Evaluating the Efficacy of Lysozyme Against Lactic Acid Bacteria Under Different Winemaking Scenarios

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This study evaluated the efficacy of lysozyme in winemaking to control lactic acid bacteria (LAB). In a winery vinification, indigenous LAB were partially and completely inhibited when lysozyme was added to red and white grape must respectively. This result was confirmed by using two selected strains of Lactobacillus brevis and Oenococcus oeni to contaminate the grape must. In the red wine microvinification, the cell population decreased only temporarily and malolactic fermentation terminated at different times, depending on the grape must pH and lysozyme dosage. In the white wine microvinification, cell mortality rates differed according to lysozyme dosage rather than pH values. During the fermentation, lysozyme activity was stable or decreased, depending on the absence or presence of grape must respectively. The study highlighted that lysozyme efficacy is strongly affected by the type of vinification.

INTRODUCTION

Lysozyme is a natural protein with bactericidal activity against Gram-positive bacteria, including lactic acid bacteria (LAB). The use of this enzyme has been shown to be an effective antimicrobial in many foods (Hughes & Johnson, 1987). In the winemaking process, hen egg lysozyme is utilised to control spontaneous LAB in many foods (Hughey & Johnson, 1987). In the winemaking process, hen egg lysozyme is utilised to control spontaneous LAB. The use of this enzyme has been shown to be an effective antimicrobial in many foods (Gerbaux et al., 1999; Delfini et al., 2004; Bartowsky, 2009). The benefits derived from the control of wine spoilage LAB are mainly the reduction in the risk of increased volatile acidity, as well as an increase in biogenic amines. The main technological interest of lysozyme is to reduce the traditional use of sulphur dioxide, which can cause health concerns in consumers (Bartowsky, 2009; Sonni et al., 2009).

It has been ascertained that the efficacy of lysozyme in inhibiting undesirable LAB varies according to species and winemaking conditions (Gerbaux et al., 1999; Gao et al., 2002). Several strains, which are potential detrimental to wine quality, are resistant to this enzyme (Delfini et al., 2004). A lytic cocktail of Streptomyces spp., assayed to control LAB and acetic acid bacteria, was described as a valid alternative to lysozyme because of its higher activity against resistant strains (Blätel et al., 2009). Moreover, lysozyme is less active in red vinification than in white, although it has been demonstrated that its activity is not compromised in the former (Bartowsky & Hensche, 2004; Delfini et al., 2004). Isabel et al. (2009) reported the use of lysozyme as being very beneficial to maintain low histamine levels and ensure Oenococcus oeni implantation in red vinification. Nevertheless, Tirelli and De Novi (2007) proved that lysozyme is unstable in young red wine, thus doubting its effectiveness in controlling malolactic fermentation (MLF). Although lysozyme does not seem to cause important changes in wine aroma, high amounts of this antimicrobial agent could increase the risk of colour instability and the formation of precipitate (Bartowsky & Hensche, 2004). Weber et al. (2007) did not exclude adverse allergic reactions to wines treated with lysozyme, but, for Kirscher et al. (2009), this risk is avoided if the wines are filtered. Moreover, the use of lysozyme involves significant additional costs for winemakers (enzyme purchase, clarification and fining procedures).

Although the use of lysozyme was approved by the Office International de la Vigne et du Vin (OIV) more than a decade ago, some questions remain open. The recommended lysozyme dosage is too generic because it does not consider that lytic activity is strongly affected by contingent winemaking conditions. Hence, from the applied point of view, a more thorough evaluation of its efficacy in defined oenological environments would be informative.

This study evaluated the effects of lysozyme added to grape must for controlling LAB growth. Before alcoholic fermentation, microbial spoilage can be favoured by the vinification of defective grapes together with high pH and temperatures (Ribéreau-Gayon et al., 2006). Two indigenous strains of Lactobacillus brevis and O. oeni were used in grape musts as contaminant bacteria in laboratory-scale winemaking trials. L. brevis is one of the most important wine spoilage LAB, while the uncontrolled growth of indigenous O. oeni strains is often undesirable. The activity of the muramidase was evaluated in terms of must pH and its dosage. The different results from the red and white vinifications are lastly discussed.

MATERIALS AND METHODS

Strains, identification and culture conditions

The strains used in the vinifications were Lactobacillus plantarum DB7 and DB13, L. hilgardii DB19, L. brevis DB12 and Oenococcus oeni DB3, isolated from grape must at the
Centro per la Sperimentazione in Vitivinicoltura, Provincia di Verona (Italy) and deposited in the culture collection of the Dipartimento di Biotecnologie of the University of Verona (Italy). These strains were isolated from fermenting juice to produce Valpolicella and Soave wine, and their identification was carried out by a classification based on morphological and biochemical tests, such as Gram staining, catalase reaction, and carbohydrate fermentations (data not shown). Coccal-shaped isolates were shown through species-specific PCR to belong to *O. oeni*, according to Zapparoli et al. (1998). Rod-shaped isolates were analysed for the identification of *Lactobacillus plantarum* by species-specific PCR according to Torriani et al. (2001). Strains were grown aerobically in FT80 at pH 5.3 (Cavin et al., 1989) and 28°C and used at the late exponential phase for biochemical and molecular assays. Purified LAB cultures were maintained at −80°C in the same medium supplemented with glycerol 25% (v/v).

**Lysozyme**

The lysozyme used in all the experiments was a food grade commercial product (Lallzyme Lyso, Lallemand Inc., Montreal, Canada).

**Analysis of must and wine**

The musts were analysed at the time of crushing, and the wine samples were collected during and after AF and MLF. Ethanol was analysed by NIR spectroscopy using Alcolyzer Wine apparatus (Anton Paar GmbH, Graz, Austria). The sugar content was determined by the Fehling method with automatic titration (Crison, Alella, Spain). Total acidity (expressed as tartaric acid) was determined by titration with 0.1 NaOH to an end point of pH 7.0. Total and free SO2 were determined iodometrically by the Ripper procedure (Ough & Amerine, 1988). L-malic acid and L-lactic acid were quantified using enzyme kits (La Roche, Basel, Switzerland). The total phenolic contents of must-wine were determined with Folin-Ciocalteu reagent, using gallic acid as standard (Singleton & Rossi, 1965). Total anthocyanins were determined by the Glories method (Glories, 1979), using malvidin-3-glucoside as standard. The colour intensity of the must-wine was determined by summing absorbances at 520 and 420 nm of diluted samples (two- to 10-fold) on aqueous solution containing 50 mM tartaric acid and 35 mM NaOH.

**Vinifications in the winery**

Two vinifications, a red and a white, to produce Valpolicella and Soave wine respectively, were carried out to evaluate the effect of lysozyme on indigenous LAB growth. Every vinification was prepared by crushing the grapes and the resulting liquid (juice) was divided into aliquots without previous must clarification. In the case of the red vinification, the solid fraction (grape pomaces) were separated from the juice, mixed and subsequently divided into the proper proportion in order to obtain homogeneous trials. The volume of each single trial was 150 L for both vinifications. The musts were sulphited by adding 50 mg/L SO2. The composition of the red grape must was: pH 3.29, reducing sugars 20.6 °Brix, total acidity 6.00 g/L as tartaric acid, acetic acid 0.09 g/L, and L-malic acid 1.58 g/L. The composition of the white grape must was: pH 3.51, reducing sugars 20.8 °Brix, total acidity 5.30 g/L as tartaric acid, acetic acid 0.03 g/L, and L-malic acid 1.84 g/L. The amount of lysozyme added to the grape musts was 150 mg/L.
The efficacy of lysozyme under winemaking conditions was evaluated using Corvina and Garganega as the red and white grape varieties respectively. After grape crushing, the musts were pasteurised in order to eliminate the indigenous microflora. The pH was corrected to 4.0 to favour rapid cell growth of the pre-cultures. Cells of L. brevis DB12 and O. oeni DB3, grown at 28°C for one week, were harvested and used to inoculate the musts (and pomaces in the red vinification). For both strains the size of inoculation was about 10^4 and 10^7 cfu/ml in the red and white microvinifications respectively. Trials were carried out in duplicate.

The must from Corvina grapes contained reducing sugars at 19.2 °Brix, total acidity of 6.56 g/L as tartaric acid, acetic acid 0.03 g/L and L-malic acid 1.72 g/L. The original pH of 3.19 was corrected to 3.60 in a fraction. The must from Garganega contained reducing sugars at 22.1 °Brix, total acidity of 6.18 g/L as tartaric acid, acetic acid 0.05 g/L and L-malic acid 1.91 g/L. In two fractions the original pH of 3.43 was corrected to 3.10 and 3.70.

The volume of each trial was 5 L for both microvinifications. The AF was induced by inoculating a commercial yeast (VRB, Lallemand) and monitored by ethanol production and sugar consumption. The fermentation was conducted in a wine cellar without controlled temperature (17-20°C).

Lysozyme activity during red grape fermentation

The fermentation of Cabernet Sauvignon grapes was carried out in the absence or presence of pomaces in order to evaluate the effect of wine polyphenols on lysozyme activity. At crushing, the pomaces were separated from the must and both fractions were divided into aliquots of 1.5 L containing only must (–GP) or must plus grape pomaces (+GP). The grape must was composed of the...
following: reducing sugars at 20.7 °Brix, total acidity 3.65 g/L as tartaric acid, and pH 3.63. After pasteurisation, AF was induced by inoculating with yeast (VRB, Lallemand) and monitored at 28°C by weight loss due to CO₂ production. Trials were carried out in triplicate. Must-wine samples were collected at different times of fermentation, centrifuged to eliminate suspended solids and then analysed. To determine the remaining lysozyme activity, samples were diluted five- and 10-fold with buffer phosphate 50 mM pH 6.0 and added to a same volume of cell suspension. This suspension was obtained at the initial stationary phase from cells of \( L. \) brevis DB12 or \( O. \) oeni DB3 grown in FT80 pH 5.3, washed once with NaCl 0.9%, once with buffer phosphate 50 mM pH 6.0, and then resuspended in the same buffer to obtain an OD₆₀₀ of about 1.0. The reaction mix (diluted must-wine sample and cell suspension, 1:1) was incubated at 28°C. Lysozyme activity was quantified by measuring the OD₅₄₀ after 24 hours using a microplate spectrophotometer (Bio-rad, Hercules, CA).

**Lysozyme quantification by HPLC**

HPLC analysis was performed with a Beckman 125S HPLC pump system coupled with a spectrophotometric detector (Beckman Coulter, Inc., Fullerton, CA). The column was a Phenomenex Gemini C18, 5 µm (Phenomenex, Torrance, CA). The solvents used were HPLC-grade acetonitrile (ACN), trifluoroacetic acid (TFA) and deionised distilled water. The HPLC method was used according to Daeschel et al. (2002): eluant A consisted of 1% ACN, 0.2% TFA and 98.8% water, and eluant B consisted of 70% ACN, 0.2% TFA and 29.8% water. The must and wine samples were filtered with MiniSart RC15 syringe filters (Sartorius,
Goettingen, Germany) before injection. The sample volume injected was 20 µL and the flow rate was 1 mL/min. The solvent gradient consisted of four steps: at min 8, solvent B from 0 to 45% in 5 min; at min 13, solvent B from 45 to 47% in 2 min; at min 15, solvent B from 47 to 100% in 15 min; at min 30 the solvent B returns to 0% in 5 min and the run time stops after 35 minutes. Detection was measured by absorbance at 280 nm and the lysozyme retention time was between 19.5 min and 20.7 min. Lysozyme standards were prepared by solubilisation in water and made from 10 to 500 mg/L.

**Statistical treatment of data**

Independent trials were conducted in duplicate for the microvinifications and in triplicate for the lysozyme sensitivity assays and the red grape fermentation; the values reported in the tables and figures are means of two and three determinations ± standard deviation respectively. Data from the red grape fermentation were analysed statistically by applying the T-test using the Graphpad Instat 3.0 program (Graphpad Software Inc., San Diego, CA).

**RESULTS**

**Efficacy of Lysozyme against spontaneous LAB**

The effects of lysozyme on the spontaneous LAB present in the grape must were evaluated during a red and a white vinification to produce Valpolicella and Soave wines respectively. Growth kinetics differed strongly between the two types of vinification (Fig. 1). In the red vinification the cell population decreased just after the addition of lysozyme, but bacterial re-growth was observed within a few days. In the trials with 150 mg/L lysozyme, the LAB rapidly reached the same cell concentration as that measured in the control, while recovery was slower in those with 300 mg/L.

As a consequence, the completion of spontaneous MLF was delayed by 10 days only in the latter, while the addition of 150 mg/L was ineffective (MLF lasted 20 days in the trials without and with 150 mg/L of lysozyme, while the fermentation lasted 30 days in the trials with 300 mg/L). Differently from situation in the red vinification, spontaneous MLF was completely hindered by lysozyme in the white vinification due to the rapid mortality of the LAB, retention time was between 19.5 min and 20.7 min. Lysozyme standards were prepared by solubilisation in water and made from 10 to 500 mg/L.

**Effect of lysozyme in microvinification**

Lysozyme efficacy in controlling the growth of *L. brevis* DB12 and *O. oeni* DB3 strains, inoculated on grape must, varied according to the type of vinification, confirming what has been observed in previous vinifications (Gerbaux et al., 1999).

In the red microvinification the treatment of grape musts at pH 3.20 and 3.60 with lysozyme (200 and 400 mg/L) caused partial mortality of the LAB, followed by a rapid re-growth (Fig. 2). Lysozyme addition did not prevent the total depletion of L-malic acid (Fig. 3), but at pH 3.60, the MLF terminated earlier than at pH 3.20. Microbial analysis revealed that MLF was carried out by *O. oeni* DB3 in all the trials, since its population dominated the wines at the end of fermentation (Fig. 4). Lysozyme caused only a transitory reduction in *O. oeni* DB3, because this species dominated at the end of MLF. In contrast, *L. brevis* DB12 persisted at the beginning of the fermentation, but its population subsequently disappeared (Fig. 4).

In the white microvinification, the addition of 200 and 300 mg/L of lysozyme completely inhibited the LAB population, with similar kinetics irrespective of the pH of the grape must (Fig. 5). In these trials the completion of MLF was hindered, while L-malic acid depletion was completed in the controls (after 38, 34 and 14 days at pH 3.10, 3.40 and 3.70, respectively). Total mortality of the LAB was also observed in grape must at pH 3.10 treated with 100 mg/L of lysozyme, but not in the other musts. *L. brevis* DB12 dominated in all the trials treated with lysozyme, while *O. oeni* DB3 was suppressed after just five days, persisting only in the controls (Fig. 6).

**TABLE 1**

Time (hours) of delay, with respect to the control (without lysozyme), necessary to obtain 50 and 100% growth in FT80 medium pH 5.3 of LAB strains that remained in contact with lysozyme (100, 200 and 500 mg/L) for 16 h.

<table>
<thead>
<tr>
<th></th>
<th>50%</th>
<th>100%</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>100</td>
<td>200</td>
</tr>
<tr>
<td><em>L. plantarum</em> DB7</td>
<td>2 ± 0</td>
<td>7 ± 1</td>
</tr>
<tr>
<td><em>L. plantarum</em> DB13</td>
<td>5 ± 0</td>
<td>5 ± 1</td>
</tr>
<tr>
<td><em>L. hilgardii</em> DB19</td>
<td>5 ± 1</td>
<td>5 ± 1</td>
</tr>
<tr>
<td><em>L. brevis</em> DB12</td>
<td>5 ± 0</td>
<td>20 ± 2</td>
</tr>
<tr>
<td><em>O. oeni</em> DB3</td>
<td>19 ± 2</td>
<td>40 ± 2</td>
</tr>
</tbody>
</table>

1 not grown
Lysozyme activity with and without grape pomaces

The fermentation of red grapes was carried out in the absence and presence of grape pomaces in order to evaluate the effect of wine polyphenols released during alcoholic fermentation on lysozyme activity.

The presence or absence of grape pomaces influenced lysozyme activity against two strains (Table 2). Without grape pomaces (−GP), the lysis of *L. brevis* DB12 and *O. oeni* DB3 decreased by up to 5 and 22% respectively. With grape pomaces (+GP), the lysozyme had less efficacy and the activity was completely inhibited before the end of AF. The quantification of residual lysozyme by HPLC revealed its disappearance in trials containing GP, starting after just six days of fermentation, while no significant variation in its amount was observed in those without GP.

DISCUSSION

The results obtained with the Valpolicella vinification confirmed previous observations by Gerbaux et al. (1999), who showed that 500 mg/L lysozyme added to Pinot Noir before AF was hardly effective or ineffective against LAB. These observations were a starting point for further study on its efficacy under red winemaking conditions.

The red wine microvinification demonstrated that lysozyme addition was not enough to completely control LAB growth. In particular, the differences observed between the pH 3.20 and 3.60 trials were attributed mainly to wine acidity rather that to the presence of lysozyme. Despite this muramidase being more active at high than at low pH, its inhibitory effect decreased under environmental conditions favourable to the growth of spoilage LAB. This evidence was confirmed by the microvinification of white grapes, where the addition of 100 mg/L of lysozyme at pH 3.70 prolonged, but did not stop, the MLF carried out by *L. brevis* DB12 with respect to the control. According to Gao et al. (2002), 125 mg/L of lysozyme was effective against *L. brevis* in Chardonnay (pH 3.8) inoculated with a similar concentration to the one we used in the white microvinification. Taking into account that these high levels of grape contamination (10^7–10^8 cfu/mL) do not occur frequently in standard winemaking conditions, the addition of 100 to 150 mg/L could be enough to prevent any attack of spoilage LAB during white grape fermentation.

The reduction of lysozyme activity in red winemaking is determined by the binding properties of wine phenols to proteins (Hagerman & Butler, 1981). The results of the red grape fermentation trials demonstrated that the presence of grape pomaces affected lytic activity against the *L. brevis* and *O. oeni* strains. A positive evaluation of the efficacy of lysozyme in red winemaking was expressed by Delfini et al. (2004), who observed the behaviour of *O. oeni* and *L. hilgardii* remaining in contact with the enzyme for 16 h in Barbera must. Nevertheless, prolonged contact of the muramidase with wine LAB, as occurs during grape fermentation, determines a gradual reduction in lytic activity due to the binding effects of wine phenols to lysozyme, as we observed in the red grape fermentation. The rate of lysozyme disappearance during AF was in relation to the increase in wine phenols extracted during the vinification. A large decrease in the amount of lysozyme in the red wine (after AF) was observed just a few days after its addition (Daeschel et al., 2002, Tirelli & De Noni, 2007). Hence, the timing of its use assumes relevance to prevent LAB contamination in red winemaking.

Lysozyme added

### TABLE 2

Results of red grape fermentation carried out in absence (−GP) and presence (+GP) of grape pomaces. Lysozyme activity against *L. brevis* DB12 and *O. oeni* DB3 was measured as percent lysis.

<table>
<thead>
<tr>
<th>Lysozyme activity (%)</th>
<th>Days of alcoholic fermentation</th>
<th>0</th>
<th>2</th>
<th>4</th>
<th>6</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>L. brevis</em> DB12</td>
<td>− GP</td>
<td>25 ± 2**</td>
<td>22 ± 3**</td>
<td>22 ± 1**</td>
<td>21 ± 0**</td>
<td>20 ± 1**</td>
</tr>
<tr>
<td></td>
<td>+ GP</td>
<td>13 ± 1</td>
<td>3 ± 1</td>
<td>3 ± 0</td>
<td>0 ± 0</td>
<td>0 ± 0</td>
</tr>
<tr>
<td><em>O. oeni</em> DB3</td>
<td>− GP</td>
<td>58 ± 2**</td>
<td>45 ± 2**</td>
<td>43 ± 2**</td>
<td>40 ± 2**</td>
<td>36 ± 3**</td>
</tr>
<tr>
<td></td>
<td>+ GP</td>
<td>49 ± 1</td>
<td>34 ± 2</td>
<td>11 ± 1</td>
<td>0 ± 0</td>
<td>0 ± 0</td>
</tr>
<tr>
<td>Residual lysozyme (mg/L)</td>
<td>− GP</td>
<td>115 ± 13</td>
<td>136 ± 10**</td>
<td>124 ± 16**</td>
<td>109 ± 14**</td>
<td>135 ± 18**</td>
</tr>
<tr>
<td></td>
<td>+ GP</td>
<td>117 ± 19</td>
<td>49 ± 5</td>
<td>0 ± 0</td>
<td>0 ± 0</td>
<td>0 ± 0</td>
</tr>
<tr>
<td>CO₂ produced (%)</td>
<td>− GP</td>
<td>0 ± 0</td>
<td>19 ± 1</td>
<td>63 ± 3</td>
<td>92 ± 1*</td>
<td>100 ± 1</td>
</tr>
<tr>
<td></td>
<td>+ GP</td>
<td>0 ± 0</td>
<td>17 ± 2</td>
<td>65 ± 1</td>
<td>96 ± 1</td>
<td>100 ± 0</td>
</tr>
<tr>
<td>Total polyphenols (mg/L)</td>
<td>− GP</td>
<td>536 ± 68'</td>
<td>540 ± 85**</td>
<td>384 ± 34**</td>
<td>390 ± 8**</td>
<td>324 ± 28'</td>
</tr>
<tr>
<td></td>
<td>+ GP</td>
<td>864 ± 127</td>
<td>1540 ± 155</td>
<td>2020 ± 14</td>
<td>2200 ± 42</td>
<td>2715 ± 219</td>
</tr>
<tr>
<td>Total anthocyanins (mg/L)</td>
<td>− GP</td>
<td>66 ± 1**</td>
<td>56 ± 8**</td>
<td>50 ± 9**</td>
<td>45 ± 5**</td>
<td>38 ± 8**</td>
</tr>
<tr>
<td></td>
<td>+ GP</td>
<td>206 ± 20</td>
<td>816 ± 5</td>
<td>1043 ± 11</td>
<td>1078 ± 3</td>
<td>967 ± 10</td>
</tr>
<tr>
<td>Colour intensity</td>
<td>− GP</td>
<td>1.5 ± 0.1**</td>
<td>1.4 ± 0.1**</td>
<td>1.3 ± 0.2**</td>
<td>0.9 ± 0.3**</td>
<td>0.8 ± 0.0**</td>
</tr>
<tr>
<td></td>
<td>+ GP</td>
<td>3.9 ± 0.1</td>
<td>9.5 ± 1.0</td>
<td>16.7 ± 0.1</td>
<td>18.0 ± 0.6</td>
<td>19.2 ± 0.2</td>
</tr>
</tbody>
</table>

**1**, **2**, significantly different for p < 0.05 and 0.01 respectively, between −GP and +GP values.
to grape must could cause a significant reduction in cells due to the low polyphenol content, as we observed at the beginning of the Valpolicella vinifications. Nevertheless, the re-growth rate is dependent on parameters that influence LAB growth, such as pH and the wine polyphenol composition responsible for the amount of insoluble enzyme. According to Tirelli and De Noni (2007), enzyme concentrations unable to affect LAB were found in wine when the flavonoid content exceeded 50 mg/L. This concentration is above the typical level of flavonoids in white table wine (40–45 mg/L) and well below the typical level in red table wine (1300–1500 mg/L) (Waterhouse, 2002). The inhibitory effects of wine phenols were clearly demonstrated by the quantification of the residual lysozyme by HPLC, despite the described method (Daeschel et al., 2002) having a detection limit (5–10 mg/L) higher than that of the enzymatic assay (Table 2).

The rapid combination of lysozyme with must colloids and suspended solids determined a significant reduction in the free enzyme just after its addition, as previously demonstrated (Delfini et al., 2004). Then, the further loss of lysozyme during AF in the presence of grape pomaces explained the decrease in activity against LAB strains. At the end of AF, the action of lysozyme against LAB could be strongly compromised when the soluble polyphenols are completely extracted.

LAB have different sensitivities to lysozyme and muramidase effects in winemaking, depending on the species/strain composition of the spontaneous population (Delfini et al., 2004; Blättel et al., 2009).

It was possible to analyse the kinetics of the cell populations of LAB strains that are differently sensitive to lysozyme by inoculating L. brevis DB12 and O. oeni DB3, mimicking a possible microbial contamination occurring during grape crushing. The transitory reduction in the O. oeni DB3 population with respect to L. brevis DB12, observed in the red wine microvinification (Figure 4), was due to its low resistance to the muramidase. The inhibition of lysozyme activity caused by wine phenol did not prevent the re-growth of the O. oeni population. However, the ineffectiveness of lysozyme can be supposed against resistant strains, such as L. plantarum DB7 and DB13 under similar experimental conditions to those utilised in this work.

The benefits of lysozyme usage after AF due to the ensured successful implantation of O. oeni MLF starter culture, as described by Isabel et al. (2009), are probably explained by the concurrence of reduced lytic activity and the different cell concentration between the inoculated culture and indigenous strains. Nevertheless, these benefits could be reduced greatly in the case that a high density of spontaneous malolactic bacteria are present at the end of AF.

In conclusion, this study used a practical approach to highlight the varying efficacy of lysozyme against LAB in red and white vinifications. New information has been presented about the opportunity of using this antimicrobial agent in grape must fermentation. Although lysozyme could be a valid alternative to SO2 for controlling LAB in white winemaking, the same cannot be said for red winemaking. Many factors concur to inhibit lysozyme activity in red winemaking, to the extent that its use could even be without effect. A better understanding is needed of the mechanisms involved in the interaction between lysozyme and red must/wine components (pH, polyphenols, SO2, ethanol, etc.).

From a technological point of view, the elaboration of prediction models will be important to define the relationship between wine composition and lysozyme dosage.

**LITERATURE CITED**


