

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30

Research Article

**Autochthonous *Oenococcus oeni* Strain to Avoid
Histamine Formation in Red Wines: A Study in Real
Winemaking Conditions**

Silvia Pérez-Magariño,^{1*} Estela Cano-Mozo,¹ Clara Albors,² Antonio Santos,³
and Eva Navascués²

¹Instituto Tecnológico Agrario de Castilla y León. Consejería de Agricultura y Ganadería. Ctra Burgos Km 119, Finca Zamadueñas. 47071 Valladolid, Spain; ²Pago de Carraovejas State Winery. Camino de Carraovejas, s/n. 47300 Peñafiel, Valladolid, Spain; and ³Department of Genetics, Physiology and Microbiology. Unit of Microbiology. Biology Faculty, Complutense University of Madrid, Madrid, Spain.

*Corresponding author: (permagsi@itacyl.es; tel: +34 983 415245)

Acknowledgments: This study is part of HEALTHWINE Project and GLOBALVITI Project (IDI-20160746 with the financial support of the CDTI-CIEN program). The authors would like to thank Agrovin S.A. for technical assistance.

Manuscript submitted Feb 19, 2020, revised Mar 19, 2020, Sept 15, 2020, accepted Nov 19, 2020

This is an open access article distributed under the CC BY license (<https://creativecommons.org/licenses/by/4.0/>).

By downloading and/or receiving this article, you agree to the Disclaimer of Warranties and Liability. The full statement of the Disclaimers is available at <http://www.ajevonline.org/content/proprietary-rights-notice-ajev-online>. If you do not agree to the Disclaimers, do not download and/or accept this article.

Abstract: The production of wines with low biogenic amine (BA) concentrations is one of the current concerns in the wine sector and strategies to avoid their formation during winemaking are of especial interest. The aim of this work was to determine the influence of selected autochthonous *Oenococcus oeni* lactic acid bacteria (LAB) on the BA content in red wines and their prevalence against the indigenous microbiota to avoid BA formation. Sixty-seven red wines were elaborated at industrial scale in real winemaking conditions in three consecutive vintages. LAB implantation and BA concentrations in every wine obtained were determined at different stages of winemaking process. The results clearly indicated that the use of selected *O. oeni* strains unable to produce BA in combination with an adapted biomass production is a good strategy to control histamine

31 production in wines. These practices, carried out over three consecutive years, were also observed
32 to ensure the persistence of this selected autochthonous *O. oeni* strain CECT 9749 against other
33 indigenous microbiota, in the whole winery. Furthermore, the analysis of BA content during wine
34 aging in barrels indicated that low BA content was maintained, resulting in healthier wines for the
35 consumer.

36 **Keywords:** autochthonous strains, biogenic amines, histamine, lactic acid bacteria, malolactic
37 fermentation, *Oenococcus oeni*

38 Introduction

39 Biogenic amines (BA) are low molecular weight organic nitrogenous compounds naturally
40 occurring in fermented foods and wines (Gardini et al. 2016). Some BA are present in grapes but
41 they are mainly formed by amino acids decarboxylation through the enzymatic activity of
42 microbial decarboxylases. In normal physiological conditions, BA are metabolized by
43 gastrointestinal enzymes (Gardini et al., 2016, Ancín-Azpilicueta et al. 2019), however, the
44 consumption of elevated BA doses could have adverse effects on consumer's health from a
45 toxicological point of view (EFSA 2011). Histamine and tyramine are considered the most toxic
46 amines, causing negative effects such as headaches, nausea, hypo- and hypertension, respiratory
47 disorders, tachycardia and various allergic disorders among others (Landete et al. 2005, Ladero et
48 al. 2010, Moreno-Arribas et al. 2010, Gardini et al. 2016).

49 Other BA, such as putrescine and cadaverine, are non-toxic polyamines but they can
50 potentiate the toxic effects of BA through the inhibition of enzymes that detoxify histamine,

51 tyramine and phenylethylamine (Straub et al. 1995, Özogul and Özogul 2019). In addition, these
52 polyamines adversely affect wine sensory quality (Tomera 1999, García-Villar et al. 2007).

53 Therefore, the presence of high BA concentrations is related to wine safety (EFSA 2011,
54 Martuscelli et al. 2013), but they may also be indicators of wine quality and hygienic conditions
55 (Del Prete et al. 2009, Gardini et al. 2016).

56 Currently, the European Union has not established a regulatory limit for histamine content
57 or any other BA in wines. However, different countries in Europe have set different
58 recommendation limits for histamine in wine ranging from 2 to 10 mg/L (Smit et al. 2008). The
59 International Organisation of Vine and Wine (OIV) recommends not to exceed the limit of 12
60 mg/L of histamine.

61 Estimating safe levels of BA in wine is difficult since it depends on several factors such
62 type of amine, concentration and the physiological conditions of consumers (individuals that do
63 not have degrading mechanisms of these compounds) and the consumption of other BA-containing
64 foods, which could increase their toxicity (Ancín-Azpilicueta et al. 2019). In addition, in humans,
65 it has been demonstrated that ethanol and acetaldehyde may enhance the toxicity of amines through
66 the inhibition of amino oxidase enzymes, responsible for amine degradation (Zimatkin and
67 Anichtchik 1999).

68 Histamine, putrescine, cadaverine, tyramine, phenylethylamine and spermidine are the
69 main BA present in wine (Moreno-Arribas and Polo 2009, EFSA 2011). Some BA, such as
70 putrescine and spermidine, can be present in grapes (Landete et al. 2005, Izquierdo-Cañas et al.
71 2008, Del Prete et al. 2009), but most of them are produced during winemaking by
72 microorganisms. The formation of BA requires the presence of amino acids, microorganisms with

73 decarboxylase activity and the favorable conditions for their growth (Smit et al. 2008, Costantini
74 et al. 2009, Moreno-Arribas et al. 2010). Some authors reported that yeasts can produce BA during
75 alcoholic fermentation (AF) (Ancín-Azpilicueta et al. 2008, Smit et al. 2008) or during wine
76 storage (Jiménez-Moreno et al. 2003, Hernández-Orte et al. 2008). Although reported results are
77 contradictory, it is assumed that the greatest amount of BA, especially histamine, is produced
78 during malolactic fermentation (MLF) through the decarboxylation activity of lactic acid bacteria
79 (LAB) that transform amino acids into BA (Soufleros et al. 1998, Lonvaud-Funel 1999, Landete
80 et al. 2005, Marcobal et al. 2006).

81 *Oenococcus oeni* is the most habitual species of LAB found after AF in both spontaneous
82 MLF, due to the growth of indigenous strains, and inoculated MLF with selected strains (Moreno-
83 Arribas et al. 2003, Nehme et al. 2010). This is due to the fact that *O. oeni* was best adapted to the
84 harsh wine conditions (high ethanol, low pH, low nutrients and SO₂) (Lonvaud-Funel 1999). Other
85 LAB species of different genera such as *Lactobacillus*, *Leuconostoc* and *Pediococcus* can also
86 grow in wine, especially if pH values are higher than 3.5 (Lonvaud-Funel 2001, Costantini et al.
87 2009). The ability of these bacterial species to produce BA seems to be strain-dependent (Berbegal
88 et al. 2017), therefore, it is important to select LAB strains non-BA producers to minimize the BA
89 content. Some authors reported that selected LAB without decarboxylase activity can be used to
90 prevent BA formation in wines (Marcobal et al. 2006). Moreover, it was demonstrated that
91 simultaneous yeast/ LAB co-inoculation was more effective to avoid BA production and for
92 obtaining wines with better sensory characteristics (Massera et al 2009, Izquierdo-Cañas et al.
93 2014).

94 The occurrence of BA in wines has been studied in the last years and winemakers are
95 looking for strategies to avoid their formation during winemaking to obtain wines with low BA
96 concentrations (Benito 2019a). Different strategies to control the production of BA or to degrade
97 BA have been suggested. The use of non-*Saccharomyces* strains that decrease malic acid content,
98 or the usage of different retention systems (not used for quality wines) have been studied in depth
99 (Benito 2019b, Rodríguez-Bencomo and Mehdi 2019). Nowadays, one of the main control
100 strategies to prevent the formation of BA in wines at industrial scale is the inoculation of selected
101 non-BA producers *O. oeni* strains (Moreno-Arribas et al. 2003, Izquierdo-Cañas et al. 2009).
102 However, the induction of MLF by these commercial LAB was not always successful in a whole
103 winery, due to different factors such as wine is a very harsh medium for LAB growth (Ruiz et al.
104 2010) or the competitive advantage of autochthonous microbiota to winery conditions. In this
105 sense, some authors suggested the use of autochthonous selected LAB of a specific wine-
106 producing area, which could improve the MLF development (Ruiz et al. 2010, Berbegal et al.
107 2017).

108 Therefore, the aim of this work was to study the influence of selected autochthonous *O.*
109 *oeni* on the BA content of red wines from Ribera del Duero during three consecutive vintages
110 (2016, 2017 and 2018) and study their prevalence against the indigenous microbiota to avoid BA
111 formation in these wines. The importance of managing the adaptation of the LAB culture and how
112 to ensure its presence in winery tanks was also studied in order to avoid histamine formation in
113 wines. The effect of LAB inoculation and wine aging in oak barrels on BA production was also
114 evaluated.

115

Materials and Methods

116 **Winemaking conditions.** Red wines were elaborated following the usual winemaking
117 process in Pago de Carraovejas State winery, located in Peñafiel (Valladolid, Spain) in Ribera del
118 Duero Geographical Indication in Spain. Vineyards, belonged to the winery, are disposed in the
119 same valley between 850- and 950-meters elevation and are cultivated under organic farming
120 conditions. Grape variety is 'Tinto Fino' ('Tempranillo') Carraovejas clone with 110 R rootstock.

121 Maturity control was checked using not only technical parameters (sugar, pH, Total acidity,
122 weight/100 grape berries) but also quality parameters such “Glories method” (Cromoenos ®) that
123 evaluates the phenolic maturity of grapes. It provides information on both, the quantity (total
124 potential in anthocyanins and tannins) and the quality (anthocyanin extractability, seed maturity)
125 of the polyphenols.

126 This work, performed during three consecutive harvests, was done with the real
127 winemaking conditions and practices used at cellar facilities. Taking into consideration the
128 differences in characteristics and grape yield accounted in each vintage, the experimental design
129 was slightly different each year. However, in order to establish a suitable point of comparison,
130 each vintage, half of the harvest was inoculated with the selected *O. oeni* CECT 9749 strain and
131 compared with the other half.

132 In this study, three consecutive years were compared: 2016 (harvest start/end, 4 Oct/29
133 Oct), 2017 (harvest start/end, 19 Sept/30 Sept) and 2018 (harvest start/end, 29 Sept/12 Oct).
134 Grapes were harvested manually and immediately transported to the winery in 12-kg boxes. Then,
135 grapes went through double selection to avoid any rot or not mature cluster. After that, clusters
136 were destemmed and lightly crushed.

137 For all vintages, the alcoholic fermentations were carried out in stainless steel tanks of
138 25,000 L and were filled in with 18,000 kg of red grapes. It was collected 396,000 kg, 324,000 kg
139 and 486,000 kg, respectively in the three vintages, and were filled 22, 18 and 27 fermentation tanks
140 (Table 1).

141 Alcoholic fermentation (AF) was carried out by using the strain *Saccharomyces cerevisiae*
142 CECT 12008 (Spanish Type Culture Collection) that was previously isolated and selected from
143 the vineyards of Pago de Carraovejas State winery. Yeast strain was inoculated in the filling of
144 each tank ensuring 10^6 cells/mL of must or mg of grape. Also, some fermentations were
145 spontaneously developed by the indigenous yeasts in each harvest, to see the influence of
146 spontaneous alcoholic fermentation in the development of the inoculated bacterial culture and/or
147 in the production of biogenic amines.

148 Alcoholic fermentation kinetics were conducted at $24^{\circ}\text{C} \pm 1^{\circ}\text{C}$. Temperature (cap and
149 liquid) and sugar content was measured daily during fermentation. Addition of SO_2 was done at
150 30 mg/L during the filling of the fermentation tanks. It was supplemented using organic nitrogen
151 without diammonium phosphate (Actimax Natura, Agrovín) at 0.50 g/L doses, at the beginning of
152 the fermentation. For color extraction, maceration was conducted along fermentation and pumping
153 with aeration twice a day for color extraction. When fermentation and maceration finished, the
154 wines were devatted and only the first pressings (55% yield) were taken.

155 **Lactic acid bacteria strain and inocula preparation.** An autochthonous *O. oeni* strain
156 CECT 9749 was used to inoculate the experimental wines of this study. This strain was isolated
157 and selected from red wines of the same winery in a previous study (Berbegal et al. 2017). This
158 non-histaminogenic strain was selected because of its prevalence in wines, high alcohol resistance

159 and high quality properties that not affects wine fruitiness. In addition, this strain was found to be
160 well-adapted to the conditions of these wines, competing effectively with other indigenous LAB
161 strains (Berbegal et al. 2017).

162 Inoculum production was performed following a strict scale-up procedure to reach $>10^9$
163 CFU/mL. The selected *O. oeni* CECT 9749 strain was initially grown in MLO broth (Zúñiga et al.
164 1993) to reach an early stationary phase. Cells were then centrifuged (8,000 rpm, 10 min), washed
165 with Ringer's solution and transferred to 10 mL culture media at a final concentration of 1×10^6
166 CFU/mL. Then, cellular concentration was scaled up through a three-stage (0.5 L, 10 L and finally,
167 80 L) procedure using a liquid production medium described in Berbegal et al. (2015), with several
168 modifications that allowed the culture to adapt to the harsh conditions of red wines: use of red
169 wine with more than 70 TPI (Total Polyphenol Index), higher alcohol degree, fermentation
170 activator enhancers as Actimax NATURA (Agrovin S.A., Alcázar de San Juan, Ciudad Real,
171 Spain) and cofactors (Mg, Mn, folic acid, ascorbic acid and retinol).

172 The industrial biomass production was performed in a fermenter with 80 L of production
173 medium under sterile conditions (Bioprocess technology Bio-pro 100 L), and it was used to
174 inoculate each tank of 25,000 L with the selected *O. oeni* strain (2×10^6 CFU/mL). A
175 microbiological analysis of the three scale-up steps was carried out to certify the correct
176 sterilization of the culture medium (Berbegal et al. 2015). This scale-up was done in Agrovin S.A.
177 facilities.

178 **Inoculation of lactic acid cultures and malolactic fermentation.** The autochthonous
179 selected *O. oeni* CECT 9749 strain was inoculated at the beginning of the AF, ensuring that free
180 SO_2 was zero. The first tanks of each harvest (2016, 2017 and 2018) were inoculated, trying to

181 maintain the same number of inoculated and non-inoculated tanks. The percentage of inoculated
182 tanks in the three consecutive harvests were 40.9%, 55.6% and 59.3%, respectively (Table 1).

183 Once AF was finished (glucose + fructose content < 1 g/L), MLF was conducted at 22°C ±
184 1°C, until malic acid was < 0.2 g/L. Then, it was considered that MLF was over.

185 **Aging in cellar.** After MLF, the wines were racked, corrected with SO₂ (approximately 0.5
186 of molecular SO₂), and later aged in French oak barrels (225 L) for 12 months in a cellar at 15°C
187 and with a controlled relative humidity of about 75%–85%.

188 **Chemicals.** Chromatographic grade reagents were provided by Carlo Erba Reagents
189 (Sabadell, Spain). 2-aminoheptanoic acid (internal standard) and diethylethoxymethylenemalonate
190 (DEEMM) were from Fluka (Sigma-Aldrich, Germany), and the remaining reagents were supplied
191 by Panreac (Madrid, Spain). Milli-Q grade water was obtained using a Millipore system (Bedford,
192 MA). The biogenic amine standards were purchased from Sigma-Aldrich (Steinheim, Germany).

193 **LAB and yeast counts and implantation.** RAPD-based genotyping was used to determine
194 the implantation of *O. oeni* CECT 9749 strain as described below. Before inoculation with *O. oeni*
195 CECT 9749, grape musts were checked in order to determine the amount of wild LABs from
196 vineyards (Figure 1A, red points). Then, wine samples were collected at three different moments
197 for LAB determination (*O. oeni* inoculation, after 48h of inoculation and in the second third of the
198 MLF) (Figure 1A black points). Samples were serially diluted and seeded onto MLO plates
199 (Zúñiga et al. 1993). Plates were counted and ten single colonies were randomly picked from plates
200 with 30-300 colonies and suspended in 10 µL of sterile milliU water (Millipore, Bedford, MA).
201 These suspensions were used in RAPD (Random Amplification of Polymorphic DNA)-PCR
202 amplification with M13 primers as described by Berbegal et al. (2017). In each electrophoresis

203 gel, comparison between the electrophoretic band profiles obtained from LAB isolates and *O. oeni*
204 CECT 9749 allowed to determine the percentage of implantation of the inoculated strain in each
205 tank (Figure 1C, 1D).

206 A similar procedure was developed in order to check the cellular densities of yeasts and the
207 implantation success of the inoculated yeast strain *S. cerevisiae* CECT 12008 (Figure 1B).
208 Samples, taken at three stages of alcoholic fermentation (corresponding to densities (1.100, 1.040
209 and 0.990 g/cm³), were seeded onto Malt Extract Agar and incubated. Yeasts colonies were
210 counted and ten of them were randomly picked from plates for genotyping. Colonies were
211 suspended in water and subjected to PCR-amplification of the interdelta regions (de Celis et al.
212 2019). The electrophoretic band pattern obtained for *S. cerevisiae* CECT 12008 was used for
213 reliable comparisons into the same electrophoresis gel.

214 **Analysis of biogenic amines.** Nine biogenic amines (histamine, tyramine, putrescine,
215 cadaverine, phenylethylamine, spermidine, agmatine, tryptamine, and isoamylamine) were
216 analyzed using the method described by Gómez-Alonso et al. (2007) with slight modifications
217 (Ortega-Heras et al. 2014). Aminoenone derivatives of amines were obtained by reaction with
218 DEEMM and after that, they were analyzed by reverse-phase high performance liquid
219 chromatography (RP-HPLC) in an Agilent Technologies LC series 1200 with a diode array
220 detection system (DAD) (Agilent, Stuttgart, Germany).

221 Samples were taken after MLF and after 12 months of aging in barrels after their respective
222 racking. BA analyses were carried out in duplicate.

223 **Analysis of oenological parameters.** Standard oenological parameters in wines were
224 determined according to the official analysis methods of OIV: pH, titratable acidity (as g/L tartaric

225 acid), and alcohol degree (% vol: mL ethanol/100 mL wine). Malic, lactic and acetic acids were
226 analyzed using enzymatic kits in an Y15 Analyser (Biosystems, Barcelona, Spain).

227 SO₂ was determined by the official analysis methods of OIV (OIV-MA-AS323-04A).

228 These oenological parameters were analyzed in samples after AF and when MLF was
229 finalized. Analyses were performed immediately after sample collection.

230 **Statistical analyses.** A one-way analysis of variance and a Fisher Least Significant
231 Difference test (LSD) at a significant level of $p < 0.05$ was performed to determine the effects of
232 inoculation and of aging in oak barrels on the BA content, using Statgraphics Centurion XVII.

233 Results

234 **Oenococcus oeni implantation.** The selected *O. oeni* strain was implanted in all the
235 inoculated wines and in almost all the non-inoculated ones. Only in eight non-inoculated wines
236 from 2016 vintage (tanks 15-S to 22-S), the selected *O. oeni* strain was not found or it was found
237 under 50% (Table 2). In 2017 and 2018 vintage, the inoculated strain was present in the whole
238 winery. The LAB implantation percentage is shown in Tables 2 to 4.

239 Figure 1 (Figure 1C and 1D) shows electrophoretic patterns (EP) of bands for ten randomly
240 picked colonies from MLO plates seeded with wines of two tanks of the 2016 vintage in order to
241 determine the percentage of *O. oeni* implantation. Overall, a strain is considered to be implanted
242 when it is clearly dominant, being the percentage of implantation superior to 80% (ie. eight of ten
243 EP are equal to the EP of the inoculated strain). In tank 16 1-I, the EP of all samples (lanes 2 to
244 11) were identical to the known EP (lane 12) of the inoculated *O. oeni* CECT 9749 strain, and
245 therefore the implantation was considered to be complete. However, in tank 16 3-I, 1 do not

246 matched with the EP of the inoculated *O. oeni* strain, and the implantation was considered to be
247 around 90%.

248 **Fermentation kinetics.** Kinetics of malic acid consumption and lactic acid production
249 followed the same pattern in all fermentations performed in the three harvests of the study. As an
250 example, Figure 1 (1C and 1D) shows malic acid/lactic acid kinetics during the progression of the
251 fermentation in two selected tanks (Tank 16 1-I and 16 3-I). In both cases, one with a good (100%)
252 implantation of the *S. cerevisiae* strain CECT 12008 (Tank 16 1-I, Figure 1B) and the other (Tank
253 16 3-I, Figure 1B), without a good implantation (0%) -similar to spontaneous-, the global kinetics
254 of malic acid consumption and lactic acid production were similar. Furthermore, this fact, jointly
255 considered with the results of Table 2, indicates that the influence of the yeast in the progression
256 malolactic fermentation was little.

257 **Biogenic amine analyses.** Tables 2 to 4 show the BA concentrations found in all analyzed
258 wines from the three vintages just after MLF and after 12 months aging in barrels. Spermidine and
259 phenylethylamine were found in low concentrations and no statistically significant differences
260 were observed between wines (data not shown). Agmatine, tryptamine and isoamylamine were not
261 detected in any wine. Therefore, the study was focused on the content of the main BA (histamine,
262 tyramine, cadaverine and putrescine) found in wines. Putrescine was the most abundant amine in
263 all the wines, with concentrations ranging between 2.9-12.0 mg/L.

264 A clear difference in histamine and putrescine contents between the inoculated and the non-
265 inoculated wines from 2016 vintage was observed in both stages, after MLF and after 12 months
266 of aging in barrels (Table 2). Non-inoculated wines presented statistically significant higher
267 histamine and putrescine concentrations than the inoculated ones. The wines with the *O. oeni* strain

268 implanted showed a reduction of 93.6 % in histamine and 30.2 % in putrescine. Furthermore, it
269 should be pointed out that those non-inoculated wines in which the selected *O. oeni* strain was also
270 implanted (tanks 10-S to 14-S), showed the lowest histamine content.

271 On the contrary, the same was not observed in wines from 2017 and 2018 vintages (Tables
272 3 and 4). The inoculation did not produce statistically significant differences in histamine and
273 putrescine concentrations. However, it should be taken into account that in these two vintages, the
274 content of these BA was very low and the presence of the *O. oeni* CECT 9749 strain was detected
275 in all the wines, not only in the inoculated ones but also in those non-inoculated.

276 No statistically significant differences were found in the tyramine and cadaverine
277 concentrations of the wines due to *O. oeni* inoculation, with the exception of the wines from 2017
278 vintage. However, the differences found in these wines were low. After MLF, the inoculated wines
279 had 1.30 mg/L of tyramine and 0.36 mg/L of cadaverine while the non-inoculated wines had 1.70
280 mg/L of tyramine and 0.49 mg/L of cadaverine (Table 3).

281 After 12 months of aging in barrels, significant increases were recorded for putrescine in
282 the non-inoculated wines from 2016 vintage. Slight increments in BA of 2017 wines were
283 observed, although, in general, they were very low.

284 **General oenological characteristics.** Mean values of the oenological parameters and the
285 standard deviation of wines from the three consecutive vintages (2016, 2017 and 2018) at the end
286 of AF and MLF are shown in Table 5. Wines from 2017 and 2018 showed a slightly higher alcohol
287 content (15.4%vol.) than 2016 wines (14.8%vol.). In addition, 2017 wines also showed slight
288 lower titratable acidity (5.2 g/L) and malic acid content (1.45 g/L) than wines obtained from 2016
289 and 2018 vintages (mean values of 5.9 g/L of tartaric acid and 1.9 g/L of malic acid). In spite of

290 these slight differences, it can also be highlighted that wines from the three vintages showed similar
291 oenological characteristics, with pH ranges around 3.6-3.8; titratable acidities between 5.2-6.1 g/L
292 of tartaric acid; and alcoholic degrees between 14.7%vol.-15.6%vol.

293 Concerning the effect of the *O. oeni* inoculation, no statistically significant differences
294 were found in most of the oenological parameters between the inoculated and the non-inoculated
295 wines in each vintage, with the exception of the lactic and acetic acid concentrations in 2018 wines.

296 All wines completed the MLF since malic acid concentrations were below 0.2 g/L. Before
297 aging, wines were corrected with SO₂ and molecular SO₂ was up of 0.5 (Table 5), allowing to
298 maintain its antibacterial activity.

299 Discussion

300 **Oenococcus oeni** implantation. To ensure *O. oeni* implantation to avoid wild
301 histaminogenic LAB development, an efficient production of the selected *O. oeni* strain was
302 achieved. Highly active cultures exerting a good malic acid enzymatic activity prevent sugar
303 consumption, avoiding increments in acetic acid and lactic acids.

304 Several studies have reported that the inoculation of bacterial starters before or during AF
305 (simultaneous inoculation of yeast and LAB) allows better control of MLF in winemaking
306 (Massera et al. 2009, Azzolini et al. 2010, Smit and DuToit 2013, Izquierdo-Cañas et al. 2014). In
307 this sense, some preliminary experiences (data not shown) were conducted during 2013, 2014 and
308 2015 harvests. We observed that, when wines were inoculated with *O. oeni* CECT 9749 at the end
309 of AF, wild LAB counts coming from grapes were high, making it impossible to ensure the
310 implantation of the inoculated strain, consequently, histamine production was not reduced.

311 Therefore, in this study, it was decided to inoculate a selected non-histaminogenic *O. oeni*
312 strain at the beginning of AF.

313 The development of this strain was successful in all the inoculated wines of the three
314 vintages. In 2016 vintage, the selected *O. oeni* strain was found in the 38% of the non-inoculated
315 wines and in the following vintages, 2017 and 2018, it was found in all the wines, independently
316 of the inoculation. This is a very interesting result since it has been observed that over the years,
317 the autochthonous selected *O. oeni* strain is predominant during the elaboration of the wines of the
318 whole winery, even in those wines that were not previously inoculated. Therefore, the addition of
319 the well-adapted culture of *O. oeni* CECT 9749 strain to the different tanks during AF seems to
320 generate a high prevalence of the strain during the three consecutive vintages studied. Similar
321 results have been obtained using selected *Saccharomyces cerevisiae* strains as starters for wine
322 fermentation, indicating that these practices could have an important incidence on microbial
323 diversity in surrounding vineyards (de Celis et al. 2019).

324 In winery real conditions, yeast and bacteria does not work separately. Inoculated and non-
325 inoculated tanks are processed using the same material (pumps, tubes, etc) and as the inoculated
326 tanks are routinely the first to be used for fermentation, the dissemination of the bacterial cultures
327 to the non-inoculated tanks is a feasible situation.

328 **Biogenic amine analyses.** The implantation results are according to the BA data of these
329 wines. The non-inoculated and non-LAB implanted wines from 2016 presented the highest
330 histamine contents (Table 3). Histamine formation occurs mainly during MLF, and no significant
331 increases were observed during aging in barrels, with some exceptions. The increase of histamine
332 was showed in the wines with very low initial concentrations, but this increase is not considered

333 significant since the histamine values were always less than 1 mg/L. The results found in
334 bibliography are contradictory respect to the BA formation mainly during MLF or during aging
335 (Soufleros et al. 1998, Lonvaud-Funel 1999, Jiménez-Moreno et al. 2003, Landete et al. 2005,
336 Marcobal et al. 2006, Hernández-Orte et al. 2008), and also are dependent on whether the MLF
337 was carried out spontaneously or by inoculating LAB (Marcobal et al. 2006, Hernández-Orte et al.
338 2008). Hernández-Orte et al. (2008) showed an increase of histamine concentration after 6 months
339 of oak aging that was higher in wines that conducted MLF with indigenous bacteria than in
340 inoculated wines. Other authors indicated that LAB inoculation did not give rise to an increase of
341 histamine during oak aging (Marcobal et al. 2006). The contradictory results found by different
342 authors could be due to the prevalence or not of the non-histaminogenic strain, since some works
343 evaluated the BA content but not the LAB strain that carried out the MLF. Taking into account our
344 results, the presence of the *O. oeni* strain non-histamine producer during MLF avoids the increase
345 of histamine values during the aging time.

346 Mean values of histamine in the inoculated and the non-inoculated wines from the three
347 vintages studied after MLF and after 12 months of aging in barrels are shown in Figure 2. The use
348 of the selected autochthonous *O. oeni* strain from the red wines of the winery has drastically
349 reduced the content of histamine in its wines. A progressive reduction in histamine content in
350 wines has been observed over the years. Histamine mean values have been reduced from 6 mg/L
351 in inoculated wines and 18 mg/L in non-inoculated ones in 2011 (Bergal et al. 2017) to values
352 < 1 mg/L in wines of 2017 and 2018 vintages. These low histamine values were also found in the
353 non-inoculated wines, but in those where the selected *O. oeni* strain has been implanted. This might
354 be due to the prevalence of the selected *O. oeni* strain against other indigenous LAB that manage

355 to avoid the growth of other populations' histamine producers. In addition, this also leads the
356 maintenance of low histamine levels during aging in barrels. Therefore, it seems that the
357 implantation of the selected *O. oeni* strain is determinant to reduce the risk of histamine formation
358 and to obtain wines with low BA concentrations. The influence of the use of selected *O. oeni* strain
359 was more significant in the histamine and putrescine content than in the cadaverine and tyramine.

360 Putrescine concentrations were also significantly higher in the non-inoculated wines than
361 in the inoculated ones but only in wines from 2016. Therefore, the indigenous microbiota that
362 conducted the MLF in these wines were responsible for producing these BA. These results are in
363 agreement with those found by other researchers who indicated that spontaneous MLF has more
364 risk to produce wines with high BA contents (Izquierdo-Cañas et al. 2008, Berbegal et al. 2017).

365 Significant increases were recorded for putrescine in the non-inoculated wines during the
366 aging process, mainly in 2016 vintage. Putrescine can be formed by the decarboxylation of
367 ornithine from the action of bacterial decarboxylase enzymes. In addition, Mangani et al. (2005)
368 have reported that *O. oeni* strains can produce putrescine, not only from ornithine, but also from
369 arginine, if they have the enzyme necessary to degrade arginine to ornithine. Arginine is one of
370 the major amino acids found in grape and wine, and may be the main responsible for the formation
371 of putrescine. The increase observed in the mean putrescine values from 5.13 to 12.0 mg/L was
372 mainly due to the great increase found in four wines (non-inoculated tanks 10-S, 12-S, 19-S and
373 21-S from 2016 vintage). These wines presented a high decrease in arginine (between 40-77%)
374 and ornithine (between 36-94%) during the aging time (data not shown). Therefore, although it
375 was not found a linear correlation between the degradation of the precursor amino acids and the
376 formation of putrescine, the decrease in the concentrations of these amino acids can lead to the

377 formation of this BA by indigenous bacteria that have survived after MLF. Putrescine can reduce
378 sensorial quality at 20-30 mg/L in red wines (Barbieri et al. 2019), but only the non-inoculated
379 wine “tank 10-S” from 2016 vintage exceeded these values.

380 In general, the inoculation did not influence the content of tyramine and cadaverine. The
381 non-inoculated wines from 2017 showed a slightly higher content of tyramine and cadaverine than
382 the inoculated ones. However, these differences are not considered important since the values are
383 low, similar to those found in 2016 vintage and lower than those found in bibliography (Marcobal
384 et al. 2006, Hernández-Orte et al. 2008, Izquierdo-Cañas et al. 2008, EFSA 2011). These BA can
385 be found in grapes, before AF and MLF and their amount is related to grape maturation degree
386 and grape variety.

387 After 12 months of aging in barrels, no statistically significant differences were found in
388 tyramine and cadaverine concentration in both the inoculated and the non-inoculated wines. Only
389 it was observed a slight increase in 2017 wines, although they were very low, being the mean final
390 concentrations of tyramine lower than 2.5 mg/L and of cadaverine lower than 0.61 mg/L. These
391 results are in accordance with those found by Marcobal et al. (2006) and Hernández-Orte et al.
392 (2008) who neither found an increase of these BA after 6 or 12 months of aging in barrels. The
393 mean values of these BA found in our wines were lower than those reported by other authors
394 (Marcobal et al. 2006, Hernández-Orte et al. 2008, Izquierdo-Cañas et al. 2008, EFSA 2011), and
395 were similar to the concentrations reported by the EFSA (2011).

396 Although the toxicological role of BA in wines is still not well-known, it is desirable to
397 avoid their formation and to obtain wines with low BA concentrations, allowing the production of
398 healthier wines and with less allergic reactions (Ancín-Azpilicueta et al. 2019). The values of

399 histamine found in the wines of this study were lower than those reported by the EFSA (2011),
400 with the exception of the wines in which the selected *O. oeni* strain was not implanted. In addition,
401 the concentrations of tyramine, putrescine and cadaverine found in all the studied wines were in
402 the range of the data reported by the EFSA (2011).

403 **General oenological characteristics.** AF was generally conducted by the inoculation of a
404 selected *S. cerevisiae* CECT 12008 strain (Tables 2 to 4), with no influence of the yeast strain on
405 the compounds and parameters analyzed.

406 The LAB inoculation neither produced changes in the oenological parameters of the wines
407 (Table 5). It was only found statistically significant differences between the inoculated wines and
408 the non-inoculated wines from 2018 in lactic and acetic acid concentrations. However, the
409 differences found in acetic acid were very low and the mean values ranged from 0.42 to 0.50 g/L.

410 On the other hand, Mendoza et al. (2011) showed an increase in volatile acidity in co-
411 inoculated musts due to bacterial sugar catabolism. This reaction is more favorable in wines with
412 high pH values. In this study, only the inoculated wines from 2018 vintage showed a slightly higher
413 acetic acid concentration than the non-inoculated ones. However, the acetic acid values of 0.5 g/L
414 do not suppose any negative effect in wines. Acclimatisation of the culture in a medium rich in
415 malic acid, activates malolactic metabolism of the culture and avoids bacterial sugar catabolism
416 during AF. Therefore, it could consider that the selected *O. oeni* strain did not produce an increase
417 in volatile acidity, even when it is inoculated during AF, which agrees with the results obtained by
418 other authors (Massera et al. 2009, Azzolini et al. 2010, Izquierdo-Cañas et al. 2014).

419 Alexandre et al. (2004) and Muñoz et al. (2014) observed that the use of simultaneous
420 inoculation of yeasts and LAB resulted in sluggish AF. However, in this study, the inoculation of

421 the selected *O. oeni* strain did not interfere in the yeast growth and the AF showed adequate
422 development in all the wines. Both, the yeast and the LAB used in this study, were selected from
423 the autochthonous microbiota of the surrounding vineyards, and did not give rise to the negative
424 effects found by other authors, such as the increase in volatile acidity or sluggish AF, since these
425 effects can be strain-dependent.

426 In spite of the high alcohol content of the wines (around 15-15.5 % vol.), the MLF was
427 developed adequately and it was finalized in all wines. So, the selected *O. oeni* strain was adapted
428 gradually to the ethanol content and wine characteristics (Jussier et al. 2006, Zapparoli et al. 2009,
429 Azzolini et al. 2010), being resistant to high alcohol content (> 14.5 % vol) and able to compete
430 with other bacterial species as *Pediococcus* or *Lactobacillus* that could reach the wines from
431 vineyards (Lonvaud-Funel 2001, López et al. 2008, Berbegal et al. 2017).

432 Most of the researches that study the influence of the inoculation of LAB starters have been
433 carried out at laboratory or pilot scale (Zapparoli et al. 2009, Azzolini et al. 2010, Mendoza et al.
434 2011, Izquierdo-Cañas et al. 2014, Muñoz et al. 2014, Ortega-Heras et al. 2014). Therefore, these
435 results are important and significant since this study has been conducted in real winemaking
436 conditions (18,000 kg of grapes in 25,000 L tanks) with sixty-seven wines in three consecutive
437 vintages.

438 Conclusions

439 The selection of a non-histaminogenic *O. oeni* strain together with an adequate adaptation
440 of the culture to wine conditions at the beginning of each harvest, is a good strategy to avoid the
441 formation of BA in red wines, mainly histamine.

442 This practice ensures the prevalence, against the indigenous microbiota, of the selected *O.*
443 *oeni* CECT 9749 strain inoculated that can also be present even in the non-inoculated wines, and
444 allowing reductions in the histamine content of all the wines of the winery.

445 The success of the selected *O. oeni* strain in the wines to avoid the production of BA,
446 mainly histamine, was maintained after 12 months of aging in barrels.

447 This procedure prevents microbial alterations during the moments of fermentation in which
448 wines are not protected with sulfur dioxide. For the same reason, sulfide addition for wine
449 protection is lower, and even can be a tool for making wines with very low or without sulfides.

450 Literature Cited

- 451 Alexandre H, Costello PJ, Remize F, Guzzo J and Guilloux-Benatier M. 2004. *Saccharomyces cerevisiae-*
452 *Oenococcus oeni* interaction in wine: current knowledge and perspectives. *Int J Food Microbiol*
453 93:141-154.
- 454 Ancín-Azpilicueta C, González-Marco A and Jiménez-Moreno N. 2008. Current knowledge about the
455 presence of amines in wine. *Crit Rev Food Sci Nutr* 48:257–275.
- 456 Ancín-Azpilicueta C, Jiménez-Moreno N and Sola-Larrañaga C. 2019. Wine. *In Innovations in Traditional*
457 *Foods*. Galanakis CM (ed), pp. 221-256. Woodhead Publishing, Elsevier Inc.
- 458 Azzolini M, Tosi E, Vagnoli P, Krieger S and Zapparoli G. 2010. Evaluation of technological effects of
459 yeast-bacterial co-inoculation in red table wine production. *Ital J Food Sci* 22:257-263.
- 460 Barbieri F, Montanari C, Gardini F and Tabanelli G. 2019. Biogenic amine production by lactic acid
461 bacteria: A review. *Foods* 8:17.
- 462 Benito S. 2019a. The management of compounds that influence human health in modern winemaking from
463 an HACCP point of view. *Fermentation* 5, 33:20 pages.
- 464 Benito S. 2019b. The impacts of *Schizosaccharomyces* on winemaking. *Appl Microbiol Biotechnol*
465 103:4291-4312.
- 466 Berbegal C, Benavent-Gil Y, Navascués E, Calvo A, Albors C, Pardo I and Ferrer S. 2017. Lowering
467 histamine formation in a red Ribera del Duero wine (Spain) by using an indigenous *O. oeni* strain as a
468 malolactic starter. *Int J Food Microbiol* 244:11–18.
- 469 Berbegal C, Benavent-Gil Y, Pardo I and Ferrer S. 2015. A novel culture medium for *Oenococcus oeni*
470 malolactic starter production. *LWT-Food Sci Technol* 64:25-31.

- 471 Costantini A, García-Moruno E and Moreno-Arribas MV. 2009. Biochemical transformations produced by
472 malolactic fermentation. *In Wine Chemistry and Biochemistry*. Moreno-Arribas MV and Polo MC
473 (ed), pp. 27-57. Springer, New York.
- 474 de Celis M, Ruiz J, Martín-Santamaría M, Alonso A, Marquina D, Navascués E, Gómez-Flechoso M, Belda
475 I and Santos A. 2019. Diversity of *Saccharomyces cerevisiae* yeasts associated to spontaneous and
476 inoculated fermenting grapes from Spanish vineyards. *Lett Appl Microbiol* 68:580-588.
- 477 Del Prete V, Costantini A, Cecchini F, Morassut M and García-Moruno E. 2009. Occurrence of biogenic
478 amines in wine: The role of grapes. *Food Chem* 112:474-481.
- 479 European Food Safety Authority (EFSA). 2011. Panel on Biological Hazards (BIOHAZ). Scientific opinion
480 on risk base control of biogenic amine formation in fermented foods. *EFSA J* 9:2393, pp. 1–93.
- 481 García-Villar N, Hernández-Cassou S and Saurina J. 2007. Characterization of wines through the biogenic
482 amine contents chromatographic techniques and chemometric data analysis. *J Agric Food Chem*
483 55:7453–7461.
- 484 Gardini F, Zogul Y, Suzzi G, Tabanelli G and Zogul F. 2016. Technological factors affecting biogenic
485 amine content in foods: a review. *Front Microbiol* 7:1218.
- 486 Gómez-Alonso S, Hermosín-Gutierrez I and García-Romero E. 2007. Simultaneous HPLC analysis of
487 biogenic amines, amino acids and ammonium ion as aminoenone derivatives in wine and beer samples.
488 *J Agric Food Chem* 55:608–613.
- 489 Hernández-Orte P, Lapeña AC, Peña-Gallego A, Astrain J, Baron C, Pardo I, Polo L, Ferrer S, Cacho J and
490 Ferreira V. 2008. Biogenic amine determination in wine fermented in oak barrels: Factors affecting
491 formation. *Food Res Int* 41:697-706.
- 492 Izquierdo Cañas PM, Gómez Alonso S, Ruiz Pérez P, Seseña Prieto S, García Romero E and Palop Herreros
493 ML. 2009. Biogenic amine production by *Oenococcus oeni* isolates from malolactic fermentation of
494 Tempranillo wine. *J Food Prot* 72:907–910.
- 495 Izquierdo-Cañas PM, García Romero E, Gómez Alonso S, Fernández González M and Palop-Herreros
496 MLL. 2008. Amino acids and biogenic amines during spontaneous malolactic fermentation in
497 Tempranillo red wines. *J Food Comp Anal* 21:731-735.
- 498 Izquierdo-Cañas PM, García Romero E, Pérez-Martín F, Seseña S and Llanos-Palop M. 2014. Sequential
499 inoculation versus co-inoculation in *Cabernet Franc* wine fermentation. *Food Sci Technol Int* 21:1-
500 10.
- 501 Jiménez-Moreno N, Torrea D and Ancín-Azpilicueta C. 2003. Changes in amine concentrations during
502 aging of red wine in oak barrels. *J Agric Food Chem* 51:5732-5737.
- 503 Jussier D, Dubé Morneau A and Mira de Orduña R. 2006. Effect of simultaneous inoculation with yeast
504 and bacteria on fermentation kinetics and key wine parameters of cool climate *Chardonnay*. *Appl*
505 *Environ Microbiol* 72:221-227.

- 506 Ladero V, Calles-Enríquez M, Fernández M and Álvarez MA. 2010. Toxicological Effects of Dietary
507 Biogenic Amines. *Curr Nutr Food Sci* 6:145-156.
- 508 Landete JM, Ferrer S, Polo L and Pardo I. 2005. Biogenic amines in wines from three Spanish regions. *J*
509 *Agric Food Chem* 53:1119–1124.
- 510 Lonvaud-Funel A. 1999. Lactic acid bacteria in the quality improvement and depreciation of wine. *Antonie*
511 *Van Leeuwenhoek* 67:317–331.
- 512 Lonvaud-Funel A. 2001. Biogenic amines in wines: role of lactic acid bacteria. *FEMS Microbiol. Lett*
513 199:9–13.
- 514 López I, López R, Santamaría P, Torres C and Ruiz-Larrea F. 2008. Performance of malolactic fermentation
515 by inoculation of selected *Lactobacillus plantarum* and *Oenococcus oeni* strains isolated from Rioja
516 red wines. *Vitis* 47:123–129.
- 517 Mangani S, Guerrini S, Granchi L and Vincenzini M. 2005. Putrescine accumulation in wine: role of
518 *Oenococcus oeni*. *Curr Microbiol* 51:6-10.
- 519 Marcobal A, Martín-Álvarez PJ, Polo MC, Muñoz R and Moreno-Arribas MV. 2006. Formation of biogenic
520 amines throughout the industrial manufacture of red wine. *J Food Prot* 69:397–404.
- 521 Martuscelli M, Arfelli G, Manetta AC and Suzzi G. 2013. Biogenic amines content as a measure of the
522 quality of wines of Abruzzo (Italy). *Food Chem* 140:590-597.
- 523 Massera A, Soria A, Catania C, Krieger S and Combina M. 2009. Simultaneous inoculation of *Malbec*
524 (*Vitis vinifera*) musts with yeast and bacteria: effects on fermentation performance, sensory and
525 sanitary attributes of wines. *Food Technol Biotechnol* 47:192-201.
- 526 Mendoza LM, Merín MG, Morata VI and Fariás ME. 2011. Characterization of wines produced by mixed
527 culture of autochthonous yeasts and *Oenococcus oeni* from the northwest region of Argentina. *J Ind*
528 *Microbiol Biotechnol* 38:1777–1785.
- 529 Moreno-Arribas MV and Polo MC. 2009. Amino acids and biogenic amines. *In Wine Chemistry and*
530 *Biochemistry*. Moreno-Arribas MV and Polo MC (ed), pp. 163-190. Springer, New York.
- 531 Moreno-Arribas MV, Polo MC, Jorganes F and Muñoz R. 2003. Screening of biogenic amine production
532 by lactic acid bacteria isolated from grape must and wine. *Int J Food Microbiol* 84:117–123.
- 533 Moreno-Arribas MV, Smit AY and du Toit M. 2010. Biogenic amines and the winemaking process. *In*
534 *Managing Wine Quality*. Reynolds AG (ed), pp. 494-522. Woodhead Publishing Limited, Cambridge,
535 UK.
- 536 Muñoz V, Beccaria B and Abreo E. 2014. Simultaneous and successive inoculations of yeasts and lactic
537 acid bacteria on the fermentation of an unsulfited Tannat grape must. *Braz J Microbiol* 45:59–66.
- 538 Nehme N, Mathieu F and Taillandier P. 2010. Impact of the co-culture of *Saccharomyces cerevisiae*-
539 *Oenococcus oeni* on malolactic fermentation and partial characterization of a yeast-derived inhibitory
540 peptidic fraction. *Food Microbiol* 27:150–157.

- 541 Ortega-Heras M, Pérez-Magariño S, Del-Villar-Garrachón V, González-Huerta C, Moro González LC,
542 Guadarrama Rodríguez A, Villanueva Sánchez S, Gallo González R and Martín de la Helguera S.
543 2014. Study of the effect of vintage, maturity degree, and irrigation on the amino acid and biogenic
544 amine content of a white wine from *Verdejo* variety. *J Sci Food Agric* 94:2073-2082.
- 545 Özogul Y and Özogul F. 2019. Chapter 1: Biogenic Amines Formation, Toxicity, Regulations in Food. *In*
546 *Biogenic Amines in Food: Analysis, Occurrence and Toxicity*, pp. 1-17. Royal Society of Chemistry,
547 London.
- 548 Rodríguez-Bencomo JJ and Mehdi A. 2019. Removal of biogenic amines from hydroalcoholic solutions by
549 functionalized silica. *Food Chem* 297:125027.
- 550 Ruiz P, Izquierdo PM, Seseña S and Palop ML. 2010. Selection of autochthonous *Oenococcus oeni* strains
551 according to their oenological properties and vinification results. *Int J Food Microbiol* 137:230–235.
- 552 Smit AY and DuToit M. 2013. Evaluating the influence of malolactic fermentation inoculation practices
553 and ageing on lees on biogenic amine production in wine. *Food Bioprocess Technol* 6:198–206.
- 554 Smit AY, DuToit WJ and DuToit M. 2008. Biogenic amines in wine: understanding the headache. *S Afr J*
555 *Enol Vitic* 29:109–127.
- 556 Soufleros E, Barrios M and Bertrand A. 1998. Correlation between the content of biogenic amines and
557 other wine compounds. *Am J Enol Vitic* 49:266–278.
- 558 Straub BW, Kicherer M, Schilcher SM and Hammes WP. 1995. The formation of biogenic amines by
559 fermentation organisms. *Z Lebensm-Unters- Forsch* 201:79–82.
- 560 Tomera JF. 1999. Current knowledge of the health benefits and disadvantages of wine composition. *Trends*
561 *Food Sci Technol* 10:129-138.
- 562 Zapparoli G, Tosi E, Azzolini M, Vagnoli P and Krieger S. 2009. Bacterial inoculation strategies for the
563 achievement of malolactic fermentation in high alcohol wines. *S Afr J Enol Vitic* 30:49-55.
- 564 Zimatkin SM and Anichtchik OV. 1999. Alcohol-histamine interactions. *Alcohol and Alcohol* 34:141–147.
- 565 Zúñiga M, Pardo I and Ferrer S. 1993. An improved medium for distinguishing between homofermentative
566 and heterofermentative lactic acid bacteria. *Int J Food Microbiol* 18:37–42.
- 567

Table 1 Experimental design of the experiences conducted in the present work during three consecutive vintages. The *O. oeni* CECT 9749 strain was used in the inoculated tanks.

| Vintage | Feature | Inoculated tanks | Non-inoculated tanks | Total |
|---------|-----------------|------------------|----------------------|---------|
| 2016 | Number of tanks | 9 | 13 | 22 |
| | Volume (kg) | 162,000 | 234,000 | 396,000 |
| | Percentage (%) | 40.9 | 59.1 | 100 |
| 2017 | Number of tanks | 10 | 8 | 18 |
| | Volume (kg) | 180,000 | 144,000 | 324,000 |
| | Percentage (%) | 55.6 | 44.4 | 100 |
| 2018 | Number of tanks | 16 | 11 | 27 |
| | Volume (kg) | 288,000 | 198,000 | 486,000 |
| | Percentage (%) | 59.3 | 40.7 | 100 |

Table 2 Biogenic amines of the wines from 2016 vintage analyzed after malolactic fermentation (MLF) and after 12 months of aging in barrels^a.

| Sample | Yeast inoculation | LAB inoculation | LAB implantation (%) | Histamine (mg/L) | | Tyramine (mg/L) | | Putrescine (mg/L) | | Cadaverine (mg/L) | |
|---------------------------|-------------------|-----------------|----------------------|------------------|-------------|-----------------|-----------|-------------------|-----------|-------------------|-----------|
| | | | | After MLF | 12 months | After MLF | 12 months | After MLF | 12 months | After MLF | 12 months |
| 16-Tank 1-I ^b | yes | yes | 100 | nd ^c | 2.00 | 1.85 | 1.38 | 3.10 | 4.03 | 0.35 | 0.43 |
| 16-Tank 2-I | yes | yes | 100 | nd | 1.39 | 1.54 | 1.36 | 3.42 | 2.99 | 0.41 | 0.33 |
| 16-Tank 3-I | yes | yes | 90 | nd | 2.40 | 1.60 | 1.45 | 2.75 | 2.92 | 0.38 | 0.37 |
| 16-Tank 4-I | yes | yes | 80 | 1.14 | 1.62 | 1.42 | 1.20 | 4.31 | 3.79 | 0.37 | 0.33 |
| 16-Tank 5-I | yes | yes | 100 | nd | 1.18 | 1.29 | 1.35 | 2.36 | 2.43 | 0.34 | 0.32 |
| 16-Tank 6-I | yes | yes | 100 | nd | 3.80 | 1.55 | 1.54 | 4.64 | 4.64 | 0.41 | 0.39 |
| 16-Tank 7-I | yes | yes | 100 | nd | 1.00 | 1.51 | 1.31 | 3.72 | 3.43 | 0.37 | 0.36 |
| 16-Tank 8-I | yes | yes | 100 | 1.11 | 1.76 | 1.54 | 1.33 | 3.42 | 3.24 | 0.35 | 0.34 |
| 16-Tank 9-I | yes | yes | 100 | 1.16 | 1.41 | 2.20 | 1.69 | 2.99 | 2.91 | 0.42 | 0.38 |
| MEAN | | | | 0.38 a, y | 1.84 a, z | 1.61 a, y | 1.40 a, y | 3.41 a, y | 3.38 a, y | 0.38 a, y | 0.36 a, y |
| Standard deviation | | | | 0.57 | 0.85 | 0.27 | 0.14 | 0.73 | 0.68 | 0.03 | 0.04 |
| 16-Tank 10-S ^b | yes | no | 100 | 2.64 | 6.88 | 2.22 | 1.91 | 4.10 | 42.5 | 0.50 | 0.44 |
| 16-Tank 11-S | yes | no | 80 | 4.32 | 5.95 | 1.32 | 1.13 | 7.69 | 7.15 | 0.39 | 0.33 |
| 16-Tank 12-S | yes | no | 100 | nd | 2.63 | 2.74 | 2.15 | 5.91 | 19.0 | 0.45 | 0.41 |
| 16-Tank 13-S | no | no | 100 | nd | 3.54 | 1.54 | 1.45 | 3.58 | 3.31 | 0.39 | 0.35 |
| 16-Tank 14-S | yes | no | 100 | 1.50 | 1.70 | 1.00 | 1.05 | 5.46 | 4.39 | 0.29 | 0.28 |
| 16-Tank 15-S | yes | no | 0 | 16.4 | 16.3 | 1.87 | 1.84 | 10.1 | 12.0 | 0.53 | 0.47 |
| 16-Tank 16-S | yes | no | 0 | 18.7 | 17.3 | 1.57 | 1.36 | 6.42 | 11.2 | 0.37 | 0.36 |
| 16-Tank 17-S | yes | no | 0 | 12.8 | 15.8 | 1.73 | 1.45 | 4.40 | 4.55 | 0.35 | 0.32 |
| 16-Tank 18-S | yes | no | 40 | 14.3 | 12.8 | 1.28 | 0.98 | 3.12 | 2.93 | 0.36 | 0.32 |
| 16-Tank 19-S | yes | no | 0 | 15.7 | 17.4 | 1.66 | 1.42 | 3.87 | 17.6 | 0.36 | 0.41 |
| 16-Tank 20-S | yes | no | 0 | 17.1 | 17.4 | 1.89 | 1.66 | 3.01 | 3.30 | 0.39 | 0.39 |
| 16-Tank 21-S | yes | no | 0 | 8.45 | 12.6 | 1.61 | 5.94 | 4.78 | 24.8 | 0.28 | 0.81 |
| 16-Tank 22-S | yes | no | 0 | 10.7 | 12.5 | 1.06 | 1.24 | 4.23 | 3.84 | 0.32 | 0.31 |
| MEAN | | | | 9.42 b, y | 11.0 b, y | 1.65 a, y | 1.81 a, y | 5.13 b, y | 12.0 b, z | 0.38 a, y | 0.40 a, y |
| Standard deviation | | | | 6.96 | 6.02 | 0.47 | 1.29 | 2.01 | 11.59 | 0.07 | 0.14 |

^aMean values with different letters in each compound indicate statistically significant differences at $p < 0.05$. Letters (a, b) indicate differences between the inoculated and the non-inoculated wines in the same aging stage and letters (y, z) indicate differences between aging stages. Histamine values > 10 mg/L (up OIV recommendation) are marked in bold.

^bI: inoculated wines; S: spontaneous MLF or non-inoculated wines. ^cnd: no detected.

Table 3 Biogenic amines of the wines from 2017 vintage analyzed after malolactic fermentation (MLF) and after 12 months of aging in barrels^a.

| Sample | Yeast inoculation | LAB inoculation | LAB implantation (%) | Histamine (mg/L) | | Tyramine (mg/L) | | Putrescine (mg/L) | | Cadaverine (mg/L) | |
|---------------------------|-------------------|-----------------|----------------------|-------------------|-----------|-----------------|-----------|-------------------|-----------|-------------------|-----------|
| | | | | After MLF | 12 months | After MLF | 12 months | After MLF | 12 months | After MLF | 12 months |
| 17-Tank 1-I ^b | yes | yes | 100 | < LQ ^c | 0.28 | 1.15 | 1.42 | 2.96 | 3.46 | 0.31 | 0.35 |
| 17-Tank 2-I | yes | yes | 100 | 0.32 | 1.42 | 0.68 | 1.08 | 2.57 | 3.31 | 0.35 | 0.40 |
| 17-Tank 3-I | yes | yes | 100 | < LQ | 0.41 | 1.60 | 1.98 | 2.87 | 3.54 | 0.37 | 0.41 |
| 17-Tank 4-I | yes | yes | 100 | < LQ | 0.35 | 1.22 | 1.57 | 2.60 | 3.11 | 0.36 | 0.41 |
| 17-Tank 5-I | yes | yes | 100 | nd ^c | < LQ | 1.23 | 1.78 | 2.77 | 3.25 | 0.26 | 0.31 |
| 17-Tank 6-I | yes | yes | 100 | nd | 0.58 | 2.03 | 2.87 | 2.95 | 3.58 | 0.39 | 0.47 |
| 17-Tank 7-I | yes | yes | 100 | < LQ | 0.61 | 1.70 | 1.91 | 2.68 | 3.46 | 0.46 | 0.53 |
| 17-Tank 8-I | yes | yes | 80 | nd | nd | 1.19 | 1.53 | 2.91 | 3.52 | 0.39 | 0.45 |
| 17-Tank 9-I | yes | yes | 100 | nd | 0.27 | 0.85 | 1.14 | 2.86 | 3.43 | 0.36 | 0.40 |
| 17-Tank 10-I | yes | yes | 100 | nd | 0.39 | 1.31 | 1.62 | 3.70 | 4.45 | 0.34 | 0.39 |
| MEAN | | | | < LQ a, y | 0.43 a, z | 1.30 a, y | 1.69 a, z | 2.89 a, y | 3.51 a, z | 0.36 a, y | 0.41 a, z |
| Standard deviation | | | | | 0.40 | 0.39 | 0.51 | 0.32 | 0.36 | 0.05 | 0.06 |
| 17-Tank 11-S ^b | yes | no | 100 | < LQ | 0.51 | 1.62 | 1.87 | 2.48 | 3.24 | 0.47 | 0.53 |
| 17-Tank 12-S | yes | no | 100 | nd | 0.28 | 1.91 | 2.49 | 2.95 | 4.80 | 0.48 | 0.58 |
| 17-Tank 13-S | yes | no | 100 | < LQ | < LQ | 2.00 | 2.17 | 2.80 | 3.74 | 0.54 | 0.56 |
| 17-Tank 14-S | yes | no | 100 | nd | 0.36 | 1.91 | 2.18 | 3.02 | 3.81 | 0.47 | 0.53 |
| 17-Tank 15-S | yes | no | 100 | < LQ | 0.43 | 1.59 | 1.78 | 2.34 | 3.31 | 0.44 | 0.50 |
| 17-Tank 16-S | yes | no | 100 | 0.44 | 0.45 | 1.63 | 1.95 | 3.19 | 3.82 | 0.59 | 0.61 |
| 17-Tank 17-S | no | no | 80 | 0.38 | 0.35 | 1.22 | 1.55 | 3.09 | 3.74 | 0.45 | 0.50 |
| 17-Tank 18-S | no | no | 100 | nd | < LQ | 1.75 | 2.26 | 3.59 | 5.13 | 0.46 | 0.52 |
| MEAN | | | | < LQ a, y | 0.80 a, z | 1.70 b, y | 2.03 b, z | 2.93 a, y | 3.95 b, z | 0.49 b, y | 0.54 b, z |
| Standard deviation | | | | | 0.43 | 0.25 | 0.30 | 0.40 | 0.67 | 0.05 | 0.04 |

^aMean values with different letters in each compound indicate statistically significant differences at $p < 0.05$. Letters (a, b) indicate differences between the inoculated and the non-inoculated wines in the same aging stage and letters (y, z) indicate differences between aging stages.

^bI: inoculated wines; S: spontaneous MLF or non-inoculated wines.

^cLQ: limit of quantification (0.20 mg/L); nd: no detected.

Table 4 Biogenic amines of the wines from 2018 vintage analyzed after malolactic fermentation (MLF) and after 12 months of aging in barrels^a.

| Sample | Yeast inoculation | LAB inoculation | LAB implantation (%) | Histamine (mg/L) | | Tyramine (mg/L) | | Putrescine (mg/L) | | Cadaverine (mg/L) | |
|---------------------------|-------------------|-----------------|----------------------|------------------|-----------|-----------------|-----------|-------------------|-----------|-------------------|-----------|
| | | | | After MLF | 12 months | After MLF | 12 months | After MLF | 12 months | After MLF | 12 months |
| 18-Tank 1-I ^b | yes | yes | 100 | <LQ ^c | <LQ | 1.53 | 1.50 | 3.61 | 3.47 | 0.45 | 0.42 |
| 18-Tank 2-I | yes | yes | 100 | 0.39 | 0.34 | 1.23 | 1.53 | 2.10 | 2.33 | 0.44 | 0.42 |
| 18-Tank 3-I | yes | yes | 100 | 0.42 | 0.40 | 2.31 | 2.32 | 4.49 | 4.26 | 0.56 | 0.52 |
| 18-Tank 4-I | yes | yes | 100 | 0.41 | 0.38 | 1.88 | 1.65 | 5.04 | 4.65 | 0.55 | 0.48 |
| 18-Tank 5-I | yes | yes | 100 | 0.32 | 0.30 | 2.13 | 2.30 | 3.03 | 2.98 | 0.52 | 0.49 |
| 18-Tank 6-I | yes | yes | 100 | 0.62 | 0.59 | 2.49 | 2.27 | 3.35 | 3.13 | 0.59 | 0.53 |
| 18-Tank 7-I | yes | yes | 100 | 0.62 | 0.57 | 2.58 | 2.35 | 3.31 | 3.10 | 0.59 | 0.53 |
| 18-Tank 8-I | yes | yes | 100 | 0.45 | 0.42 | 1.54 | 1.57 | 3.21 | 3.02 | 0.45 | 0.41 |
| 18-Tank 9-I | yes | yes | 100 | 0.50 | 0.51 | 1.56 | 1.61 | 3.21 | 3.18 | 0.44 | 0.42 |
| 18-Tank 10-I | yes | yes | 100 | 0.41 | 0.51 | 2.58 | 1.85 | 4.63 | 3.22 | 0.47 | 0.57 |
| 18-Tank 11-I | yes | yes | 100 | 0.38 | 0.40 | 2.31 | 2.26 | 4.54 | 4.31 | 0.57 | 0.51 |
| 18-Tank 12-I | yes | yes | 100 | 0.42 | 0.41 | 2.15 | 2.17 | 3.74 | 3.62 | 0.56 | 0.51 |
| 18-Tank 13-I | yes | yes | 100 | 0.46 | 0.45 | 2.49 | 2.41 | 3.93 | 3.76 | 0.58 | 0.54 |
| 18-Tank 14-I | yes | yes | 100 | 0.31 | 0.30 | 2.93 | 2.92 | 4.35 | 4.65 | 0.56 | 0.64 |
| 18-Tank 15-I | yes | yes | 100 | 0.58 | 0.60 | 2.65 | 2.44 | 3.46 | 3.29 | 0.59 | 0.50 |
| 18-Tank 16-I | yes | yes | 100 | 0.26 | 0.28 | 3.30 | 3.15 | 4.04 | 3.87 | 0.67 | 0.61 |
| MEAN | | | | 0.42 a, y | 0.42 a, y | 2.23 a, y | 2.14 a, y | 3.75 a, y | 3.55 a, y | 0.54 a, y | 0.51 a, y |
| Standard deviation | | | | 0.12 | 0.12 | 0.56 | 0.49 | 0.75 | 0.66 | 0.07 | 0.07 |
| 18-Tank 17-S ^b | yes | no | 100 | 0.32 | 0.40 | 1.52 | 1.54 | 3.01 | 3.12 | 0.45 | 0.47 |
| 18-Tank 18-S | no | no | 100 | 0.28 | 0.88 | 1.79 | 2.44 | 4.55 | 5.03 | 0.41 | 0.46 |
| 18-Tank 19-S | yes | no | 100 | 0.50 | 0.53 | 1.63 | 1.59 | 3.25 | 3.25 | 0.51 | 0.46 |
| 18-Tank 20-S | no | no | 100 | 0.46 | 0.74 | 1.83 | 1.94 | 4.60 | 4.27 | 0.50 | 0.45 |
| 18-Tank 21-S | no | no | 100 | 0.45 | 0.72 | 2.05 | 2.00 | 4.39 | 4.30 | 0.48 | 0.47 |
| 18-Tank 22-S | no | no | 100 | 0.56 | 0.94 | 2.19 | 2.16 | 4.52 | 4.12 | 0.49 | 0.45 |
| 18-Tank 23-S | yes | no | 100 | 0.41 | 0.75 | 2.58 | 2.55 | 4.63 | 4.50 | 0.47 | 0.45 |
| 18-Tank 24-S | no | no | 100 | 0.50 | 0.61 | 2.70 | 2.74 | 4.71 | 4.37 | 0.40 | 0.42 |
| 18-Tank 25-S | yes | no | 100 | 0.52 | 0.55 | 3.02 | 3.03 | 4.60 | 4.09 | 0.52 | 0.45 |
| 18-Tank 26-S | no | no | 100 | 0.42 | 0.49 | 2.43 | 2.38 | 3.48 | 3.62 | 0.61 | 0.55 |
| 18-Tank 27-S | yes | no | 100 | 0.26 | 0.44 | 2.11 | 1.86 | 4.30 | 3.96 | 0.56 | 0.48 |
| MEAN | | | | 0.42 a, y | 0.64 b, z | 2.17 a, y | 2.20 a, y | 4.19 a, y | 4.06 b, y | 0.49 a, y | 0.47 a, y |
| Standard deviation | | | | 0.10 | 0.18 | 0.47 | 0.47 | 0.62 | 0.56 | 0.06 | 0.03 |

^aMean values with different letters in each compound indicate statistically significant differences at $p < 0.05$. Letters (a, b) indicate differences between the inoculated and the non-inoculated wines in the same aging stage and letters (y, z) indicate differences between aging stages.

^bI: inoculated wines; S: spontaneous MLF or non-inoculated wines.

^cLQ: limit of quantification (0.20 mg/L).

Table 5 Mean values and standard deviation of oenological parameters of the inoculated and the non-inoculated wines from the three vintages ^a.

| | | After alcoholic fermentation | | | | After malolactic fermentation | | | Before aging | |
|---------------------|-----------------|------------------------------|----------------------------------|------------------|-------------------|-------------------------------|-------------------|----------------------|-----------------------------|---------------------------|
| | | pH | Titrateable acidity ^b | Malic acid (g/L) | Lactic acid (g/L) | Lactic acid (g/L) | Acetic acid (g/L) | Alcohol ^b | Free SO ₂ (mg/L) | Molecular SO ₂ |
| 2016 vintage | | | | | | | | | | |
| inoculated | Mean | 3.71 | 6.02 | 1.92 | 0.28 | 1.26 | 0.40 | 14.8 | 52 | 0.80 |
| | SD ^c | 0.10 | 0.40 | 0.16 | 0.09 | 0.14 | 0.04 | 0.5 | 3 | 0.11 |
| non-inoculated | Mean | 3.72 | 5.65 | 1.87 | 0.25 | 1.17 | 0.40 | 14.7 | 51 | 0.71 |
| | SD | 0.07 | 0.47 | 0.19 | 0.15 | 0.11 | 0.04 | 0.4 | 4 | 0.10 |
| 2017 vintage | | | | | | | | | | |
| inoculated | Mean | 3.78 | 5.20 | 1.59 | 0.56 | 1.21 | 0.43 | 15.4 | 49 | 0.66 |
| | SD | 0.07 | 0.42 | 0.59 | 0.16 | 0.10 | 0.06 | 0.4 | 4 | 0.09 |
| non-inoculated | Mean | 3.82 | 5.25 | 1.27 | 0.66 | 1.24 | 0.47 | 15.6 | 51 | 0.64 |
| | SD | 0.11 | 0.31 | 0.46 | 0.34 | 0.13 | 0.09 | 0.2 | 4 | 0.11 |
| 2018 vintage | | | | | | | | | | |
| inoculated | Mean | 3.65 | 6.08 | 1.91 | 0.57 b | 1.60 b | 0.50 b | 15.4 | 49 | 0.56 |
| | SD | 0.10 | 0.59 | 0.64 | 0.39 | 0.23 | 0.10 | 0.3 | 8 | 0.10 |
| non-inoculated | Mean | 3.59 | 5.99 | 2.05 | 0.24 a | 1.31 a | 0.42 a | 15.4 | 47 | 0.57 |
| | SD | 0.10 | 0.48 | 0.66 | 0.07 | 0.18 | 0.07 | 0.3 | 9 | 0.12 |

^aMean values with different letters in each compound and vintage indicate statistically significant differences at $p < 0.05$. Values without letters do not show statistically significant differences.

^bTitrateable acidity (as g/L tartaric acid); alcohol (% vol: mL ethanol/100 mL wine).

^cSD: standard deviation.

American Journal of Enology and Viticulture (AJEV). doi: 10.5344/ajev.2020.20010

AJEV Papers in Press are peer-reviewed, accepted articles that have not yet been published in a print issue of the journal or edited or formatted but may be cited by DOI. The final version may contain substantive or nonsubstantive changes.

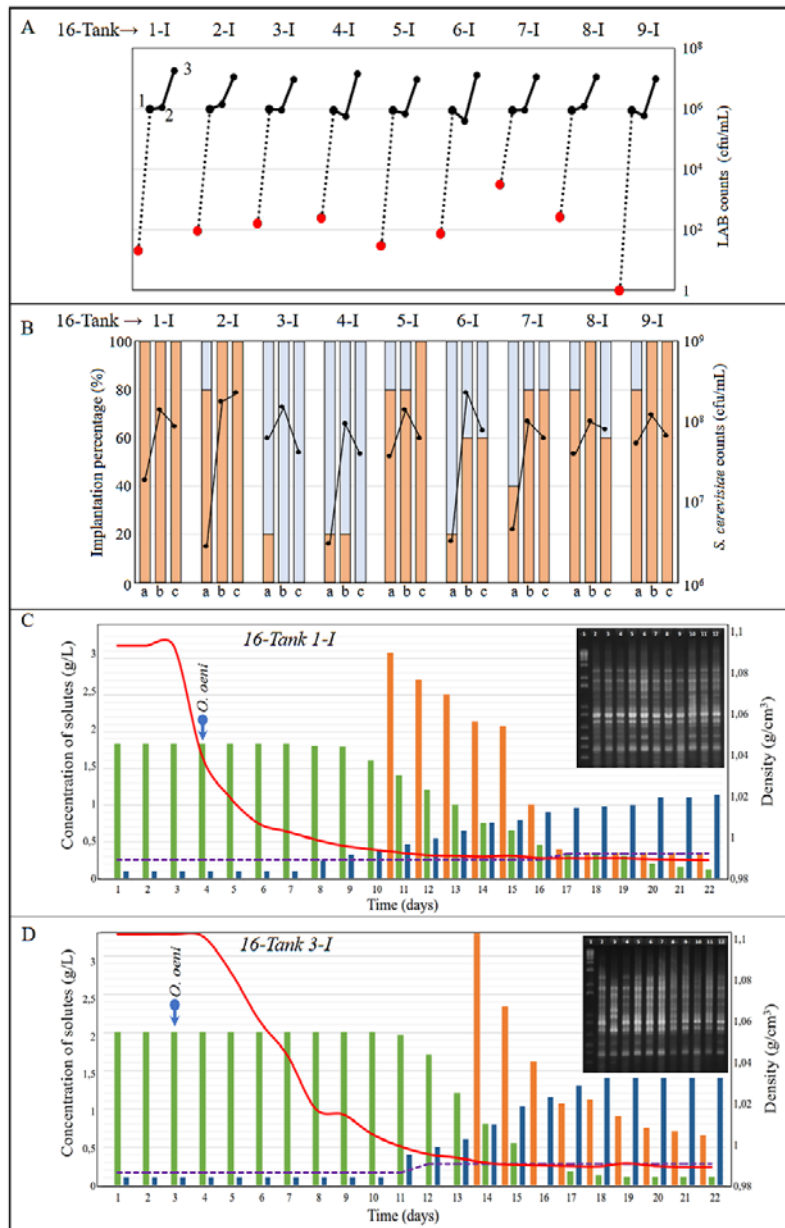


Figure 1 Microbiological control in 9 tanks from the 2016 vintage (Table 2) to exemplify the evolution of the inoculated tanks with the strains *O. oeni* CECT 9749 and *S. cerevisiae* CECT 12008. **(A)** LAB cell counts: three sampling times were used to follow *O. oeni* CECT 9749 development: 1, Inoculation of *O. oeni*; 2, LAB counts at 48h after inoculation; and 3, LAB counts at 2/3 of MLF. Additionally, LAB counts before inoculation were determined (red dots). **(B)** Progress of AF: *S. cerevisiae* cell counts and implantation of the selected yeast strain: three samples, at different stages of the AF, were taken to follow *S. cerevisiae* CECT 12008 development. Orange bars indicate the implantation percentage of the strain CECT 12008, whereas light blue bars indicate the implantation percentage of other wild *S. cerevisiae* strains. **(C, D)** Fermentation kinetics during AF and MLF in two tanks (16-Tank 1-I and 16-Tank 3-I) inoculated with *O. oeni* CECT 9749. Arrows indicate the moment of each inocula addition. Malic acid (green), lactic acid (blue), glucose + fructose (orange), density (red) and acetic acid (violet). G+F concentrations above 3 g/L are not presented in order to facilitate the representation. RAPD-fingerprinting of LAB colonies isolated from each wine tank have been embedded into C and D. Lane 1: molecular weight markers; Lanes 2-11: RAPD band patterns of ten randomly picked colonies; Lane 12: RAPD band pattern of *O. oeni* CECT 9749.

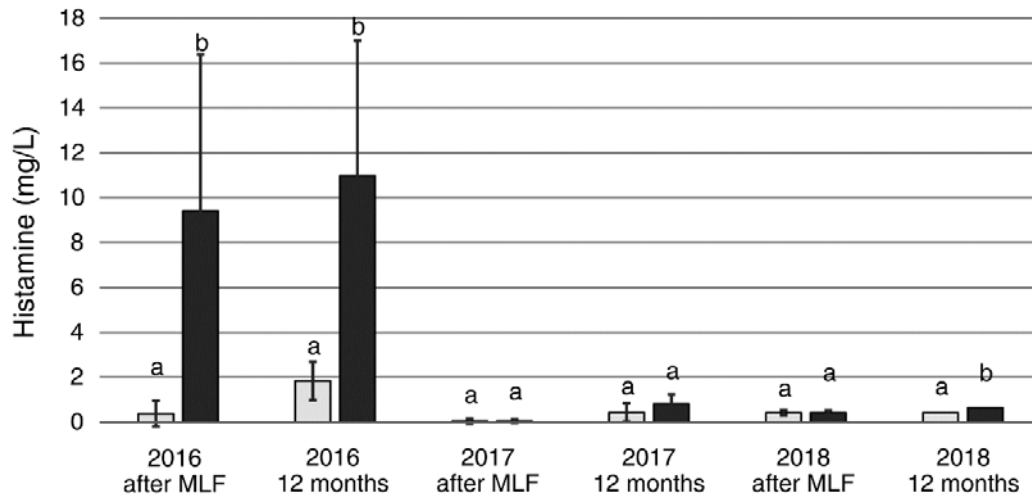


Figure 2 Mean values of histamine (mg/L) in the LAB inoculated (light grey) and the non-inoculated wines (dark grey) from the three vintages studied after MLF and after 12 months of aging in barrels. Mean values with different letters indicate statistically significant differences at $p < 0.05$.