanol	Tronchoni et al., 2017	
	quality of wines		formation	- SO <sub>2</sub>	Rossouw et al., 2015; 2018	
				- Antimicrobial peptides	Albergaria and Arneborg, 2016	
				(saccharomycin; killer	Gonzàlez et al., 2018	
				toxins)	Englezos et al., 2019b	
				- VOCs (higher alcohols)		
				- Cell-to-cell contact		
Torulaspora	- Low acetic acid production	Benito, 2018a	- Fast nutrient uptake	- Toxic compounds production	Prior et al., 2019	
delbrueckii	- Reduction of ethanol content			- Ethanol	Su et al., 2020	
	- Increase of glycerol content			- Antimicrobial peptides (Tdkt	Simonin et al., 2018	
	- Release of mannoproteins and			Killer toxin)	Villaba et al., 2016	
	polysaccharides					
	- Production of high levels of fruity					
	esters, thiols, and terpenes and lower					
	amounts of higher alcohols					

Metschnikowia	- Low acetic acid production	Morata et al., 2019a	- Iron immobilization	- Toxic compounds production	Sipiczki, 2006, Sipiczki, 2020
pulcherrima	- Ethanol content reduction		(pulcherrimic acid production)	- Ethanol	Gore-Lloyd et al., 2019
	- Increase of glycerol content		- high growth rate and a short lag	- Lytic enzymes	Kunchen et al., 2019
	- Production of high levels of esters		phase	- VOCs (ethyl alcohol and	Cordero-Bueso et al., 2017
	and higher alcohols (isobutanol and		- Biofilm formation	ethyl acetate)	Pawlikowska et al., 2019
	phenylethanol)				Contarino et al., 2019
	- Enhances varietal and fruity aromas				Saravanakumar et al., 2009
Lachancea	- Increase of titratable acidity	Benito, 2018b	- Fast nutrient uptake	- Toxic compounds production	Fiori et al., 2014
thermotolerans	- Ethanol content reduction	Morata et al., 2019b	- Short lag phase and ability to	- Ethanol	Farbo et al., 2018
	- Increase of glycerol content	Porter et al., 2019	survive until the end of	- Lactic acid	Shekhawat et al., 2019
	- Production of high levels of ethyl		fermentation	- Killer toxins	Rubio-Bretón et al., 2018
	lactate, ethyl hexanoate and 2-		- Filamentous growth/ flocculation	- VOCs (2- phenylethanol)	Vilela, 2018
	phenylethanol				Nally et al., 2018
	- Reduction of phenylethyl acetate				
Pichia kluyveri	- Ethanol content reduction	Varela and	- Biofilm formation	- Toxic compounds production	Cordero-Bueso et al., 2017
	- Increase of glycerol content	Borneman, 2017		- Ethanol	Sipiczki, 2016
	- Production of thiols (hotrienol and			- Killer toxins	Prior et al., 2019
	linalool oxide)			- Lytic enzymes	Labbani et al., 2015
	- enhance varietal aromas			- VOCs (isoamyl acetate,	Mewa-Ngongang et al., 2019a,b
				isoamyl alcohol, 2-phenyl	Englezos et al. (under preparation)

### ethylacetate and 2-

#### phenylethanol)

Sta	rmerella	- Ethanol content reduction	Englezos et al.,	- Relative fast amonium uptake	- Toxic compounds production	Prendes et al., 2018
bac	eillaris	- Increase of glycerol content	2017		- Ethanol	Lemos Junior et al., 2016, 2020
		- Production of high levels of linalool,			- Killer toxins	Englezos et al., 2018, 2019b
		organic acids			- Lytic enzymes	Binati et al., 2020
					- VOCs (benzyl alcohol)	Roca Mesa et al., 2020
						Bagheri et al., 2018

### Lactic Acid

#### Bacteria

0					D: 1 / 1 2015 2010
Denococcus oeni	- Decrease of titratable acidity		- Better adaptation to wine	- Toxic compounds production	Dimopoulou et al., 2015, 2018
	- Diacetyl production – 'buttery'		conditions	- Lactic acid	Bastard et al., 2016
	flavour		- Biofilm formation	- BLIS	Knoll et al., 2008
	- Release of secondary metabolites				Lasik-Kurdyś and Sip, 2019
	that impact on wine aroma and				
	flavor				
actiplantibacillus	- Titratable acidity decrease	Krieger-Weber et	- Competitive exclusion	- Toxic compounds production	Rubio-Breton et al., 2018
olantarum	- Diacetyl production – 'buttery'	al., 2020	- Nutrient competition	- Lactic acid	Melkonian et al., 2019
	flavour	Brizuela et al., 2019	- Biofilm formation	- Bacteriocins	Sielder et al., 2020
					Tremonte et al., 2017
					Bartle et al., 2019

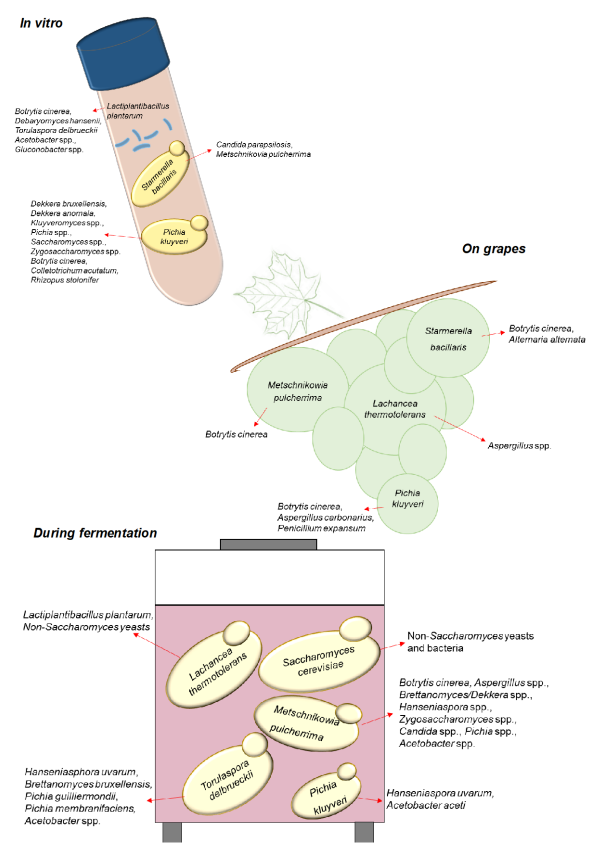
- Positive effects on the organoleptic
- properties of wine due to a diverse

array of enzymes

- Improvement of color in red wines

## 1190 Figure captions

## 1191 Fig. 1



1192

## 1193 Fig. 2

