Effect of Enzyme Addition in the Making of Pedro Ximenez Sweet Wines Using Dynamic Pre-fermentative Maceration

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This work evaluated the effect of two commercial pectolytic enzymes on some oenological parameters before and after enzymatic treatments with pre-fermentative dynamic maceration at room temperature for three hours. The study was conducted on an industrial scale using musts from sun-dried grapes of the Pedro Ximenez variety. Statistical analysis and sensory rating showed that the resulting wines varied significantly in total soluble solids (*Brix) and in the final sensorial controls. On the other hand, the enzyme treatment had no effect on the content of total polyphenols (TPI) and other chemical characteristics of the obtained musts. A higher qualitative level was observed with regard to aroma and gustative quality compared to the control. The results also demonstrate that total juice yield improved after enzyme addition together with dynamic maceration.

INTRODUCTION

Commercially available enzymes have been widely used in the oenological industry in wine-producing countries to improve important characteristics of wines, such as aroma and colour. Furthermore, enzymes are used extensively in the fruit-processing industry and other food sectors (Kashyap et al., 2001). The use of pectolytic enzymes has been shown to be suitable to improve the extraction of colour in red wines (Revilla & González-San José, 2003; Bautista-Ortín et al., 2005), aroma compounds (Canal-Lluberes, 1990; Castro-Vázquez et al., 2002; Cabaroglu et al., 2003) and soluble polysaccharides (Ayestaran et al., 2004; Doco et al., 2007) from the skins and pulp of the grapes. Enzymes may be used to increase the grape must yield during pressing, facilitate the settling of musts, and improve clarification and filtration. Numerous papers have been published on the use of enzymes in the production of white, red and rosé wines (Cruess et al., 1951; Ducruet et al., 1997; Pardo et al., 1999; Salinas et al., 2003). Studies on the use of enzymes in sweet wines are relatively limited in comparison to those performed on white and red wines.

The principal enzyme groups used in winemaking are pectinase, cellulose, hemicellulase, oxidoreductase, protease and β-glycosidase. Evidently, the enzymatic preparations and their principal and secondary activities are key factors in the results obtained in the products (Pimenta-Braz et al., 1998; Guérin et al., 2009). Enzymes in wine production is controlled by Commission Regulation EC 606/09, 2009). The general use of enzymes is regulated by the European Union according to Commission Regulations No. 1331 and 1332 (2008). Recently, the International Organisation of Vine and Wine revised a monograph on enzymatic preparations applied to grapes and their derivatives (OIV, 2009).

Enzyme use in must treatments is related to skin maceration as an additional and complementary process. Skin contact or maceration has been used to increase the aroma in white varietal wines through the extraction of aromatic compounds and non-volatile sugar-bound glycosidic conjugates (Selli et al., 2003), since aroma precursors are located mainly in the grape skin (Mateo & Jimenez, 2000; Ganga et al., 2001; Sánchez Palomo et al., 2006). The skin contact technique also produces an extraction of phenolic components from grape solids in white and red wines (Hernanz et al., 2007) and is the most important process that characterises the elaboration of rosé and red wines. Phenolic compounds from grapes and wines are receiving increasing interest because of diverse health benefits attributed to them (Moreno et al., 2007; Parker et al., 2007; Katalinić et al., 2010).

Pedro Ximenez varietal sweet wines are produced almost exclusively in the Montilla-Moriles Designation of Origin in the south of the province of Córdoba, Spain. It is a region situated close to the Guadalquivir River valley, where average diurnal temperatures in the summer can rise above 40°C and with ambient humidity values of between 30% and 40%. The grapes are carefully hand-picked and spread out on straw mats or on bands of plastic material over a cleared area. Wine production begins with the natural dehydration of the grape bunches by direct exposure to sunlight for about five to 10 days to become raisins, with constant turning-over of the bunches at the optimum degree of maturity. In the traditional Pedro Ximenez process and depending on available equipment, the extraction of must is performed with horizontal presses, but a second step is usually necessary to improve juice yield from the mash and, as a result, high pressure has to be applied using hydraulic presses. The dark must obtained has a high sugar content because of water evaporation from the grapes, producing a very low yield (~29%, w/v, of must). This may vary from year to year. Once the must has been collected, ethanol is added to prevent the start of alcoholic fermentation. After the stabilisation and maturation period, the sweet wine is aged in oak vessels.

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The sun-drying process develops new aromatic compounds such as ethyl acetate, acetoin, γ-butyrolactone, benzyl alcohol, isobutanol, 2-phenylethanol and isoamyl alcohols through anaerobic metabolism (Franco et al., 2004; Márquez et al., 2008) or 2-furaldehyde and 5-methyl-2-furaldehyde by the Maillard reactions of sugars. The wine quality is greatly influenced by aroma and flavour compounds, which come from the grapes, the sun-drying process or from later during wine ageing, because no fermentation is carried out in these traditional sweet wines. Therefore, it is important to take full advantage of the development of aromas and to increase aroma compounds during winemaking of these sweet wines, particularly in the light of the fact that Pedro Ximenez is consider a neutral aromatic variety. Consequently, it is necessary to carefully select commercially available pectolytic enzymes and to know their effect on each grape variety, which may have a different composition according to the specific winemaking conditions applied. In this regard, pre-fermentative dynamic and rapid maceration was used with enzyme addition to enhance the aroma and polyphenol contents in the musts obtained, while improving the pressability of the sun-dried grape mash.

The principal objective of this research was to evaluate, on an industrial scale, the influence standard analytical parameters have on two commercial pectolytic enzymes, compared to a non-enzyme-treated control during novel pre-fermentation dynamic maceration of sun-dried Pedro Ximenez grapes. The sweet wines obtained were characterised by means of chemical analyses and sensorial comparison. Furthermore, this work could contribute to an oenological interest in modifying the traditional winemaking process to improve the quality of these sweet wines.

MATERIALS AND METHODS

Grape samples

The experimental trials were conducted with healthy grapes from Vitis vinifera L. cv. Pedro Ximenez, a white variety, all from the same vineyard situated in the Montilla-Moriles region (Andalucia, Spain). The grapes, harvested manually, were transported in 20 kg plastic boxes to the sun-drying area. The main analytical parameters controlled on reception of the sun-dried grape samples were the pH values (from 4.10 to 4.24) and °Brix (from 43 to 51.8). The lot samples of the sun-dried grapes were divided randomly into three categories (control, enzyme VV and enzyme EM) in the winery. Four different, representative replicates of each type were processed on an industrial scale during one harvest period.

Enzymes used

Two commercial preparations in granular form were used in the experiments. Both enzymes were purified of cinnamyl esterase activity and did not contain genetically modified organisms. The enzymes were diluted in 1:10 parts of distilled water. The dosage tested (2 g per 100 kg of sun-dried grapes) was recommended by the manufacturers. Additional experiments were carried out with an alternative dose at 3 and 4 g per 100 kg of sun-dried grapes.

Vinozym® Vintage FCE (VV) from Novozymes A/S (Bagsvaerd, Denmark) is indicated for the extraction of phenolic and aromatic compounds and has an enzymatic activity of 5 000 FDU/g at 20°C, and contains mainly polygalacturonase activity. This preparation is obtained from Aspergillus niger and A. aculeatus.

Endozym® Muscat (EM) from Pascual Biotech (Paris, France) has an enzymatic activity of 12 500 FDU/g at 20°C. This enzyme is obtained from A. niger and formulated for use in aromatic grape varieties and musts that are difficult to clarify. The preparation contains pectinase (pectinesterase and polygalacturonase), but has other secondary activities such as cellulose and hemicellulase. The following data about its enzymatic activities was supplied by the manufacturer: cellulase (70 units CMC/g), polygalacturonase (4 450 units PG/g), pectinesterase (650 units PE/g) and pectinlyase (10 000 units PL/g).

Reagents

The main reagents used for the analysis were hydrochloric acid 37%, anhydrous sodium carbonate, potassium thiocyanate and sodium hydroxide from Scharlau (Barcelona, Spain). The formaldehyde 37-38% w/w and Folin-Ciocalteau reagent were from Panreac (Barcelona, Spain). The gallic acid came from Fluka (Sigma-Aldrich, Madrid, Spain). All the reagents used were of analytical grade or better.

Experiments

The skins and pulp of crushed but non-destemmed grapes were macerated in their own juice prior to pressing. The enzyme solution was dosed directly during the grape mash and homogenised in the rotating macerator to achieve the correct distribution. A schematic diagram of the process followed is shown in Fig. 1. Dynamic skin maceration was carried out for 3 h in a closed stainless steel macerator tank with about 3 000 kg per tank at room temperature. An automatic cycle was followed for the maceration, according to which the tank turns to each side a number of times, and the time in motion was 88 min (49% of the total period). The temperature of the grape mash was checked before and after the cycle and remained between 28 and 36°C during all experiments. There were no indications of intracellular fermentation during and after the maceration process. The time of maceration and enzyme dosage were previously optimised in laboratory trials. Sulphur dioxide was not added in the maceration trials. Control experiments were treated in the same way except for the enzyme addition. The grape mash was then pressed in horizontal pneumatic presses.

The must samples (mixture of two aliquots from the bottom and upper zone of the tank) were collected in the following phases of the experiment: Sample A once the macerator had been filled, before enzyme addition, and sample B after the dynamic maceration process with the enzyme action. The must samples were taken and analysed immediately. After analysis, these must samples were fortified with ethanol up to 9% v/v, and cooled in a refrigerator at 6°C for approximately 24 h to produce a static sedimentation of solids. The sweet wines obtained were then separated from the precipitated solids and subjected to the final analyses. All the wines were stored in black glass bottles in the dark at 15 ± 3°C until the sensory trials.

Physicochemical determinations

Residual total soluble solids

The residual Brix degree of the grape pomace (skin, seeds, pulp and stems) was obtained by weighing a sample of press-leached pomace after pressing in horizontal pneumatic presses. Thus, four randomly sampled aliquots of 125 g were taken and these
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Sun-dried grapes

Sample A

Enzyme addition

Sample B

Crusher

Maceration dinamic tank

Storage tanks

Pneumatic press

FIGURE 1
General flow scheme of the production process of Pedro Ximenez sweet wine in the experiments conducted on industrial scale.

amounts were combined into a composite sample (500 g) for each experiment. Following this, the pomace sample was submerged in distilled water (1 L) and 1 g of potassium metabisulphite was added. The mixture was then macerated for 72 h in a refrigerator at 5°C; this mash was filtered through filter paper of 200 mm in diameter (Filter-Lab, Barcelona, Spain) and the clear liquid was measured as °Brix by refractometry with a digital refractometer model PR-201a at 20°C (Atago & Co, Kyoto, Japan).

Oenological determinations

The total soluble solids (TSS) of the must were measured as °Brix using the refractometer previously described. Titratable acidity (TA) and pH values were measured by potentiometric titration up to pH = 7 with Titromatic 25 (Crison Instruments, Alella, Spain). The easily assimilable nitrogen content (EAN) was analysed following the method described by Giannessi and Matta (1978), based on the blockage of the aminic function of the amino acids with formaldehyde and acidimetric titration with a potentiometer micro-pH 2002 (Crison Instruments, Alella, Spain). The total polyphenols were determined according to a spectrophotometric method with the Folin-Ciocalteu reagent in alkaline medium (Singleton & Rossi, 1965) using a spectrophotometer UV-1800 (Shimadzu, Kyoto, Japan). A calibration curve was constructed with solutions of gallic acid of known concentration. The total polyphenol index (TPI) is the absorption at 280 nm by direct measurement of the diluted sample at 1:100 with distilled water, using quartz cells of 10 mm path length (Ribéreau-Gayon et al., 2006). Determination of turbidity by the measurement of absorbance at 620 nm. The usual remaining analyses (volatile acidity, reducing sugars, alcoholic strength and iron) were carried out according to the official methods of the European Community (Commission Regulation, 1990). All analyses were done in triplicate and the results were expressed as mean values.

Sensorial analysis

A comparative discrimination sensorial analysis was performed by a triangle test series (ISO, 2004) and by descriptive sensorial tests (Stone & Sidel, 2004) after four months of maturation. In order to quantify the differences, aroma (quality and intensity), flavour (quality and intensity), astringency and herbaceous notes were evaluated, as well as the final sensorial equilibrium to obtain a preferential organisation among the samples. A panel of ten expert wine tasters who were familiar with this type of sweet wine carried out the trials after specific training about the study. The key attributes for this sweet wine had been identified previously, when it is young, to define the quality degrees and criteria to score each parameter. For the discrimination and descriptive tests, two and three replicates respectively were carried out at different times. Sensory trials were performed at 22 ± 1°C in a laboratory room with independent cabins. Samples were numbered with three digits and randomly presented to avoid bias due to the order of presentation. Qualitative references for astringency and herbaceous excess were prepared with a Pedro Ximenez sample as base sweet wine by adding 4 g/L of commercial tannin (AEB Ibérica, Barcelona, Spain) for astringency and 0.05 mL/L of cis-3-hexen-1-ol (Sigma-Aldrich, Madrid, Spain) for herbaceous excess. A point scale, based on the score sheet of the international wine competition of the OIV (1994), was chosen for scoring the parameters to be evaluated. Appearance attributes (colour intensity, hue or cloudiness) were not evaluated as the samples were not considered to be finished sweet wines.

Statistical methods

The statistical tests performed were analysis of variance (one-way ANOVA) of the effect of enzymes on the parameters analysed expressed as differential variation in percentage for the musts. The variation index was defined as the relative difference of the studied parameter according to the equation:

\[ \Delta \text{ parameter} (\%) = \frac{|I - I_0|}{I_0} \times 100 \]

where \( I_0 \) and \( I \) are the values before (sample A) and after enzymatic treatment with dynamic maceration (sample B) respectively. At the same time, a Tukey HSD test at the 5% significance level (\( P < 0.05 \)) was used to separate the means. Multivariate methods were
used to study the results of the sensory analyses. All the statistical tests were done using the software package SPSS® (version 12 for Windows; SPSS Inc., Chicago, USA), and PanelCheck V1.3.2 software (http://www.panelcheck.com) was also used for the sensory analyses.

RESULTS AND DISCUSSION

Effect of enzymes on must parameters

The optimised conditions for enzymatic maceration were applied on an industrial plant scale. The variation in some oenological parameters was studied before and after treatments in these very sweet musts according to the process shown in Fig. 1. The extraction of the must compounds from the grapes into the must is mainly a diffusion process. Thus, the joint action of the enzymes and dynamic maceration helps the better extraction of desirable compounds from the grape mash. The enzymes acted on the skins, stems and the must. The results of the parameter changes in the enzyme-treated and control musts are shown in Table 1. In general, the chemical parameters of the musts produced from enzymatically macerated grapes were very similar to those of the musts from grapes in which maceration took place without enzymes.

Contrary to what was expected, the differential values for the TPI showed similar patterns in all the experiments. The enzyme preparations did not affect the polyphenol contents of the samples under the conditions studied. However, dynamic maceration produced an increase in TPI values in all the treatments, in contrast to traditional elaboration without maceration (data not shown), in despite of the short maceration times assayed. These results suggest that skin contact increases the phenolic contents in musts and wines equally. The similar TPI values in the control and the enzyme-treated musts could be explained by the fact that pre-fermentative maceration in the absence of ethanol results in the relatively low extraction of tannin compounds and flavan-3-ols (González-Manzano et al., 2004; Pinelo et al., 2006). Likewise, other authors have also found techniques least effective in extracting polyphenolic compounds from the pomace (Marais & Rapp, 1988) or reported that the use of enzymes in maceration has not improved colour parameters or polyphenols compounds (Haigh & Gump, 1994; Fernandez-Zurbano et al., 1999; Bautista-Ortín et al., 2005; Alvarez et al., 2006). The TPI values found in the enzymatically macerated samples can be explained because high molecular weight polyphenols could be precipitated during the maceration process. The oxidisable polyphenols in the must can be polymerised and precipitated by chemical or enzymatic oxidation (Colagrande, 1999) by naturally occurring polyphenoloxidases from the musts. The increase of polyphenols and colour compounds in the resulting musts may be related principally to the dehydration levels of the grapes (Serratosa et al., 2008) and with time of maceration. As a result, the formation of brown pigment in musts from sun-dried Pedro Ximenez grapes during the drying process can be explained by the Maillard reaction between sugars and amino acids (Moreno et al., 2007). The extraction capacity of enzymes depends on the composition of the enzyme preparations and their activities, among other factors such as temperature or the

TABLE 1

Data of studied parameters and percentage differences (A) obtained from the analysis of the must samples before and after maceration treatment, mean ± standard deviation (SD). A different letter after the mean values within a row refers to significant differences at $P < 0.05, n = 4$.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Control Before</th>
<th>Control After</th>
<th>Enzyme EM Before</th>
<th>Enzyme EM After</th>
<th>Enzyme VV Before</th>
<th>Enzyme VV After</th>
<th>Significance</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Brix</strong></td>
<td>51.0 ± 0.8</td>
<td>52.0 ± 0.8</td>
<td>45.23 ± 2.0</td>
<td>48.15 ± 1.9</td>
<td>49.20 ± 3.2</td>
<td>52.3 ± 3.8</td>
<td>$P &lt; 0.001$</td>
</tr>
<tr>
<td>$\Delta$ [%]</td>
<td>2.01 ± 0.71 a</td>
<td>6.09 ± 0.67 b</td>
<td>5.89 ± 0.78 b</td>
<td>$P &lt; 0.001$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Titratable acidity (g/L)</strong></td>
<td>5.21 ± 0.2</td>
<td>5.10 ± 0.4</td>
<td>4.39 ± 0.3</td>
<td>4.33 ± 0.7</td>
<td>4.97 ± 0.5</td>
<td>5.11 ± 0.6</td>
<td>$P &lt; 0.001$</td>
</tr>
<tr>
<td>$\Delta$ [%]</td>
<td>-2.33 ± 6.76 a</td>
<td>-2.63 ± 10.16</td>
<td>2.36 ± 6.80 b</td>
<td>$P &lt; 0.001$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Volatile acidity (g/L)</strong></td>
<td>0.33 ± 0.03</td>
<td>0.36 ± 0.03</td>
<td>0.32 ± 0.02</td>
<td>0.36 ± 0.05</td>
<td>0.31 ± 0.05</td>
<td>0.36 ± 0.04</td>
<td>$P &lt; 0.001$</td>
</tr>
<tr>
<td>$\Delta$ [%]</td>
<td>8.41 ± 4.28</td>
<td>10.47 ± 7.96</td>
<td>14.69 ± 4.97</td>
<td>$P &lt; 0.001$</td>
<td></td>
<td></td>
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</tr>
<tr>
<td><strong>EAN (mg/L)</strong></td>
<td>535 ± 35</td>
<td>575 ± 39</td>
<td>444 ± 55</td>
<td>512 ± 38</td>
<td>508 ± 70</td>
<td>559 ± 60</td>
<td>ns</td>
</tr>
<tr>
<td>$\Delta$ [%]</td>
<td>6.74 ± 5.69</td>
<td>13.00 ± 12.44</td>
<td>9.12 ± 7.30</td>
<td>ns</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Reducing sugar (g/L)</strong></td>
<td>607 ± 37</td>
<td>647 ± 38</td>
<td>523 ± 34</td>
<td>581 ± 15</td>
<td>576 ± 54</td>
<td>621 ± 55</td>
<td>ns</td>
</tr>
<tr>
<td>$\Delta$ [%]</td>
<td>6.07 ± 4.37 a</td>
<td>9.91 ± 5.01 b</td>
<td>7.33 ± 4.60 b</td>
<td>$P = 0.012$</td>
<td></td>
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</tr>
<tr>
<td><strong>TPI (UA)</strong></td>
<td>31.78 ± 0.5</td>
<td>51.88 ± 3.3</td>
<td>28.13 ± 2.4</td>
<td>43.65 ± 10.3</td>
<td>31.55 ± 4.5</td>
<td>54.68 ± 7</td>
<td>ns</td>
</tr>
<tr>
<td>$\Delta$ [%]</td>
<td>38.56 ± 3.05</td>
<td>33.74 ± 10.87</td>
<td>42.35 ± 1.75</td>
<td>ns</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Turbidity (UA)</strong></td>
<td>0.22 ± 0.03</td>
<td>0.19 ± 0.03</td>
<td>0.13 ± 0.06</td>
<td>0.15 ± 0.05</td>
<td>0.19 ± 0.03</td>
<td>0.17 ± 0.05</td>
<td>$P = 0.027$</td>
</tr>
<tr>
<td>$\Delta$ [%]</td>
<td>-11.33 ± 10.29 ab</td>
<td>12.90 ± 6.76 b</td>
<td>-16.74 ± 19.75 a</td>
<td>$P = 0.027$</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Expressed as tartaric acid. a Expressed as acetic acid. b Easily assimilable nitrogen. c Total polyphenol index. d Turbidity is the absorbance values (AU) at 620 nm. e Not significant.
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conditions of the treatment. In red varieties, the colour extraction from skins is increased when the enzyme preparations have high cellulase and hemicellulase activities. Thus, the use of enzymes in red varieties increases the polyphenol content and mainly the anthocyanin content of the resulting wines (Sacchi et al., 2005; Kelebek et al., 2007; Romero-Cascales et al., 2007).

At the end of the maceration time, the total soluble solids as degrees Brix and the reducing sugar content of the enzyme-treated musts were higher than in the initial samples. The enzymatic musts had higher reducing sugars than the control. Enzymatic action makes the cell walls more permeable, which causes the extraction of soluble solid matter from the grape mash. The presence of polysaccharide-degrading enzymes (cellulolytic, hemicellulases and polygalacturonase activities) in the preparations contributes to enhancing the reducing sugars in the medium. The highest effect (6.09 ± 0.67%) was observed in the release of Brix when the enzyme EM was added. Previous works have shown that there are big increases in the amounts of arabinose, rhamnose, ribose, xylose and galactose after enzyme treatment (Nee et al., 1999).

With regard to turbidity, the enzyme EM increased the turbidity of the must samples, probably due to the higher activity of this enzyme; in contrast, the turbidity was diminished in the control and enzyme VV (polygalacturonase) treatments. Macerated enzyme treatments produce lower viscosity in musts due to the main activity of pectinase (Ough et al., 1975); the enzyme preparations also contain other side activities in different concentrations, such as pectin glycosidase, which can reduce or enhance this effect. The difference in turbidity of the samples was probably due to the development of the maceration process and the activities of the enzymes used. In previous works, changes in the turbidity of enzyme-treated musts were related to the cultivar or harvest studied (Lao et al., 1996).

The treatments also had a generally negative effect on the titratable acid content of the musts obtained in both the enzymatic and control samples. The dissolution of cations in the must caused by the maceration process possibly produced a reduction in titratable acidity by the neutralisation or salinisation of tartaric acid (Ribereau-Gayon et al., 2006). In contrast, the use of pectolytic enzymes in white Albillo musts showed an increase in titratable acidity (Pérez-Magariño & González-San José, 2000).

The volatile acidity contents were similar in the enzyme-treated musts and in the control, thus were not affected by the enzymatic treatment of the musts, although a slight increase in the values was observed due to maceration. The increase in volatile acidity in the must is considered a negative effect in pre-fermentation treatments because of bacterial contamination. This depends largely on the initial health of the grapes, on the temperature of the must and on the oxygen available during maceration. The values of volatile acidity at the beginning and the end of the maceration step were typical (~0.35 g/L) for healthy grape harvests for this type of wine.

EAN estimates the quantity of assimilable nitrogen in juice, which includes nitrogen from NH₄⁺ and the α-amino acids, except proline. The EAN contents are important for the later alcoholic fermentation of the must. In these wines, fortification up to 9% v/v after pressing prevents alcoholic fermentation. In the traditional winemaking process, little or no fermentation occurs; therefore, this sweet wine is basically grape juice with alcoholic fermentation. However, enzyme addition affected the EAN values in relation to the control samples; there was an increase in EAN after maceration of the samples, including the control. This observation is in agreement with previous works on the Chardonnay variety, which showed an increase in amino acids in macerated musts (Guitart et al., 1997). The optimisation of the process in relation to time of maceration, temperature range, suitable enzyme preparations and doses is essentially to maximise the synergic effects and to obtain the desirable results; the preliminary trials were carried out in the

### TABLE 2

Chemical composition of final sweet wines obtained (control and enzyme-treated wines); values are the mean ± standard deviation (SD). A different letter after the mean values within a row refers to significant differences at P < 0.05, n = 4.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Control sweet wines</th>
<th>EM sweet wines</th>
<th>VV sweet wines</th>
<th>Significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alcoholic strength (% v/v)</td>
<td>9.03 ± 0.1</td>
<td>8.87 ± 0.2</td>
<td>8.91 ± 0.1</td>
<td>ns</td>
</tr>
<tr>
<td>Brix</td>
<td>40.00 ± 0.3 a</td>
<td>42.27 ± 0.9 b</td>
<td>43.93 ± 0.5 c</td>
<td>P = 0.001</td>
</tr>
<tr>
<td>Titratable acidity (g/L)</td>
<td>4.28 ± 0.1 b</td>
<td>3.84 ± 0.1 a</td>
<td>4.15 ± 0.2 ab</td>
<td>P = 0.014</td>
</tr>
<tr>
<td>pH</td>
<td>4.38 ± 0.1</td>
<td>4.42 ± 0.1</td>
<td>4.35 ± 0.1</td>
<td>ns</td>
</tr>
<tr>
<td>Reducing sugars (g/L)</td>
<td>0.37 ± 0.03</td>
<td>0.34 ± 0.02</td>
<td>0.36 ± 0.02</td>
<td>ns</td>
</tr>
<tr>
<td>Volatile acidity (g/L)</td>
<td>438.0 ± 5.3 a</td>
<td>488.3 ± 22.6 b</td>
<td>515.7 ± 9.3 b</td>
<td>P = 0.002</td>
</tr>
<tr>
<td>Titratable acidity (g/L)</td>
<td>29.70 ± 2.0</td>
<td>31.67 ± 2.5</td>
<td>31.33 ± 3.5</td>
<td>ns</td>
</tr>
<tr>
<td>TPP</td>
<td>1,052 ± 42</td>
<td>1,050 ± 140</td>
<td>1,047 ± 72</td>
<td>ns</td>
</tr>
<tr>
<td>Total iron (mg/L)</td>
<td>14.97 ± 0.1 b</td>
<td>13.50 ± 0.9 a</td>
<td>13.67 ± 0.4 ab</td>
<td>P = 0.035</td>
</tr>
<tr>
<td>Iron(II) (mg/L)</td>
<td>7.70 ± 0.1 a</td>
<td>9.50 ± 0.5 b</td>
<td>10.53 ± 0.7 b</td>
<td>P = 0.001</td>
</tr>
<tr>
<td>Iron(III) (mg/L)</td>
<td>7.27 ± 0.1 b</td>
<td>4.00 ± 0.6 a</td>
<td>3.13 ± 0.7 a</td>
<td>P &lt; 0.001</td>
</tr>
</tbody>
</table>

* Expressed as tartaric acid. † Expressed as acetic acid. ‡ AU at 280 nm. § Expressed as mg/L of gallic acid. ‡ Not significant.
Residual degrees Brix of the pressed grape pomace from the control and enzyme-treated musts. Different letters between the column bars refer to significant differences at $P < 0.05, n = 4$.

Comparative sensory scores of the control and enzyme-treated finished sweet wines. The asterisks indicate that the mean differs significantly at $P < 0.01 (**)$ and $0.001 (***)$. 

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Figure 4
Principal components analysis (PCA) of the wine samples and their attributes.

Laboratory (results not shown). The temperature of the grape pomace and the duration of skin contact of the enzymes with the must are key factors that contribute to the correct development of the process. With regard to the reaction temperature, the performance of the pectinases probably was suboptimal taking into account that the temperature reaction was in the range 28 to 36°C. The temperature of the must slows down or speeds up the maceration process with the enzymes. For cold maceration at temperatures of around 20°C, the applied reaction time must be longer and higher enzyme dosages can be necessary. Nevertheless, reduced activities can be compensated for by higher dosages of enzymes or longer maceration times. With the aim of obtaining the maximum effect, additional experiments were carried out by increasing the dose of both the enzymes tested (3 g and 4 g per 100 kg of sun-dried grapes) and keeping the other parameters (room temperature and maceration time). The enzymatic liquefaction of the mash prevented the accurate recording of the data and the must samples could not be analysed. As a consequence, low enzyme dosages were sufficient for complete pectin degradation in these must samples. Low enzyme concentrations were used in the experiments. Much more significant effects may be observed on an industrial scale. It is important to avoid an excess maceration time or enzyme quantity, which could involve complete liquefaction of the mash or pomace with pectin hydrolysis, and this could result in pressing problems. Another factor to consider, although it was not studied, would be the enzyme activity naturally present in these musts that could be increased by the maceration process adding up to the effect of the exogenous enzymes. Thus, taking this into account, the differences found between the control and enzyme-treated musts could be reduced. Nevertheless, to our knowledge, the endogenous enzyme potential and its activities in Pedro Ximenez musts before and after sun-drying process have not been studied.

Effect of enzymes on sweet wines parameters

Table 2 shows the analytical parameters of the final sweet wines obtained. In general, the chemical parameters of the wines produced from enzymatically macerated musts were very similar to those of the wines from control grapes without enzyme addition. All the wines had similar pH (around 4.38), volatile acidity, TPI and total polyphenols. While the degrees Brix and reducing sugars were enhanced due to the enzyme addition and maceration process, a decrease was observed in the titratable acidity of the wines. The reduction in titratable acidity in the macerated musts was due to tartaric acid neutralisation by the potassium liberated from the skins (Ribéreau-Gayon et al., 2006). Both enzymes showed a similar response, thus the analytical values of the final wines were similar. Kelebek (2007) had similar findings for dry red wine from the Oküzgözü variety, which showed no significant differences in density, ethanol, pH, total acidity and reducing sugar, although slight differences were detected in volatile acidity.

The values for total iron varied according to the treatment utilised, thus the total iron in the enzymatic wines was lower (13.50 and 13.67 mg/L) than in the control wines (14.97 mg/L). This could be due to the content of iron(III) in the enzyme-treated wines, which is responsible for the precipitation of colour compounds.
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Manhattan Plots: aroma quality
Manhattan Plots: astringency
Manhattan Plots: herbaceous

Prod. sign. (2-way ANOVA):
- ns
- p<0.05
- p<0.01
- p<0.001

FIGURE 5
Manhattan plots of the attributes indicating variations from PCA on the data of individual assessors. Vertical axes represent the principal components and their cumulative variance. Horizontal axes represent the panel.

linked to metals such as iron. The enzyme-treated macerated musts were kept under oxidised conditions, at which the iron(II) may have been oxidized to iron(III) by dissolved oxygen. During maceration, process differences between the enzyme-treated musts and controls were more pronounced, while these analytical differences diminished in the wines that were obtained.

Residual sugar content
The data on the residual sugar content (expressed as degrees Brix) of the pressed pomace was used as a method for evaluating the extractability of the enzymes with regard to must sugar content (Fig. 2). The results revealed that the control samples were less leached than the pomace from the enzyme samples ($P < 0.001$) and between the two enzymes ($P = 0.035$). Despite the low dosage used, the use of enzymes has produced an improvement in the press capacity during grape processing and prevents extremely hard pressing to obtain a suitable must yield. The residual sugar of the pressed grape pomace confirms the positive effect on the volume of free-run must obtained. Similar findings have been reported previously (Ough & Crowel, 1979; Van Oort & Canal-Llauêres, 2002). The residual sugar content of the pressed pomace was still important even when using enzymes, because of the high TSS concentration (between 43 and 52 °Brix) of the initial sun-dried grapes used. Evidently, the degree of ripening and extractability of the skin cell walls of sun-dried grapes, in combination with the processing methods used, will influence the quality and final characteristics of the resulting wines. Therefore, enzyme addition can have advantageous results in the making of these very sweet wines because it produces an improvement in the effective use of the equipment, and a reduction in the time necessary for processing and in the loss of must. In addition, the use of various enzyme preparations together with different activities in the treatment of the must can produce synergistic additional effects to optimise the use of these products.

Sensory analyses
As a first step, the preliminary results of the triangle tests (results not shown) demonstrate that there was a statistically significant difference ($P < 0.01$), indicating that the control wines could be distinguished from the enzyme-treated wines. In order to determine the magnitude of the difference between the samples, the panel performed a descriptive sensorial analysis. To this end, the sensory attributes evaluated were: astringency and herbaceous notes, general equilibrium, flavour and aroma (quality and intensity) of the samples. The principal differences between the sensory profiles of the three sweet wines are presented in Fig. 3. These results
indicate that the enzymes increased aroma compounds and the general quality impression of these sweet wines. The sensory tests indicate that the quality of the sweet wine is improved when these enzymes are used. Therefore, the flavour intensity and quality, aroma intensity and general equilibrium of the enzyme-treated sweet wines showed significant differences against the control \((P < 0.001)\). All the wines were evaluated in the same way regarding aroma quality, showing no significant defects. The wines made with enzymes were the highest rated on aroma intensity, partly because the typical aromas that predominated were more intense. The sweet wines elaborated with enzyme EM were punctuated better in the sensorial trials. The general sensorial characteristics that dominated corresponded to the peculiarities of these sweet wines when they are young, mostly grape-based aromas with notes of raisins, figs, dates and honey, caramel or toffee. Astringency and herbaceous characters are considered defects in these types of sweet wines that develop due to excessive yields from the pressing process and from the use of non-ripe grapes with high concentrations of C-6 compounds.

Fig. 4 provides a graphic illustration of the principal components analysis (PCA) of the samples and their studied attributes. As can be seen, PC1 explains 99.5% of the variance, while PC2 explains 0.5%. The three types of wines are identified according to their score attributes and by showing how the attributes contributed to the variation in the sensory data. This confirms what was shown in the spider plot (Fig. 3).

The information about the performance of the assessors, which can be provided by PanelCheck program (Tomic et al., 2010), is show in Fig. 5. The Manhattan plots show the variation for one attribute across all assessors. Therefore, every one of the assessors had an explained variance of close to 99% for all the significant attributes with PC1, except with regard to flavour intensity, which shows great variability among the assessors. It also shows the assessors’ performance in the non-significant attributes “aroma quality” and “herbaceous”.

The cellulytic and hemicellulytic secondary activities may contribute to the release of bound aroma precursors from the skins and pulp of dried grapes (Van Oort & Canal-Llabèrè, 2002). In addition, the residual glycosidic activity in the enzymes probably liberates the aroma compounds in the resulting musts during maceration; this effect could be higher if the \(\beta\)-glucosidase from the enzyme preparations was not inhibited by glucose (Maicas & Mateo, 2005). The \(\beta\)-glucosidase action and its efficacy depends on various factors, such as the origin of the enzymes, the concentration of glucose and ethanol and the pH of the medium. In most cases, the pH values of the sun-dried musts were near to the activity optimum, with a pH range between 4.5 and 6.0 (Acuña-Arguelles et al., 1995).

CONCLUSIONS

In summary, the enzyme preparations assayed, combined with dynamic maceration, increased the degrees Brix of the musts obtained, while at the same time improving the total juice yield in the industrial scale experiments. In contrast, these enzyme preparations did not affect the TPI values of the resulting musts and wines significantly. With regard to the effect of the enzymes on the sensory properties, the sweet wines obtained with the enzyme treatments were appreciated more in the sensory evaluations. Therefore, a more detailed study of the enzyme activities of commercial preparations will be of great interest to understand their influence on sensory findings, and to study the impact of new aromatic compounds formed and their contribution to wine aroma and flavour profiles. This technique, pre-fermentative enzyme addition together with short dynamic maceration under controlled conditions, appears to be interesting for the production of fortified sweet wines made with sun-dried grapes, particularly from the Pedro Ximénez variety.

LITERATURE CITED


