

Detection	Linearity range (mg/L)	LOD (mg/L)	LOQ (mg/L)	Repeatability (white wine, N = 5) RSD %	Repeatability (red wine, N = 5) RSD %
UV 280	5 - 250	1.86	6.20	0.62	5.54
FLD	1 - 250	0.18	0.59	0.68	2.37

Table 2

LABELLING OF POTENTIAL ALLERGENIC SUBSTANCES IN EUROPE

Today, eggs and their derivatives are regarded as potential allergens. In the first European directive about the labelling of allergenic ingredients in food (Directive 2000/13/EC), lysozyme used in winemaking was provisionally excluded. In 2007 the European Commission issued the directive 2007/68/EC including the final list of ingredients which must be indicated on the label of food stuffs including alcoholic beverages as they are likely to cause adverse reactions in susceptible individuals. Now lysozyme is part of the list and therefore needs to be determined in the final product. All wines which will be labelled and marketed after 31 May 2009 have to be labelled accordingly. As a consequence the International Organisation for Wine and Vine (OIV) published a standard method³ for measurement of lysozyme in wine, based on a HPLC method with fluorometric detection, developed at the Food Science Department of the University of Bologna⁴.

DETERMINATION OF LYSOZYME BY HPLC

Early methods for the determination of lysozyme used microbiological tests based on enzyme activity. But as these methods were not very robust and very sensitive to phenolic or colloid interactions, alternative HPLC methods were developed. HPLC-UV methods⁵⁻⁷ did not reach the sensitivity required to detect residual lysozyme in marketed wines in the low concentrations, possibly sufficient to cause allergenic responses in sensitive consumers. The new validated HPLC-FLD method is based on the separation of the components on a polymer based TSK-GEL Phenyl-5PW reversed phase column with 1000 Å pore size, which is especially suited for the separation of proteins. Fluorometric detection of the lysozyme resulted in an increased sensitivity compared to UV based HPLC methods. It allows the quantification of lysozyme independently of the enzyme activity.

Fluorometric detection (FLD) of the enzyme without derivatisation is based on the intrinsic fluorescence of its phenylalanine, tyrosine and tryptophan residues. It is dominated by tryptophan because of its higher molar extinction⁸. Egg white lysozyme has six tryptophan residues, resulting in a much higher fluorescence signal intensity compared to the common UV detection at 280 nm. For spectrofluorometric detection an excitation wavelength of 276 nm and an emission wavelength of 345 nm were applied.

HPLC ANALYSIS OF WHITE, ROSE AND RED WINE AFTER THE ADDITION OF LYSOZYME

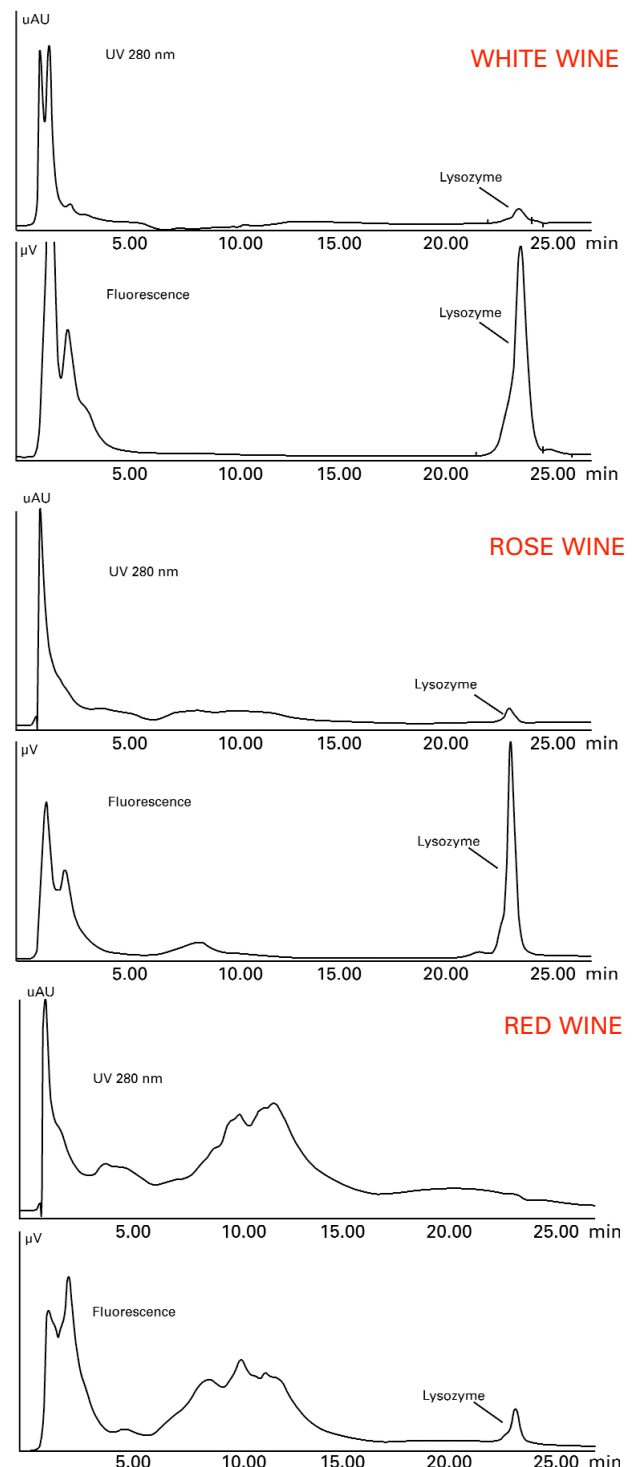


Figure 2

HPLC analysis of three Italian wines after the addition of lysozyme: A: white, B: rosé and C: red

Limits of detection (LOD) and of quantification (LOQ) are up to 11-fold lower compared to UV at 280 nm (Table1). Figure 1 shows the UV280 and fluorescence chromatograms of a model solution spiked with lysozyme to a final concentration of 150 mg/l.

In the original work⁴ it was found that a matter of concern in analysis of lysozyme in wines is the portion of enzyme bound by phenols or other compounds. This happens especially in red wines. Fluctuations related to these interactions can be reduced by acidification of the sample. Sample pre-treatment with HCL enhanced the recovery of lysozyme in red wine. The highest recovery was reached with 10 N HCL at the ratio of 1:10 (HCL : wine). To reach the recoveries reported in the original paper, HPLC injection immediately followed acidification and filtration. Nevertheless repeatability of the method for red wines is worse than for white wines, due to higher concentrations of interfering substances.

We analysed white, red and rosé Italian wine to show the versatility of this method.

MATERIAL & METHODS

HPLC system: Binary pump, 20 µl loop, column oven, photodiode array detector, fluorescence detector (Jasco, Tokyo, Japan)

Column: TSKgel Phenyl-5PW RP reversed phase column, 4.6 mm ID x 7.5 cm L (Tosoh Bioscience, Stuttgart, Germany)

Eluents: A: 98.8 % Water/1% Acetonitrile/0.2% TFA;
B: 70% Acetonitrile/29.8% Water/0.2%TFA

Gradient: 100% A for 3 min, to 35% B in 7 min, maintained for 5 min, to 59.5 % B 0in 12 min, to 100% B in 2 min, maintained for 5 min, to 100% A in 2 min, maintained for 10 min.

Flow rate: 1 ml/min

Temperature: 30°C

Detection: UV @ 280nm, Fluorescence @ 276_{ex}/345_{em} Gain=10

Injection vol.: 20 µl

Sample: Trebbiano di Romagna white wine, Montepulciano di Abruzzo rosé wine, Nero d'Avola red wine, each spiked 24 hours prior to analysis with lysozyme to a final concentration of 160 mg/l.

Sample pre-treatment: Samples were acidified (10:1) with HCL (10 M), filtrated after 5 min. using a 0.22 µm disposable filter and injected directly after filtration.

RESULTS

We analysed three different types of wine from Italy to which lysozyme had been added in the laboratory, with the aim to testing the method. The white wine was a Trebbiano di Romagna, the rosé a Montepulciano di Abruzzo and the red one a Nero d'Avola. The chromatograms are shown in Figure 2. Red wine strongly binded the protein (probably due to the higher presence of tannins and pectins) and a very low amount of free lysozyme was found after 24 hours from the addition. However, free lysozyme tends to progressively increase with time and the final amount of free lysozyme in red wines is usually about 40 - 60% of the added quantity. The asymmetry (fronting) observed in the chromatograms of the samples might be related to the portion of lysozyme which has reacted with other components, such as tannins, sulphur dioxide or pectins. The exact nature of this portion is still unknown and will be the subject of next studies. The peak shape progressively degrades while the lysozyme is interacting with the matrix.

WINE	LYSOZYME (mg/l)
White wine	147
Rose wine	135
Red wine	6.5

➤ Table 2

Results of the HPLC-FLD analysis of three different types of wine spiked with lysozyme (final amount: 160 mg/L)

HPLC-FLD analysis of underivatized hen's egg lysozyme in wines can be applied to all types of wine. It shows a high sensitivity and is a straight forward method for quantification of free amounts of enzyme during winemaking.

REFERENCES:

- F. E. Cunningham et al., *World's Poul. Sci. J.*, 47, 141-163 (1991)
A. Amati et al., *Vitic. Enol. Sci.*, 51, 59-62 (1996)
Resolution OENO 8/2007
C. Riponi et al., *Am. J. Enol. Vitic.*, 58, 405-409 (2007)
R. D. Marchal et al., *J. Agric. Food Chem.*, 48, 3225-3231 (2000)
M. A. Daeschel et al., *Am. J. Enol. Vitic.*, 53, 154-157 (2002)
L. Pellegrino & A. Tirelli, *Int. Dairy J.*, 10, 435-442 (2000)
C. Formoso & L. Foster, *J. Biol. Chem.*, 250, 3738-3745 (1975)

Headquarters

JSB International
Tramstraat 15
5611 CM Eindhoven
T +31 (0) 40 251 47 53
F +31 (0) 40 251 47 58

Zoex Europe
Tramstraat 15
5611 CM Eindhoven
T +31 (0) 40 257 39 72
F +31 (0) 40 251 47 58

Sales and Service

Netherlands
Apolloweg 2B
8239 DA Lelystad
T +31 (0) 320 87 00 18
F +31 (0) 320 87 00 19

Belgium
Grensstraat 7
Box 3 1831 Diegem
T +32 (0) 2 721 92 11
F +32 (0) 2 720 76 22

Germany
Max-Planck-Strasse 4
D-47475 Kamp-Lintfort
T +49 (0) 28 42 9280 799
F +49 (0) 28 42 9732 638

UK & Ireland
Cedar Court,
Grove Park Business Est.
White Waltham, Maidenhead
Berks, SL6 3LW
T +44 (0) 16 288 220 48
F +44 (0) 70 394 006 78

info@go-jsb.com
www.go-jsb.com



With courtesy of **TOSOH**

