## Wine Analysis Made Easy



## ENOLOG

Since its inception in 1981, BioSystems has offered reliable, efficient analytical systems to laboratories worldwide.

Our head offices in Barcelona occupy $16,000 \mathrm{~m}^{2}$ and are staffed by a young, highly qualified team of employees devoted to the research, development, production and marketing of a wide variety of instruments and reagents of utmost quality and outstanding features.

Building on our teamwork and interest in new markets and business units, BioSystems has developed a new system for wine analysis.

Thanks to the high-level scientific expertise of BioSystems staff, we continue to create technologically innovative products that meet the growing needs of laboratories.

We are also conducting ongoing research to improve the procedures used to obtain raw materials and optimize reagent manufacturing.

All research and manufacturing processes are governed by stringent control standards, and our quality systems are regulated by various European and international standards.

At BioSystems we stress the need for innovation and work tirelessly to gain your confidence and loyalty.
We are fully committed and determined to serve you better than anyone else, knowing that this is no easy challenge.

Your satisfaction is the reason for our work and our enthusiasm.

Sincerely yours,


[^0]Managing Director

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## Acetaldehyde

## Enzymatic analysis for acetaldehyde determination

## ADVANTAGES

Stable working reagent for 3 weeks
Ready-to-use dedicated reagent
Liquid calibrator included in the kit


Acetaldehyde is one of the components of the oxidative chain of alcoholic fermentation. Acetaldehyde is also formed in the wine aging process by ethanol oxidation. Acetaldehyde concentration is closely related to $\mathrm{SO}_{2}$ content. This combination is responsible for antioxidant activity.

This is the reason why acetaldehyde is one of the main quality control parameters for wine.

Acetaldehyde in the sample yields NADH (by the following reaction), which can be measured by spectrophotometry.

$$
\text { Acetaldehyde }+\mathrm{NAD}^{+} \xrightarrow{\text { ALDH }} \text { Acetic Acid + NADH }+\mathrm{H}^{+}
$$

| Kit volume: | 50 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 340 nm |
| Limit of linearity: | $200 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $0.1 \mathrm{mg} / \mathrm{L}$ |

Ref. 12820

## ENOLOGQ

## Acetic Acid

Enzymatic method for acetic acid determination

## E

## ADVANTAGES

Stable working reagent for 1 month Ready-to-use dedicated reagent Liquid calibrator included in the kit

Acetic acid is produced during both alcoholic and malolactic fermentations and helps enbance flavors and aromas. When the wine is aerated or remains in contact with air, acetic acid bacteria can multiply, leading to a problem known as "acetic spoilage". The characteristic aroma of this spoilage is due to ethyl acetate.

Acetate in the sample consumes NADH (by the following reaction), which can be measured by spectrophotometry.


| Kit volume: | 100 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 340 nm |
| Limit of linearity: | $1.3 \mathrm{~g} / \mathrm{L}$ |
| Limit of detection: | $0.03 \mathrm{~g} / \mathrm{L}$ |

Ref. 12810

## Ammonia

## Enzymatic method for ammonia determination



## ADVANTAGES

Stable liquid reagent until the expiration date Ready-to-use dedicated reagent Liquid calibrator included in the kit

Low nitrogen levels have been related to slow fermentation or sulfide production. Conversely, high levels can lead to microbial instability and production of ethyl carbonate.

Ammonia in the sample consumes NADH (according to the following reaction), which is then assayed by spectrophotometry.


| Kit volume: | 100 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 340 nm |
| Limit of linearity: | $200 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $3 \mathrm{mg} / \mathrm{L}$ |

Ref. 12809

# Anthocyanins 

Colorimetric analysis for the assay of anthocyanins

## ADVANTAGES

Stable liquid reagent until the expiration date
Ready-to-use dedicated reagent
Liquid calibrator included in the kit
Anthocyanins are the tinted pigments in grapes, with the word coming from the Greek root "antos" (flower) and "kyanos" (blue). These pigments are found in both the skin and the pulp. Anthocyanins can actually have other colorations based on pH and also on their interrelation with other polyphenols. These combinations with other polyphenols can belp further stabilize wine color; hence, bigh interest in analyzing it is warranted.

Anthocyanins are water-soluble pigments that provide the characteristic red color of wine. At 520 nm and under certain conditions, the color is proportional to anthocyanin concentrations. The proposed method determines ionized and ionizable anthocyanins present in the sample. Anthocyanins polymerized with tannins or other compounds cannot be assayed with this method.

| Kit volume: | 100 mL |
| :--- | :--- |
| Method: | End point with reading at 520 nm |
| Limit of linearity: | $1386 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $12 \mathrm{mg} / \mathrm{L}$ |

## Ascorbic Acid

## Enzymatic method for ascorbic acid determination



## ADVANTAGES

Stable working reagent for 10 days
Ready-to-use dedicated reagent
Calibrator included in the kit. Once reconstituted, stable for 20 days

Ascorbic acid is a compound found in ripe grapes at very low levels compared with other acids ( $30-60 \mathrm{mg} / \mathrm{L}$ ). It disappears rapidly when grapes are crushed, leading to early oxidation of must. Due to its reducing properties, ascorbic acid is used as an effective antioxidant and can be used in wines and must at amounts no higher than $100 \mathrm{mg} / \mathrm{L}$.
Ascorbic acid in the sample lowers MTT in the presence of PMS electron carrier, forming dehydroascorbic acid and MTT-formazan that can be assayed by spectrophotometry. In a second determination, ascorbic acid is eliminated from the sample by oxidation to dehydroascorbic acid (ascorbate oxidase [AO]) and other reducing substances (Xred) are measured. The difference between the results obtained from the two reactions is the ascorbic acid concentration. 1,2
Ascorbic Acid + Xred + MT $\xrightarrow{\text { PMS }}$ Dehydroascorbic Acid + Xox + MTT-Formazan Ascorbic Acid $+1 / 2 \mathrm{O}_{2} \xrightarrow{\mathrm{AO}}$ Dehydroascorbic Acid

| Kit volume: | 90 mL |
| :--- | :--- |
| Method: | Two-reagent differential, reading at 560 nm |
| Limit of linearity: | $150 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $1 \mathrm{mg} / \mathrm{L}$ |

## Calcium

## Colorimetric analysis for calcium determination

## ADVANTAGES

Stable two-reagent liquid until the expiration date
Ready-to-use dedicated reagent
Liquid calibrator included in the kit
Calcium is present in wine at concentrations of 6 to 165 $\mathrm{mg} / \mathrm{L}$. The concentrations may be bigher, depending on the soil characteristics, some deacidification processes, etc. Instability due to calcium tartrate appears at 4 to 7 months of fermentation and depends largely on alcohol content, pH , temperature, etc. Controlling these precipitates is key to ensuring wine quality.

Calcium in the sample reacts with 2,7-[bis(2-arsonophenylazo)]-1,8-dihydroxynaphthalene-3,6-disulfonic acid (Arsenazo III). The color increase is directly proportional to the calcium concentration of the sample.

Calcium (Ca) + Arsenazo III $\xrightarrow{\mathrm{pH}=6.5}[$ Ca-(Arsenazo III) $]$

| Kit volume: | 80 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 635 nm |
| Limit of linearity: | $180 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $2 \mathrm{mg} / \mathrm{L}$ |

## Catechins

## Colorimetric analysis for the assay of catechins

## ADVANTAGES

Stable liquid reagent until the expiration date Stable working reagent for 4 months Ready-to-use dedicated reagent Liquid calibrator included in the kit

Catechins are phenolic compounds from the family of flavonoids belonging to the flavanol subgroup. They are reducers and prevent anthocyanin oxidation, keeping them from being precipitated. They are also responsible for the bitterness, astringency, yellow bue, structure and stability of the wine. When catechins are polymerized, they form procyanidins that gradually form complexes with proteins, peptides and polysaccharides as the wine ages. This softens and clarifies wines.

Catechins in the sample react with the chromogen 4-(dimethylamino)-cinnalmaldehyde in the presence of ethanol and an acidic medium, forming a colored complex that can be assayed by spectrophotometry.

| Catechins + DMACA |  |
| :--- | :--- |
| Kit volume: | 100 mL | [Catechins -DMACA]

## Citric Acid

Enzymatic method for citric acid determination

## ADVANTAGES

Stable liquid reagent until the expiration date Stable working reagent for 1 month Ready-to-use dedicated reagent Liquid calibrator included in the kit

Citric acid is an organic acid naturally present in wine that contributes to total wine acidity. Its content is higher in white wine, as the content in red wine drops during malolactic fermentation yielding volatile acids. The permissible legal limit is $1 \mathrm{~g} / \mathrm{L}$, and its concentration must be controlled by wine exporters.
Citrate in the sample yields oxaloacetate due to the action of the enzyme known as lyase citrate. All oxaloacetate from citrate in the sample is converted into L-malic acid by the enzyme L-malate dehydrogenase. This enzyme uses NADH as a coenzyme and is oxidized to NAD+. The disappearance of NADH may be read by spectrophotometry.


| Kit volume: | 50 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 340 nm |
| Limit of linearity: | $400 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $11 \mathrm{mg} / \mathrm{L}$ |

## Color

## Colorimetric analysis for color determination



## ADVANTAGES

Stable liquid reagent until the expiration date Ready-to-use dedicated reagent

Wine color plays a major role in the impression of quality. Color is also an important indicator in many winemaking processes. Regular use of this test allows enologists to document and confirm their own impressions.

The wine sample is diluted in a buffer solution that does not alter color-related properties. Absorbance reading at 420 nm , 520 nm and 620 nm allows the chromatic characteristics to be calculated.

## ENOLOGQ

## Copper

## Colorimetric analysis for copper determination

## ADVANTAGES

Stable liquid reagent until the expiration date
Ready-to-use dedicated reagent
Liquid calibrator included in the kit


Copper is a metal that clearly originates in the process of vinegrowing. The main source is phytosanitary treatments of vineyards to prevent mildew. During harvest, the copper content may be 4 to $6 \mathrm{mg} / \mathrm{L}$. During fermentation its concentration decreases to $0.2-0.3 \mathrm{mg} / \mathrm{L}$ due to the formation of copper sulfides or the presence of yeasts that fix the copper contained in the medium. The International Organisation of Vine and Wine (OIV) has set a maximum acceptable limit of copper of $1 \mathrm{mg} / \mathrm{L}$.

Any copper in the sample reacts with 4-(3,5-dibromo-2-pyridylazo)- N -ethyl-N-sulfopropylaniline (PAESA) sodium salt in acidic medium and in the presence of a reducer. The color increase is directly proportional to the copper concentration of the sample.

$$
\text { Copper (Cu) }+2 \text { 2PAESA } \xrightarrow[\text { reducing agent }]{\text { pH-4. }}\left[\mathrm{Cu}(\text { PAESA })_{2}\right]
$$

| Kit volume: | 100 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 560 nm |
| Limit of linearity: | $7 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $0.4 \mathrm{mg} / \mathrm{L}$ |

Ref. 12814

## $\mathrm{CO}_{2}$

## Enzymatic method for $\mathrm{CO}_{2}$ determination

## ADVANTAGES

Stable liquid reagent until the expiration date
Ready-to-use dedicated reagent
Liquid calibrator included in the kit

Carbon dioxide is a natural gas produced during fermentation that is dissolved in wines. The addition of $\mathrm{CO}_{2}$ during preparation directly affects the aroma and taste of wine and can enhance freshness and acidity in the mouth, softening the sweetness. However, it can also intensify bitterness and astringency.

According to the coupled reactions described below, carbon dioxide $\left(\mathrm{CO}_{2}\right)$ in the sample consumes NADH analogue cofactors that can be assayed by spectrophotometry at 405 nm .


| Kit volume: | 50 mL |
| :--- | :--- |
| Method: | Single-reagent fixed time, <br> reading at 405 nm |
| Linearity: | $1500 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $55 \mathrm{mg} / \mathrm{L}$ |

## D-Gluconic Acid / D-Gluconolactone

## Enzymatic method for D-gluconic acid / D-gluconolactone determination

ADVANTAGES
Stable liquid reagent until the expiration date
Ready-to-use dedicated reagent
Liquid calibrator included in the kit

D-gluconic acid is an indicator of grape deterioration and sanitary condition.

D-gluconic acid in the sample yields NADPH (by the following reaction), which can be measured by spectrophotometry.


D-gluconolactone can be determined according to the same principle after alkaline hydrolysis.


| Kit volume: | 100 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 340 nm |
| Limit of linearity: | $2 \mathrm{~g} / \mathrm{L}$ |
| Limit of detection: | $0.003 \mathrm{~g} / \mathrm{L}$ |

Ref. 12832
Ref. 12811

## D-Glucose / D-Fructose

## Enzymatic method for D-glucose / D-fructose determination

## ADVANTAGES

Stable liquid reagent until the expiration date Working reagent stable until the expiration date
Ready-to-use dedicated reagent
Liquid calibrator included in the kit


This test indicates the best moment for grape harvesting and allows alcoholic fermentation to be monitored. It is widely used to determine the dryness of the wine before bottling.

D-fructose and D-glucose in the sample generate NADH (by the following reaction), which can be measured by spectrophotometry. The configuration of these reagents allows D-glucose/D-fructose (total sugars) to be determined if the enzyme PGI is added or D-glucose to be determined if it is not.

| D-Fructose | HK $\longrightarrow$ Fructose-6-Phosphate + ADP |
| :---: | :---: |
| D-Clucose | HK $\longrightarrow$ Clucose-6-Phosphate + ADP |
| Fructose-6-Pho | $\mathrm{PGI} \longrightarrow$ Clucose-6-Phosphate |
| Clucose-6-Phosphate | $\xrightarrow{\text { G6P-DH }}$ Cluconate-6-Phosphate+NADPH $+\mathrm{H}^{+}$ |
| Kit volume: | 120 mL |
| Method: | Two-reagent differential determination reading at 340 nm |
| Limit of linearity: | $8 \mathrm{~g} / \mathrm{L}$ |
| Limit of detection: | D-Glucose: $0.01 \mathrm{~g} / \mathrm{L}$ D-Glucose/D-Fructose: $0.01 \mathrm{~g} / \mathrm{L}$ |

Ref. 12800

## 新 D-Lactic Acid

## Enzymatic method for D-lactic acid determination

## ADVANTAGES

Stable liquid reagent until the expiration date
Ready-to-use dedicated reagent
Liquid calibrator included in the kit

The excess of bacteria that are producing D-Lactic acid can inhibit alcoholic fermentation, converting some sugars into D-lactic acid. This is one of the main problems in the winemaking process. Levels above $0.3 \mathrm{~g} / \mathrm{L}$ of D-lactic acid indicate bacterial contamination.

D-lactic acid in the sample yields NADH (by the following reaction), which can be measured by spectrophotometry.

D-Lactate $+\mathrm{NAD}^{+} \xrightarrow{\text { D-LDH }}$ Pyruvate + NADH

| Kit volume: | 100 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 340 nm |
| Limit of linearity: | $0.25 \mathrm{~g} / \mathrm{L}$ |
| Limit of detection: | $0.004 \mathrm{~g} / \mathrm{L}$ |

## Free Sulfite

## Colorimetric analysis for free sulfite determination



## ADVANTAGES

Stable liquid reagent until the expiration date Stable working reagent for 9 months Ready-to-use dedicated reagent Liquid calibrator included in the kit

Most sulfur dioxide added to the must or wine combines with different organic compounds. This is the predominant fraction in wine; however, there is another fraction that is not combined, namely, free SO2. Although it is present in lower amounts, its antiseptic and antioxidant properties are superior to those of combined sulfite.

Any free sulfites in the sample react with 4,4'- (4-iminocyclohexa-2,5-dienylidene) methyl) dianiline (pararosaniline) dye in the presence of formaldehyde and in acidic medium. The color increase of the sample is directly proportional to the free sulfite concentration.

$$
\mathrm{SO}_{2}+\text { Pararosaniline } \xrightarrow{\mathrm{pH}=1.0} \text { Pararosaniline-Formaldehyde- } \mathrm{SO}_{2}
$$

| Kit volume: | 400 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 560 nm |
| Limit of linearity: | $150 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $3 \mathrm{mg} / \mathrm{L}$ |

## ЄNOLOGQ

## Clycerol

## Colorimetric analysis

for glycerol determination

## ADVANTAGES

Stable one-reagent liquid until expiration date
Ready-to-use dedicated reagent
Liquid calibrator included in the kit


Glycerol is an indicator of the quality of finished wine and is extremely important for the mouthfeel. High glycerol concentrations add sweetness, body and fullness to the wine.

Glycerol in the sample yields (by the following reaction), a colored complex that is assayed by spectrophotometry.


| Kit volume: | 100 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at $500 \pm 20 \mathrm{~nm}$ |
| Limit of linearity: | $20 \mathrm{~g} / \mathrm{L}$ |
| Limit of detection: | $0.24 \mathrm{~g} / \mathrm{L}$ |

Ref. 12812

## Histamine

## Enzymatic method for the assay of histamine

## ADVANTAGES

Stable liquid reagent until the expiration date
Ready-to-use dedicated reagent
Liquid calibrator included in the kit

## FOODQUALITY

histamine


Histamine is a biogenic amine, a chemical compound formed by the action of microorganisms on amino acids present in foods.

Histamine is present particularly in fermented foods such as wines, cheese and meats, as well as fish. High amounts of bistamine in food can cause organoleptic alterations as well as trigger undesirable effects once consumed and, therefore, bistamine concentrations should be controlled. Although it is true that there are currently no global regulations, acceptable limits for histamine concentrations in wines are around 10 ppm , even though lower amounts are recommended in the case of export to other countries.

By means of the coupled reactions described, histamine in the sample yields a colored complex that is quantitated by spectrophotometry ${ }^{1,2,3}$.


| Kit volume: | 100 mL |
| :--- | :--- |
| Method: | Two-reagent differential, reading at 420 nm |
| Limit of linearity: | $2.1 \mathrm{a} 160 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $2.1 \mathrm{mg} / \mathrm{L}$ |

Ref. 12829

## Iron

Colorimetric analysis for iron determination


## ADVANTAGES

Stable liquid reagent until the expiration date
Ready-to-use dedicated reagent
Liquid calibrator included in the kit

Metal components in wine can originate in grapes or the machinery used to make wine. A bigh iron content can cause clouding due to a lack of solubilization, thus affecting the color and clarity of the wines.

Any iron in the sample reacts with 3-(2-pyridyl)-5,6-bis (4-phenyl-sulfonic)-1,2,4-triazine (ferrozine) sodium salt in acidic medium and in the presence of a reducing agent. The color increase is directly proportional to the iron concentration of the sample.


## L-Lactic Acid

## Enzymatic method for L-lactic acid determination



## ADVANTAGES

Stable liquid reagent until the expiration date Ready-to-use dedicated reagent Liquid calibrator included in the kit

L-lactic acid is the product of the metabolism of malic acid during the malolactic fermentation. L-lactic acid is perceived as less acidic and softer on the palate compared to malic acid.

L-lactic acid in the sample yields NADH (by the following reaction), which can be measured by spectrophotometry.

L-Lactate + NAD $+\xrightarrow{\text { L-LDH }}$ Pyruvate + NADH

| Kit volume: | 100 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 340 nm |
| Limit of linearity: | $3 \mathrm{~g} / \mathrm{L}$ |
| Limit of detection: | $0.02 \mathrm{~g} / \mathrm{L}$ |

Ref. 12817
Ref. 12802


## L-Malic Acid

## Enzymatic method for L-malic acid determination

## ADVANTAGES

Stable liquid reagent until the expiration date Stable working reagent for 4 months Ready-to-use dedicated reagent Liquid calibrator included in the kit

L-malic acid is responsible for the sharply acidic, green apple flavor in wine. It's fermentation yields L-lactic acid and causes perceived acidity to soften.

L-malic acid in the sample yields NADH (by the following reaction), which can be measured by spectrophotometry. The equilibrium of this reaction moves toward L-malic acid formation. The enzyme glutamate-oxaloacetate transaminase (GOT) causes the equilibrium to shift by eliminating oxaloacetate, which is converted into L-aspartate in the presence of L-glutamate.


| Kit volume: | 100 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 340 nm |
| Limit of linearity: | $4 \mathrm{~g} / \mathrm{L}$ |
| Limit of detection: | $0.016 \mathrm{~g} / \mathrm{L}$ |

Ref. 12803

## Polyphenols

## Colorimetric analysis for polyphenols determination

## ADVANTAGES

Stable liquid reagent until the expiration date Ready-to-use dedicated reagent Liquid calibrator included in the kit

Phenol components significantly enhance the antioxidant properties, color and mouthfeel of red wines. The importance of these phenol components in sensory perception requires assay at all stages of the winemaking process.

Any polyphenols in the sample react with Folin-Ciocalteu's reagent in basic medium. The color increase is directly proportional to the polyphenols concentration of the sample.

Polyphenols + Folin-Ciocalteu's Reagent (RF) $\xrightarrow{\text { pH=10.9 }}$ [Polyphenols - FC]

| Kit volume: | 80 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 670 nm |
| Limit of linearity: | $3000 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $60 \mathrm{mg} / \mathrm{L}$ |

Ref. 12815

## Potassium

## Enzymatic method <br> for potassium determination

## ADVANTAGES

Stable liquid reagent until the expiration date Ready-to-use dedicated reagent Liquid calibrator included in the kit

The amount of potassium in grape must varies between 600 and more than $2500 \mathrm{mg} / \mathrm{L}$ in certain varieties of red wine. During véraison, soil potassium moves toward the fruit where it forms soluble potassium bitartrate. Alcohol and low temperatures can reduce its solubility, leading to precipitation.

Potassium in the sample consumes NADH (by the following reaction), which can be measured by spectrophotometry.

| $\text { Phosphoenolpyruvate + ADP } \underset{\mathrm{P}^{+}}{\mathrm{PK}^{+}} \text {Pyruvate + ATP }$ |  |
| :---: | :---: |
| $\text { Pyruvate + NADH + H+ } \xrightarrow{\text { LDH }} \text { Lactate }+ \text { NAD }^{+}$ |  |
| Kit volume: | 80 mL |
| Method: | Two-reagent differential determination reading at 340 nm |
| Limit of linearity: | $1500 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $8 \mathrm{mg} / \mathrm{L}$ |

Ref. 12823

## Primary Amino Nitrogen (PAN)

## Colorimetric analysis <br> for primary amino nitrogen determination

## ADVANTAGES

Stable liquid reagent until the expiration date Stable working reagent for 12 months
Ready-to-use dedicated reagent Liquid calibrator included in the kit

Nitrogen compounds (molecules containing a primary amine nitrogen) in must and wine play a key role in fermentation and the potential of microbial stability.

Any molecules in the sample that contain a primary amino nitrogen react with o-phthaldialdehyde (OPA) in the presence of a reducing agent in basic medium, yielding a chromogen that is assayed spectrophotometrically.

$$
\mathrm{OPA}+\mathrm{NH}_{2} \mathrm{R} \xrightarrow[\text { reducing agent }]{\mathrm{pH}=9.5} \mathrm{OPA}-\mathrm{NH}_{2} \mathrm{R}
$$

| Kit volume: | 100 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 340 nm |
| Limit of linearity: | $400 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $1 \mathrm{mg} / \mathrm{L}$ |

Ref. 12807

## Pyruvic Acid

Enzymatic method<br>for pyruvic acid determination

## ADVANTAGES

Stable liquid reagent until the expiration date
Stable working reagent for 2 months
Ready-to-use dedicated reagent
Liquid calibrator included in the kit

Pyruvic acid is an organic acid naturally present in wine and one of the acids that most influences its body and mouthfeel. Pyruvic acid is a result of the fermentation process and contributes to the organoleptic properties of wine, but must be controlled because selective sulfite-binding shortens the life of the wine.

Pyruvate in the sample yields oxalacetate due to the action of the enzyme known as D-lactate dehydrogenase. This reaction consumes NADH that is oxidized to NAD + and the disappearance of the latter can be measured by spectrophotometry.


| Kit volume: | 100 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 340 nm |
| Limit of linearity: | $400 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $6 \mathrm{mg} / \mathrm{L}$ |

Ref. 12826

## Sucrose / D-Glucose / D-Fructose

## Enzymatic method for sucrose or total sugar determination

## ADVANTAGES

Stable liquid reagent until the expiration date
Stable working reagent for 3 months
Ready-to-use dedicated reagent
Liquid calibrator included in the kit

Precise anabsis of sucrose or total sugar is important for many winecellars in two winemaking operations.

Sparkling wine (cava, champagne, etc.) production: the process may vary according to the method used, but basically consists of adding sucrose once alcobolic fermentation has been carried out in order to achieve a secondary fermentation that produces CO2, which is retained in the wine.

Chaptalization: a technique that consists of adding sucrose to the must when, for various reasons, the grape does not ripen sufficiently and lacks glucoselfructose. This enhances alcoholic fermentation and yields a product with a higher alcohol content. This technique is not approved in all countries.

Sucrose, D-fructose and D-glucose in the sample generate NADPH (by the following reaction), which can be measured by spectrophotometry. The configuration of these reagents allows sucrose or sucrose/D-glucose/D-fructose (total sugars) to be determined.


| Kit volume: | 60 ml |
| :--- | :--- |
| Method: | One-reagent end point or two-reagent <br> differential determination, reading at 340 nm |
| Limit of linearity: | Sucrose $4 \mathrm{~g} / \mathrm{L}$, Total sugar: $8 \mathrm{~g} / \mathrm{L}$ |
| Limit of detection: | Sucrose $0.08 \mathrm{~g} / \mathrm{L}$, Total sugar $0.07 \mathrm{~g} / \mathrm{L}$ |

Ref. 12819


Tartaric Acid
Colorimetric analysis for tartaric acid determination

## ADVANTAGES

Stable liquid reagent until the expiration date
Ready-to-use dedicated reagent
Liquid calibrator included in the kit

Tartaric acid is the main acid of wine that can become insoluble, forming various salts. This acid produces the fruity aromas and freshness of wines and is the most commonly used acidifier.

Any tartaric acid in the sample reacts with vanadium salt in acidic medium, forming a colored complex that is assayed by spectrophotometry.

$$
\text { Tartaric Acid (TART) + Vanadium Salt (V) } \xrightarrow{\mathrm{pH}=2.5}[\text { TART-V }]
$$

| Kit volume: | 100 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 522 nm |
| Limit of linearity: | 0.06 to $6 \mathrm{~g} / \mathrm{L}$ |
| Limit of detection: | $0.06 \mathrm{~g} / \mathrm{L}$ |

## Total Acidity

## Colorimetric analysis

 for the assay of total acidity
## ADVANTAGES

Stable liquid reagent until the expiration date Ready-to-use dedicated reagent Liquid calibrator included in the kit

Total acidity should be determined in mustto ensure goodfermentation, as well as in wine after fermentation because it is a key factor for the storage and stability of wine over time. Low acidity means that microbial alterations and wine with defects and of poorer quality is more likely. Low acidity can cause microbial instability that results in wine defects and overall decrease in quality. Wine should have an adequate total acidity value consistent with the other components to achieve good balance. This value can be between 3 and $7 \mathrm{~g} / \mathrm{L}$.
Total acidity is the sum of assayable acids in wine or must, such as malic acid, tartaric acid, lactic acid, etc., except for carbonic acid and sulfurous acid. Th is reagent determines the total acidity, expressed as $\mathrm{g} /$ Loftartaric acid. Acids in the sample modify the pH in the reaction mixture that, in the presence of the bromothymol blue (BTB) indicator, can be measured spectrophotometrically.

| Kit volume: | 100 ml |
| :--- | :--- |
| Method: | Two-reagent differential determination reading at 620 nm |
| Linearity: | $12 \mathrm{~g} / \mathrm{l}$ |

Ref. 12808

## GNOLOGQ

## Total Sulfite

## Colorimetric analysis for total sulfite determination

## ADVANTAGES

Stable liquid reagent until the expiration date
Ready-to-use dedicated reagent
Liquid calibrator included in the kit

Sulfite is the main preservative of wines and musts, due to its antiseptic properties on yeasts and bacteria; it also has antioxidant properties. According to Council Regulation (EC) No 1493/1999 and Council Regulation (EC) No 1622/2000, the sulfur dioxide content of wine is limited, as it is considered to be a slightly toxic substance from the point of view of its effects on human physiology.

Total sulfites in the sample react with 5-5'-dithio-2-nitrobenzoic (DTNB) acid in basic medium. Cleavage of the disulfide bond (R-S-S-R) of DTNB by a sulfite molecule yields the 5 -mercaptan2 -nitrobenzoate molecule, which absorbs at 405 nm . The color increase of the sample is directly proportional to the total sulfite concentration of the sample.

$$
\mathrm{SO}_{2}+\mathrm{R}-\mathrm{S}-\mathrm{S}-\mathrm{R}(\mathrm{DTNB}) \xrightarrow{\mathrm{pH}=8.2} \mathrm{R}-\mathrm{S}-\mathrm{SO}_{2}+\mathrm{S}-\mathrm{R}
$$

| Kit volume: | 200 mL |
| :--- | :--- |
| Method: | Two-reagent differential determination <br> reading at 405 nm |
| Limit of linearity: | $400 \mathrm{mg} / \mathrm{L}$ |
| Limit of detection: | $1 \mathrm{mg} / \mathrm{L}$ |

## Control Wine (white and red)

Multiparameter control

Control Wine (white and red) is a wine ( $10 \times 5 \mathrm{~mL}$ ) that contains various components at adequate concentrations for quality control in laboratories. The product is designed for intra-laboratory quality control and is supplied with acceptable value intervals.

Traceability is only ensured when the reagents and measurement procedures recommended by BioSystems are used.

| Component | U |
| :--- | :---: |
| Acetic acid | $\mathrm{g} / \mathrm{L}$ |
| Ammonia | $\mathrm{mg} / \mathrm{L}$ |
| D-Gluconic acid | $\mathrm{g} / \mathrm{L}$ |
| D-Clucose/D-Fructose | $\mathrm{g} / \mathrm{L}$ |
| D-Glucose | $\mathrm{g} / \mathrm{L}$ |
| Clycerol | $\mathrm{g} / \mathrm{L}$ |
| L-Lactic acid | $\mathrm{g} / \mathrm{L}$ |
| L-Malic acid | $\mathrm{g} / \mathrm{L}$ |
| Primary Amine Nitrogen | $\mathrm{mg} / \mathrm{L}$ |
| Polyphenols | $\mathrm{mg} / \mathrm{L}$ |
| Tartaric acid | $\mathrm{g} / \mathrm{L}$ |
| Calcium | $\mathrm{mg} / \mathrm{L}$ |
| Citric acid | $\mathrm{mg} / \mathrm{L}$ |
| Histamine | $\mathrm{mg} / \mathrm{L}$ |
| Iron | $\mathrm{mg} / \mathrm{L}$ |



Ref. 12821 Ref. 12822

## Sulfite Control

Sulfite (I and II) Control is a synthetic liquid material that contains stabilized sulfite at adequate concentrations for quality control in laboratories. It does not contain preservatives that could interfere with the measurements.

The concentration values assigned to each level are shown in the attached tables. The values are traceable to the unit of mass. Traceability is ensured only by using the measurement reagents and procedures recommended by BioSystems. The acceptable ranges suggested have been prepared based on prior experience in interlaboratory variability and are provided only as a guideline, as each laboratory should establish its own precision parameters.

| Component | Level | Value | Limits | Unit |
| :--- | :---: | :---: | :---: | :---: |
| Sulfite | । | 40 | $36-44$ | $\mathrm{mg} / \mathrm{L}$ |
| (free and total) | ॥ | 80 | $72-88$ | $\mathrm{mg} / \mathrm{L}$ |

Ref. 12827


## Multical

## Multiparameter calibrator

MULTICAL is a multiparameter calibrator with five synthetic matrix liquid levels $(5 \times 10 \mathrm{~mL})$. It contains various analaytes at adequate concentrations for the calibration of the measurement procedures.

The traceability of the results in samples to reference materials or systems of higher metrological hierarchy is only ensured when the reagents and measurement procedures recommended by BioSystems are used.

| Parameter | U | $\mathbf{1}$ | $\mathbf{2}$ | $\mathbf{3}$ | $\mathbf{4}$ | $\mathbf{5}$ |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| Acetic acid | $\mathrm{g} / \mathrm{L}$ | 0.15 | 0.30 | 0.60 | 0.90 | 1.20 |
| Ammonia | $\mathrm{mg} / \mathrm{L}$ | 23 | 45 | 90 | 135 | 180 |
| Citric acid | $\mathrm{mg} / \mathrm{L}$ | 45 | 90 | 180 | 270 | 360 |
| D-Gluconic acid | $\mathrm{g} / \mathrm{L}$ | 0.20 | 0.40 | 0.80 | 1.20 | 1.60 |
| D-Glucose | $\mathrm{g} / \mathrm{L}$ | 0.90 | 1.80 | 3.60 | 5.40 | 7.20 |
| D-Glucose/D-Fructose | $\mathrm{g} / \mathrm{L}$ | 0.90 | 1.80 | 3.60 | 5.40 | 7.20 |
| Clycerol | $\mathrm{g} / \mathrm{L}$ | 0.113 | 0.225 | 0.450 | 0.675 | 0.900 |
| D-Lactic acid | $\mathrm{mg} / \mathrm{L}$ | 0.028 | 0.056 | 0.113 | 0.169 | 0.225 |
| L-Lactic acid | $\mathrm{g} / \mathrm{L}$ | 0.34 | 0.68 | 1.35 | 2.03 | 2.70 |
| L-Malic acid | $\mathrm{g} / \mathrm{L}$ | 0.45 | 0.90 | 1.80 | 2.70 | 3.60 |
| PAN | $\mathrm{mg} / \mathrm{L}$ | 45 | 90 | 180 | 270 | 360 |
| Total sugar | $\mathrm{g} / \mathrm{L}$ | 0.90 | 1.80 | 3.60 | 5.40 | 7.20 |

Traceability: aqueous reference standard

IONS MULTICAL. 5 levels with 10 mL . Multiparameter calibrator with five synthetic matrix liquid levels that contain various metals at adequate concentrations to calibrate the measurement procedures.

The concentration values assigned to each component and their traceability is ensured by using the reagents and measurement procedures recommended by BioSystems.

| Parameter | U | $\mathbf{1}$ | $\mathbf{2}$ | $\mathbf{3}$ | $\mathbf{4}$ | $\mathbf{5}$ |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| Calcium | $\mathrm{mg} / \mathrm{L}$ | 20.3 | 40.5 | 81.0 | 121.5 | 162.0 |
| Copper | $\mathrm{mg} / \mathrm{L}$ | 0.8 | 1.6 | 3.2 | 4.7 | 6.3 |
| Iron | $\mathrm{mg} / \mathrm{L}$ | 3.4 | 6.8 | 13.5 | 20.3 | 27.0 |
| Potassium | $\mathrm{mg} / \mathrm{L}$ | 34 | 68 | 135 | 203 | 270 |

Traceability: aqueous reference standard

Ref. 12818
Ref. 12841


## Casein

## ELISA method

## ADVANTAGES

Fast, standard method
High sensitivity
Liquid reagent, stable until the expiration date Easy sample preparation

## 部部

BioSystems has addition solutions (spike solutions) to validate the method or to be used as controls.

Ref. 14151 Casein Spike Solution

Casein is an allergenic protein present in cow's milk and dairy products made from cow's milk. The presence of traces of these proteins must be labeled due to the risk it poses to the health of people with allergies, as set forth in the legislation. In addition to foods that naturally contain casein, there may be traces of these proteins in processed foods due to cross-contamination or the use of additives. Caseins are used as clarifier or fining agent in the winemaking process.

Casein reagent is a sandwich enzyme-linked immunosorbent assay (ELISA) for the quantitative analysis of casein traces in samples of wine, juice, cookies, meat products, chocolate and other food products. Any casein in the sample binds to an antibody fixed on the surface of the wells. In a second incubation, another peroxidase-conjugated antibody binds to the casein previously bound to the well. A final incubation with a peroxidase substrate (TMB) develops color based on the presence of the analyte. The reaction is stopped with sulfuric acid or stop solution. The resulting absorbance change is read at 450 nm and is proportional to the casein concentration present in the sample.

| Presentation: | 96 wells |
| :--- | :--- |
| Method: | Sandwich ELISA |
| LOD: | 0.04 ppm |
| Range of measurement: | $0-0.2-0.6-2-6 \mathrm{ppm}$ |

## GNOLOGQ

## "High-Sensitivity" Histamine

## ELSA method

## ADVANTAGES

High sensitivity
Liquid reagent, stable until the expiration date
Easy sample preparation

Histamine is a biogenic amine present in certain food with bigh concentrations of protein or foods exposed to fermentation processes. Histamine is created by certain microorganisms that affect the amino acid histidine. Histamine intake by sensitive individuals produces undesireable effects, such as headaches or skin reactions; hence, it should be controlled.

High-sensitivity ELISA of histamine is a competitive enzymelinked immunoabsorbent assays for the quantitative analysis of histamine in wine, fish, cheese and meat.

Histamine in the sample is quantitatively derivatized to N -acylhistamine by using an acylating reagent. The microplate wells are coated with histamine. In a first incubation, acylated histamine in the sample or reference standard competes with fixed histamine to bind to anti-histamine antibodies.

In a second incubation, a peroxidase-labeled immunoglobulin conjugate binds to antibodies previously bound to the surface of the wells. Lastly, tetramethylbenzidine (TMB) is added to each well as a substrate for the enzyme and, once color develops, the enzyme reaction is stopped with sulfuric acid or stop solution. The product formed is measured at 450 nm and is inversely proportional to histamine concentrations in the sample.

| Presentation: | 96 wells |
| :--- | :--- |
| Method: | Competitive ELISA |
| LOD: | 0.15 ppb |
| Range of measurement: | $0-0.5-1.5-5-15$ and 50 ppb |

Ref. FCE3100

## Lysozyme

## ELISA method

## ADVANTAGES

Fast, standard method
High sensitivity
Liquid reagent, stable until the expiration date
Easy sample preparation


Lysozyme is an allergenic protein contained in eggs and egg products. As set forth by law, the presence of traces of this protein should be labeled due to the risk posed to the health of allergic individuals. In addition to foods that naturally contain lysozyme, there may be traces of this protein in processed foods due to crosscontamination or the use of additives. Lysozyme is used as a preservative in the winemaking process.

Lysozyme reagent is a sandwich enzyme-linked immunosorbent assay (ELISA) for the quantitative analysis of casein traces in wine and cheese samples. The lysozyme in the sample binds to an immobilized antibody on the surface of the wells. In a second incubation, another peroxidase-conjugated antibody binds to the lysozyme previously bound to the well. A final incubation with a peroxidase substrate (TMB) develops color based on the presence of the analyte. The reaction is stopped with sulfuric acid or stop solution. The resulting absorbance change is read at 450 nm and is proportional to the lysozyme concentration present in the sample.

| Presentation: | 96 wells |
| :--- | :--- |
| Method: | Sandwich ELISA |
| LOD: | 2 ppb |
| Range of measurement: | $0-25-50-100-250 \mathrm{ppb}$ |

## Ovalbumin

## ELISA method

## ADVANTAGES

Fast, standard method
High sensitivity
Liquid reagent, stable until the expiration date
Easy sample preparation


BioSystems has addition solutions (spike solutions) to validate the method or to be used as controls.

Ref. 14154 Ovoalbumin Spike Solution

Ovalbumin is an allergenic protein contained in eggs and egg products. As set forth by law, the presence of traces of this protein should be labeled due to the risk posed to the health of allergic individuals. In addition to foods that naturally contain ovalbumin, there may be traces of this protein in processed foods due to cross-contamination or the use of additives. Ovalbumin is used as a clarifier finding agent in the winemaking process.

Ovalbumin reagent is a sandwich enzyme-linked immunosorbent assay (ELISA) for the quantitative analysis of casein traces in wine and food samples. Ovalbumin in the sample binds to an immobilized antibody on the surface of the wells. In a second incubation, another peroxidase-conjugated antibody binds to the ovalbumin previously bound to the well. A final incubation with a peroxidase substrate (TMB) develops color based on the presence of the analyte. The reaction is stopped with sulfuric acid or stop solution. The resulting absorbance change is read at 450 nm and is proportional to the ovalbumin concentration present in the sample.

| Presentation: | 96 wells |
| :--- | :--- |
| Method: | Sandwich ELISA |
| LOD: | 4 ppb |
| Range of measurement: | $0-25-100-250-500 \mathrm{ppb}$ |

Y15 / Y25 / Y350 are Open Analyzers.
In conjunction with the reagent line, the BioSystems Analyzers make it possible to monitor the entire vinification process.
The system adjusts to the various sample types that the enologist needs to analyze.

## (1) 15

TECHNICAL SPECIFICATIONS
Ref. 83106


## (4) 25



Random Access Automatic Analyzer.
Direct photometric reading on the reaction rotor.

| Test rate | 240 tests/hour |
| :---: | :---: |
| Cooled reagent positions | 30 |
| Positions for uncooled racks | 3 (multipurpose rack) |
| Number of samples per rack | 24 |
| Maximum number of samples | 72 |
| Sample tubes | $\emptyset 13 \mathrm{~mm}, \emptyset 15 \mathrm{~mm}$ (max. height 100 mm ) |
| Pediatric vials | $\emptyset 13 \mathrm{~mm}$ |
| Number of reagents per rack | 10 |
| Max. number of uncooled reagents | 20 |
| Reagent bottles | 20 mL and 50 mL |
| Programmable reagent volume | $10 \mu \mathrm{~L}-440 \mu \mathrm{~L}$ |
| Programmable sample volume | $2 \mu \mathrm{~L}-40 \mu \mathrm{~L}$ |
| Removable methacrylate rotor |  |
| Number of wells | 120 |
| Automatic pre- and post-dilutions |  |
| Dilutions using a single calibrator |  |
| Reaction volume range | $180 \mu \mathrm{~L}-800 \mu \mathrm{~L}$ |
| Measurement range | De-0.05 A to 3.6 A |
| Basic filter drum setting | $340,405,420,520,560,600,620,635,670 \mathrm{~nm}$ |
| Dimensions | $1080 \times 695 \times 510 \mathrm{~mm}$ (Lx W x H) |
| Weight | 73 kg |

## (4) 350



BA400 is a high-capacity instrument ( 400 tests/hour) that sets the standard for a new generation of analyzers based on an LED optical system and smart functionality that offers top-notch performance for large enology laboratories.

TECHNICAL SPECIFICATIONS
Ref. 83400
LED TECHNOLOGY


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- Certified Management

System

- EN ISO 9001


[^0]:    Antonio Elduque

