

Microwave Plasma Atomic Emission Spectroscopy (MP-AES)

Measuring metals in wine and food handbook



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Atomic Spectroscopy Solutions

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MP-AES – How it works

How Microwave Plasma – Atomic Emission Spectroscopy works

Atomic spectroscopy describes a number of analytical techniques used to determine the elemental composition of a sample by examining its electromagnetic spectrum, or its mass spectrum. The techniques which identify an analyte element by its electromagnetic spectrum include flame atomic absorption spectroscopy (FAAS), inductively coupled plasma optical emission spectroscopy (ICP-OES) and microwave plasma atomic emission spectroscopy (MP-AES). Those which identify an element by its mass spectrum include inductively coupled plasma mass spectrometry (ICP-MS), and triple quadrupole inductively coupled plasma mass spectrometry (ICP-QQQ).

Atomic excitation

Microwave plasma atomic emission spectroscopy is an atomic emission technique. It uses the fact that once an atom of a specific element is excited, it emits light in a characteristic pattern of wavelengths – an emission spectrum, as it returns to the ground state. Sources for atomic emission include the microwave plasma (MP) and the inductively coupled argon plasma (ICP) both of which are high temperature sources, and therefore excellent excitation sources for atomic emission spectroscopy. The nitrogen-fuelled microwave plasma reaches temperatures nearing 5,000 K. At these temperatures, atomic emission is strong, producing excellent detection limits and linear dynamic range for most elements.

Inside a MP-AES instrument, microwave energy from a robust and reliable industrial magnetron is used to form a plasma from nitrogen that can be extracted from compressed air by Agilent's Nitrogen Generator (Figure 1). Effectively, the MP-AES runs on air.



Figure 1. The nitrogen generator extracts nitrogen from compressed air to fuel the plasma.

Using a magnetic field rather than an electric one for excitation generates a robust plasma – capable of handling a wide range of sample types.

An optimized microwave waveguide creates concentrated electromagetic fields at the torch (Figure 2). An axial magnetic field and a radial electrical field focus and contain the microwave energy to create a plasma.



Figure 2. A microwave waveguide creates concentrated electromagnetic fields around the torch.



Sample introduction

Just like a flame AA instrument, an aerosol is created from a liquid sample using a nebulizer and a spraychamber. The aerosol is then introduced into the centre of the hot plasma.

The aerosol dries, decomposes and is then atomized. The atoms continue to be excited and emit light at wavelengths characteristic for each element as they return to lower energy states (Figure 3).



Figure 4. The emission from the plasm is directed into the optical system.

Optical system

Emission from the plasma is directed into a fast scanning monochromator (Figure 4). The selected wavelength range is imaged onto the high efficiency CCD detector. This measures both spectra and background simultaneously for optimum precision.

Quantification

Just like flame AAS, MP-AES quantifies the concentration of an element in a sample by comparing its emission to that of known concentrations of the element, plotted on a calibration curve. The final result is the concentration of the element in the sample.

The benefits of MP-AES

Low cost of ownership

Gas supply is one of the highest costs associated with elemental analysis. Because the MP-AES runs on air, it vastly reduces the cost of ownership and eliminates the need for ongoing supply of flammable or expensive gases. There's no gas cylinders or lamps to buy and no standby operating costs. When a MP-AES is off, no gas or power is used.

Safety

Removing the requirement for flammable gases means no gas leaks and ordering and transporting of cylinders. Removing all of these risks makes your laboratory a safer workplace.

Better performance than AA

An MP-AES has high sensitivity with detection limits down to sub ppb levels and is faster and has a wider linear dynamic range than conventional flame Atomic Absorption (AA) for a typical multi-element analysis.

Handling difficult matrices

The robust, magnetically excited microwave plasma source of a MP-AES handles difficult matrices with ease, including fuels and organic solvents, geochemical samples, fertilizers and foods. A vertically positioned torch gives the best performance with difficult samples, and features end-on axial viewing for excellent detection limits.

Remote operation

Requiring only electricity, an MP-AES can be located at a sampling point, instead of in a lab. This allows measurement turnaround to be much quicker, delivering timely data that could deliver huge benefits, such as preventing environmental spills or incorrectly manufactured products.

Fast & easy to use

Agilent's MP-AES instruments feature easy-to-use, application-specific software applets that automatically load a pre-set method so you can start analysis immediately without method development or alignment, and with minimal training.

The instrument's torch loader automatically aligns the torch and connects gases for fast start up and reproducible performance.

Why switch from FAAS to MP-AES?

Reducing ongoing operating costs, increasing safety, improving analytical performance and simplifying operation are some of the key challenges facing current users of Flame Atomic Absorption Spectroscopy (FAAS). With the introduction of the Agilent Microwave Plasma-Atomic Emission Spectrometer (MP-AES), these challenges have been overcome, making it the ideal instrument for laboratories looking to transition away from FAAS to a more powerful, less expensive and safer technique. Additionally, with the extra performance of the MP-AES, the sample preparation process can also be significantly simplified, saving time and money.

The Agilent 4210 MP-AES features a waveguide design and torch that is capable of running samples with total dissolved solids up to 3% without compromising detection limits.

Reduced running costs

The largest contributor to ongoing running costs for spectroscopy is gases. FAAS uses a combination of air and acetylene, or nitrous oxide and acetylene for the flame. While air can be provided by an air compressor, the acetylene and nitrous oxide is supplied in cylinders and regularly needs to be replenished as it is consumed.

The 4210 MP-AES uses nitrogen extracted straight from air to sustain the plasma. The Agilent 4107 Nitrogen Generator coupled to an air compressor supplies all the free nitrogen required at greater than 99.5% purity. This leads to dramatic reductions in operating costs over the life of the instrument.

The potential cost saving of using the 4210 MP-AES for the determinaton of Ca, Mg, Na and K in fruit juice is illustrated by comparing a FAAS purchased with an air compressor and 1 year of consumables to an MP-AES purchased with air compressor, nitrogen generator, SPS 4 autosampler, and 1 year of consumables (Figure 1).



Figure 1. <u>Online cost estimator</u> showing the cost savings of using the MP-AES for the determination of Ca, Mg, Na and K in fruit juice.

The analysis requirements were assumed to be 500 samples per week and 4 elements per sample.

The calculation assumes that the FAAS is run without an autosampler and that 3 elements are determined with air/acetylene and 1 element with nitrous oxide/ acetylene. In this example the results show an estimated cost saving of greater than US \$220,000 over a 7 year evaluation period¹. A global average gas cost was used in this calculation and results will vary from country to country.

Improved safety

Another major concern for FAAS users is the safety aspects related to the use of acetylene and nitrous oxide, from the storage and handling of cylinders, to the use of a flame in the instrument. Presence of a naked flame is of concern in all labs, particularly those that handle organic solvents, and for this reason it is not possible to run FAAS unattended.

It is also common to change burners for optimal performance to determine the full range of elements when running FAAS. While Agilent's FAAS instruments are fully interlocked to ensure the correct burners are used with the correct method, care must be taken when handling burners which can remain hot after use.

These issues are eliminated with the 4210 MP-AES. The requirements for acetylene and nitrous oxide can be avoided, along with the storage and handling concerns, and there is no need for burner change-over because of the increased performance of the higher temperature nitrogen plasma.

Improved analytical performance

The plasma of the 4210 MP-AES operates at around 5000 K which results in improved detection limits when compared to FAAS. The improvements in detection limit means that it's possible to analyse elements like phosphorus, which have very high detection limits on FAAS.

Table 1 shows the instrument detection limits (IDL) on MP-AES and FAAS for elements in a rice flour sample. The lower detection limits for P, Cu and Fe allow the major, minor and trace elements to be determined in one sample measurement.

| Element | 4210 Typical IDL 10 sec read μg/L | FAAS Typical IDL μg/L |
|---------|--------------------------------------|--------------------------|
| Са | 0.04 | 0.4 |
| Mg | 0.1 | 0.27 |
| Na | 0.1 | 0.26 |
| К | 0.6 | 0.76 |
| Р | 66 | 26000 |
| Fe | 1.7 | 7.3 |
| Pb | 2.5 | 14 |
| Cu | 0.5 | 1.2 |
| Mn | 0.2 | 1.0 |
| | | |

 Table 1. Comparison of typical instrument detection limits for the 4210 MP-AES and FAAS

1 This example is intended to help you compare the running costs and savings of the MP-AES vs. flame AA. The applied formulas and parameters are correct to the best of our knowledge, but we cannot guarantee the results. Savings may vary depending on factors such as local gas and electricity costs, operator costs, number and types of elements. For this calculation operator labor costs were set to US\$25/hour and electricity costs were set to US\$25.18 per kW. The design of the waveguide and torch in the 4210 MP-AES, combined with the mass flow control and humidification of the nebulizer gas line, gives excellent long term stability in samples with a complex matrix common in mining and environmental samples. The introduction of elevated salt solutions into an air acetylene FAAS burner over an extended period, such as an 8 hr work day, will require maintenance to avoid blockage. If this routine maintenance is not performed, it can lead to signal drift.

Long term stability of the 4210 MP-AES was tested with a 2% TDS solution of digested rice flour. The results are shown in Figure 2.





The 4210 MP-AES also features a greater linear dynamic range than FAAS. Table 2 gives the linear calibration range and correlation coefficient for major elements in a fruit juice sample for the 4210 MP-AES. Also shown is the optimum working range on FAAS for the same elements. The calibrations for FAAS used the default New Rational model. The greater linear range of the MP-AES compared to FAAS reduces the need to perform dilutions on over range samples which simplifies the analysis. Reducing the dilutions also means that if there are trace contaminants that need to be determined, it may still be possible to detect them. Furthermore, the improved linearity means that fewer calibration standards are required for an accurate calibration curve.

Simplifying sample preparation

A factor which greatly influences sample preparation procedures on FAAS is the presence of interferences. Presence of compounds that cannot be broken down in the low temperature flame lead to chemical interference and elements like Na and K can suffer from ionization interference.

Table 2. Linear concentration range of the 4210 MP-AES and optimum concentration range of FAAS for fruit juice samples

| Element | 4210 MP-AES linear concentration range (mg/L) | Linear correlation coefficient on MP-AES calibration | FAAS optimum working range (mg/L) |
|------------|---|---|--------------------------------------|
| Ca 422.673 | 0 – 20 | 0.9999 | 0.01 - 10 |
| Mg 518.360 | 0 – 100 | 0.99988 | 0.15 - 20 (for Mg 202.6) |
| Na 589.592 | 0 - 20 | 0.99996 | 0.01 – 2.0 |
| K 769.897 | 0 – 100 | 0.99968 | 1 - 6.0 |

Table 3. Typical sample preparation requirements for FAAS and MP-AES

| Element | Possible chemical interferences | FAAS specific sample preparation | MP-AES specific sample preparation |
|---------|--|---|------------------------------------|
| Са | Refractory compounds Ionization effects | Lanthanum releasing agent Cesium ionization buffer | None |
| Mg | Ionization effects | Cesium ionization buffer | None |
| Na | Refractory compounds Ionization effects | Lanthanum releasing agent Cesium ionization buffer | None |
| К | Ionization effects | Cesium ionization buffer | None |

Table 4. Certified Reference Material (CRM) recoveries of major elements in grapefruit juice, analyzed by MP-AES. No ionization suppressant was required and excellent accuracy for K was achieved. Additionally, no lanthanum nitrate was added and excellent recoveries for Ca was achieved

| Grapefruit Juice T0842QC | Certified Value (mg/L) | | Found (mg/L) | %Recovery |
|-----------------------------|------------------------|---------------|--------------|-----------|
| | Assigned Value | Range | | |
| Calcium | 145.6 | 123.6 - 167.6 | 158.3 ± 3.2 | 108.7 |
| Magnesium | 92.5 | 77.5 - 107.4 | 91.1 ± 0.6 | 98.5 |
| Potassium | 1102 | 979 – 1225 | 1100 ± 14.7 | 99.8 |

Various strategies for dealing with these interferences are well established. It is common to add releasing agents such as strontium or lanthanum to overcome chemical interferences, or alternatively the hotter nitrous oxide flame can be used. Ionization effects are usually overcome by adding an ionization buffer to the solution, such as sodium, potassium or cesium. Another strategy is to extract the elements of interest into an organic phase in order to remove the interfering elements. As a result the sample must be individually prepared for each element in the sample.

With the hotter plasma source of the 4210 MP-AES these chemical interferences are eliminated. This means that the element specific sample preparation required on FAAS is not needed which greatly simplifies the sample preparation process. As an example the elements covered in a fruit juice analysis are shown above with a comparison of the sample preparation required for each element (Tables 3 and 4).

Conclusion

Agilent's 4210 MP-AES is the ideal instrument for customers looking to transition from Flame Atomic Absorption Spectroscopy (FAAS) to another technique. By using nitrogen as the source gas for the plasma, running costs are greatly reduced, and by removing the requirement for hazardous nitrous oxide and acetylene safety is greatly increased. Additionally the higher temperature nitrogen plasma atomization/ionization source improves detection limits, linear range, and long term stability, and allows the sample preparation process to be greatly simplified.

MP-AES Accessories

Expanding capabilities with accessories

Agilent offers a range of accessories for the MP-AES instruments. They add additional capabilities to the instrument, allowing you to optimise the setup for specific applications.



Advanced Valve System (AVS 4)

This four port switching valve enables high sample throughput and prolongs component lifetime due to the reduced exposure of sample introduction components to harsh samples.

Multimode Sample

Introduction System (MSIS) An innovative accessory for both ICP-OES and MP-AES that allows sample introduction via either vapor generation or nebulization modes, or both modes at the same time. Allows sub ppb detection of As, Se, and Hg.





SPS 4 Autosampler Our fastest ever autosampler, the SPS 4 allows the unattended multi-element analysis of up to 360 samples.



4107 Nitrogen Generator The nitrogen generator extracts nitrogen from compressed air, to fuel the plasma of an MP-AES instrument. It provides a continuous supply of nitrogen.



IsoMist A temperature controlled spraychamber that offers improved stability for volatile organic solvents.



External Gas Control Module (EGCM) The EGCM injects air into the plasma to minimize carbon build up, reduce background in organic applications and enable sulfur determinations.

Wine

MP-AES offers the simple, fast, automated analysis of the elements in wine – with no sample preparation required. The International Organisation of Vine and Wine (OIV) has approved the use of <u>MP-AES</u> for the analysis of iron, copper, potassium, calcium and manganese in wine samples. These elements are monitored regularly to control the growth of the vineyard, the grape quality, the aromatic loss, and to determine the risk of oxidation. MP-AES offers the advantages of:

- No sample preparation is required as on-line dilution of the wine sample is used so it's a straightforward and simple analysis
- MP-AES can be used to measure: copper 0.05-1 mg/L, iron 1-10 mg/L, potassium 15-1200 mg/L, calcium 1-100 mg/L, and manganese 0.25-4 mg/L
- 100+ samples per day can be easily measured
- No gases need be purchased and only nitrogen is required for the analysis.
 This makes it a safe technique for use in oenological labs

An MP-AES can also be used to measure plant and soil samples, being able to determine:

- The nutritional balance of the vines during the growing season
- Vine elemental nutrient needs for the coming season, so foliar applications can be planned
- The agro-oenological potential of soils being considered for planting



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Routine analysis of total arsenic in California wines using the Agilent 4200/4210 MP-AES

Application note Food Safety



Introduction

Arsenic (As) is a naturally occurring element found throughout the world. The environmental levels of As have been increasing due to natural sources, such as volcanic activity, and anthropogenic sources, such as smelting. The continuous release of As into the ecosystem has formed an accumulation of the element in the food chain.

Wine is a globally consumed beverage where total levels of As are regulated between 100-200 μ g L⁻¹, depending on the country in question [1]. However, there are countries, such as the United States, that do not regulate levels

of all elements in wine. This necessitates investigating total As levels in wine produced in the United States to identify potential contamination, beyond the levels regulated by other countries.

Measuring total arsenic levels in wine with various spectrometric techniques typically deliver insufficient sensitivity due to the element's relatively high ionization potential. However, the use of vapor generation techniques to form volatile forms of As allow for a more sensitive detection of As.

This study investigates the use of the Agilent 4200 Microwave Plasma-Atomic Emission Spectrometer (MP-AES) coupled with the Multimode Sample Introduction System (MSIS) accessory to measure total As in wine samples from the California region. This application is also applicable for Agilent's 4210 MP-AES instrument.

The MP-AES offers high sensitivity, with superior performance in comparison to Flame Atomic Absorption Spectroscopy. The instrument uses nitrogen to sustain the plasma, either extracted from the ambient air (using a nitrogen generator) or supplied via a nitrogen Dewar. Compared with acetylene-based instruments, the MP-AES is much safer to run, as no flammable gases are required. Operating costs are also significantly less. The addition of the MSIS accessory assists with the production of hydride species which are separated from the liquid and introduced to the plasma, delivering better performance and lower detection limits, than with conventional nebulization.

Experimental

Instrumentation

All measurements were performed using the Agilent 4200 MP-AES fitted with the MSIS accessory, MicroMist glass nebulizer and Easy-fit torch.

The As 188.979 nm line was selected for analysis and the read time optimized in the easy to use MP Expert software that controls the instrument. The instrument operating and method settings are given in Table 1. The Agilent MP Expert software allows for correction of background and spectral interferences. The Fast Linear Interference Correction (FLIC) background correction (refer to Figure 1) was used to correct for the complex structured background by modeling with the solution blank of 10% HCI and 5% ethanol as the interferent.



Figure 1. FLIC background correction applied to As (188.979 nm) calibration standard. Background and analyte peak (blue line), FLIC model applied to background and analyte peak (green line).

Table 1. Agilent 4200 MP-AES operating and method conditions.

| Parameter | Setting |
|-------------------------|---------------------------------|
| As wavelength (nm) | 188.979 |
| Pump speed (rpm) | 20 |
| Sample pump tubing | Black-black |
| Hydride reagent tubing | Black-black |
| MSIS waste tubing | Black-white |
| Read time (sec) | 20 |
| Number of Replicates | 3 |
| Sample uptake delay (s) | 40 |
| Nebulizer flow (L/min) | 0.45 |
| Fast pump during uptake | On |
| Stabilization time (s) | 20 |
| Background correction | FLIC |
| Calibration Fit | Linear |
| Nitrogen gas source | Agilent 4107 Nitrogen Generator |

Samples

Forty commercially available wines, originating from various areas in California, were analysed in this study (Table 2). The wine varieties included: Pinot noir, Merlot, Cabernet Sauvignon, Rosé, Chardonnay, white Zinfandel, Sauvignon blanc, a white blend, sparkling wine and portstyle wine.
 Table 2. Wine samples used in the study, with corresponding sample number and regional origin.

| Sample# | Wine type | Region |
|---------|--------------------------------|-------------------------------------|
| 1 | Cabernet Sauvignon | South Coast |
| 2 | Cabernet Sauvignon | Sonoma County |
| 3 | Cabernet Sauvignon | Lodi/Woodbridge Grape Commission |
| 4 | Cabernet Sauvignon | South Coast |
| 5 | Cabernet Sauvignon | Sierra Foothills |
| 6 | Cabernet Sauvignon | Napa County |
| 7 | Cabernet Sauvignon | Sierra Foothills |
| 8 | Cabernet Sauvignon | Napa County |
| 9 | Cabernet Sauvignon | Greater Bay Area |
| 10 | Cabernet Sauvignon | North Coast |
| 11 | Cabernet Sauvignon | North Coast |
| 12 | Cabernet Sauvignon | South Central Coast |
| 13 | Cabernet Sauvignon | Greater Bay Area |
| 14 | Cabernet Sauvignon | Sonoma County |
| 15 | Cabernet Sauvignon | Lodi/Woodbridge Grape Commission |
| 16 | Cabernet Sauvignon | North Central Coast |
| 17 | Cabernet Sauvignon | South Central Coast |
| 18 | Cabernet Sauvignon | South Coast |
| 19 | Cabernet Sauvignon | South Central Coast |
| 20 | Cabernet Sauvignon | Sonoma County |
| 21 | Cabernet Sauvignon | Sierra Foothills |
| 22 | Cabernet Sauvignon | South Central Coast |
| 23 | Cabernet Sauvignon | North Central Coast |
| 24 | Cabernet Sauvignon | North Coast |
| 25 | Cabernet Sauvignon | Greater Bay Area |
| 26 | Cabernet Sauvignon | Napa County |
| 27 | Cabernet Sauvignon | Sierra Foothills |
| 28 | Pinot Noir | Appellation Central Coast |
| 29 | White Blend | San Joaquin County |
| 30 | Rosé | Contra Costa County |
| 31 | Rosé | Lodi/San Joaquin County |
| 32 | Chardonnay | Santa Barbara County |
| 33 | White Zinfandel | Napa and Sonoma |
| 34 | Chardonnay | Central Coast |
| 35 | Chardonnnay | Napa County |
| 36 | Merlot | Napa County |
| 37 | White Zinfandel | Napa and Lodi |
| 38 | Sauvignon Blanc | Oakville/Napa County |
| 39 | California Brut sparkling wine | Sonoma County |
| 40 | Petite Sirah port-style wine | Clarksburg/Yolo County |

Sample preparation

All wine samples were diluted by a factor of three with Millipore ultrapure water (Milli-Q[™] Water System, Darmstadt, Germany) then further diluted with hydrochloric acid (34-37.5%, Environmental Grade, Alfa Aesar, Ward Hill, MA, USA) for a total acid concentration of 10%. This resulted in a final 3.3-fold wine dilution. Each wine was prepared in triplicate with the exception of sample 37, which was prepared with 5 replicates due to high residual sugar content.

Four wine samples were selected and prepared as spiked samples for additional analysis. Low and high concentration spikes were used at 10 μ gL⁻¹ and 50 μ gL⁻¹ and prepared in triplicate.

Calibration standards and reagents

A 1,000 μ gL⁻¹ single element calibration standard of As (VHG Labs, Manchester, NH, USA) was used to prepare working standards at 5, 10, 25, 50 and 100 μ gL⁻¹. As (III) and As (V) calibration standards from SPEX CertiPrep (Metuchen, NJ, USA) were used as check standards to validate the method at 20 μ gL⁻¹ each. All calibration standards were matrix-matched with a 10% hydrochloric acid (34-37.5% Environmental Grade, Alfa Aesar, Ward Hill, MA, USA) and 5% ethanol (v/v) (200 proof, Gold Shield Distributors, Hayward, CA) solution.

Two separate solutions were created for hydride generation. The first consisted of 1.2% Sodium Borohydride (NaBH₄, 98%, J.T. Baker, Center Valley, PA, USA) and 1.0% Sodium Hydroxide (NaOH, ACS Grade, EMD Chemicals Inc., Gibbstown, NJ) in Millipore ultrapure water (Milli-Q[™] Water System, Darmstadt, Germany). The second was a 1:1 solution of HCI (34-37.5%, Environmental Grade, Alfa Aesar, Ward Hill, MA, USA) and Millipore ultrapure water (Milli-Q[™] Water System, Darmstadt, Germany).

A reduction solution of 25% (w/v) potassium iodide (ACS Grade, BDH Chemicals, West Chester, PA) was used to reduce As species prior to analysis. As (III) and As (V) are the most prevalent forms of As in wines. This reduction step aims to change the valence state of As from As (V) to As(III), because As (V) does not readily form a metal hydride. The potassium iodide solution was added to all samples and standards to create a final concentration of 1%. Best results were obtained when it was added at least 3 hours prior to analysis. The setup of the MSIS (in Vapor Generation mode) for this analysis is displayed in Figure 2. The sample and 50% HCl solution were mixed using a 'tee' fitting after the peristaltic pump and the combined sample/ HCl line was connected to the bottom of the MSIS. The reductant, NaOH/NaBH₄ solution, line was attached to the top of the MSIS. The unused sample line to the nebulizer was blocked during analysis as conventional nebulization was not required.



Figure 2. The MSIS setup for Vapor Generation mode.

Results and discussion Calibration

The five-point calibration curve for As (188.979 nm) is shown in Figure 3. It shows excellent linearity with a calibration coefficient of greater than 0.999 and less than 6% error on each calibration point.



Figure 3. Calibration curve for As 188.979 nm showing excellent linearity across the 5-100 $\mu g L^{\rm 1}$ concentration range, illustrating the wide analytical range of the instrument.

Method Detection Limit

The Method Detection Limit (MDL) for As was determined from the analysis of ten replicate measurements of the blank solution. The results displayed in Table 3, show that the calculated MDL (confidence interval of 99.5%) for As was 0.34 µgL⁻¹.

| Table 3. The calculated MDL and standard deviation results for | |
|--|--|
| As 188.979 nm. | |

| Element | Concentration (μ gL ⁻¹) |
|-------------|--|
| Mean (n=10) | 0.01 |
| SD | 0.10 |
| MDL (0.995) | 0.34 |
| MDL (0.99) | 0.29 |
| MDL (0.95) | 0.19 |

Method validation

To check the validity of the method, Quality Control (QC) samples were run after the calibration and throughout the analytical run. The Continuing Calibration Verification (CCV) and Continuing Calibration Blanks (CCB) were measured every 10 samples. The initial calibration verification block included a 20 μ gL⁻¹ As (III) and As (V) sample. This was done to validate the calibration and confirm the efficiency of the KI reduction step of As (V) to As (III) prior to analysis. All CCV recoveries were within ± 10% of the assigned values (Table 4).

Table 4. QC recoveries of CCB, CCV and 20 $\mu g L^{\cdot 1} As$ (V) and As (III) samples.

| Solution | Concentration (µgL ^{.1}) | Recovery (%) |
|---|------------------------------------|--------------|
| CCB, (mean, n=7) | 0.46 | - |
| 25 μ g L ⁻¹ CCV, (mean, n=7) | 23.94 | 96 |
| 20 µg L ^{.1} As (V) | 20.59 | 103 |
| 20 µg L ^{.1} As (III) | 19.92 | 99.6 |

Analysis of wine samples

The method described above was applied to the analysis of 40 Californian wine samples. The results obtained for each sample are the average of 3 replicates and can be found in Table 5, along with the standard deviation and relative standard deviation (%RSD). As concentrations in the wine samples ranged from below the MDL to 48.81 μ gL⁻¹, well below the range of 100-200 μ gL⁻¹ regulated in many countries.

 Table 5. The quantitative results for total As (188.979 nm) concentration in 40

 Californian wines using the 4200 MP-AES.

| Sample | mple Mean Sta Concentration (μgL ⁻¹) De (mean, n=3) | | Relative Standard Deviation (%) |
|--------|---|------|---------------------------------------|
| 1 | 1.03 | 0.88 | 0.85 |
| 2 | <mdl< td=""><td>0.71</td><td>N/A</td></mdl<> | 0.71 | N/A |
| 3 | 1.48 | 2.12 | 1.43 |
| 4 | 1.43 | 0.84 | 0.59 |
| 5 | <mdl< td=""><td>0.21</td><td>N/A</td></mdl<> | 0.21 | N/A |
| 6 | <mdl< td=""><td>0.40</td><td>N/A</td></mdl<> | 0.40 | N/A |
| 7 | <mdl< td=""><td>0.77</td><td>N/A</td></mdl<> | 0.77 | N/A |
| 8 | <mdl< td=""><td>0.74</td><td>N/A</td></mdl<> | 0.74 | N/A |
| 9 | <mdl< td=""><td>0.74</td><td>N/A</td></mdl<> | 0.74 | N/A |
| 10 | 43.81 | 1.13 | 0.03 |
| 11 | 2.92 | 3.55 | 1.22 |
| 12 | <mdl< td=""><td>1.17</td><td>3.95</td></mdl<> | 1.17 | 3.95 |
| 13 | <mdl< td=""><td>3.91</td><td>1.90</td></mdl<> | 3.91 | 1.90 |
| 14 | 6.63 | 1.14 | 0.17 |
| 15 | 6.09 | 1.45 | 0.24 |
| 16 | 3.24 | 1.89 | 0.58 |
| 17 | <mdl< td=""><td>0.22</td><td>N/A</td></mdl<> | 0.22 | N/A |
| 18 | 2.33 | 1.33 | 0.57 |
| 19 | <mdl< td=""><td>2.11</td><td>14.06</td></mdl<> | 2.11 | 14.06 |
| 20 | <mdl< td=""><td>1.52</td><td>6.70</td></mdl<> | 1.52 | 6.70 |
| 21 | 0.75 | 0.92 | 1.23 |
| 22 | <mdl< td=""><td>2.51</td><td>N/A</td></mdl<> | 2.51 | N/A |
| 23 | 10.16 | 1.15 | 0.11 |
| 24 | <mdl< td=""><td>1.63</td><td>N/A</td></mdl<> | 1.63 | N/A |
| 25 | <mdl< td=""><td>1.74</td><td>N/A</td></mdl<> | 1.74 | N/A |
| 26 | <mdl< td=""><td>1.66</td><td>N/A</td></mdl<> | 1.66 | N/A |
| 27 | <mdl< td=""><td>0.22</td><td>N/A</td></mdl<> | 0.22 | N/A |
| 28 | 3.96 | 0.52 | 0.06 |
| 29 | 1.37 | 1.16 | 0.85 |
| 30 | <mdl< td=""><td>0.71</td><td>N/A</td></mdl<> | 0.71 | N/A |
| 31 | <mdl< td=""><td>0.47</td><td>N/A</td></mdl<> | 0.47 | N/A |
| 32 | 27.04 | 1.21 | 0.04 |
| 33 | 31.17 | 4.72 | 0.15 |
| 34 | 4.53 | 0.37 | 0.08 |
| 35 | 9.86 | 0.23 | 0.02 |
| 36 | 3.44 | 0.46 | 0.13 |
| 37 | 17.84 (mean, n=5) | 0.63 | 0.04 |
| 38 | 11.23 | 0.28 | 0.02 |
| 39 | 9.76 | 0.24 | 0.02 |
| 40 | 2.64 | 0.90 | 0.34 |

Analysis of spiked wine samples

Four wines, representing various total As concentrations, were selected for a spike recovery study. Two spikes were completed for each sample at 10 and 50 μ gL⁻¹. This equates to spikes of 33 and 165 μ gL⁻¹ in the sample. The recoveries for each spike were within \pm 10%, with the exception of one which had a recovery of 111%. Results seen in Table 6, show excellent recovery for As using the MP-AES at low and high μ gL⁻¹ levels.

Conclusions

The Agilent 4200 MP-AES coupled with the MSIS accessory provided an easy and accurate analysis of total As in wine. The MSIS technology increased sensitivity to levels lower than single digit μ gL⁻¹. The resulting MDL would be approximately 100 times lower when compared to using direct nebulization. All 40 wine samples analyzed were found to have As concentrations less than the levels regulated by most countries worldwide.

The complex background on the calibration and samples was easily corrected with Agilent's Fast Linear Interference Correction (FLIC), improving analytical accuracy.

The nitrogen-based plasma significantly reduces operating costs when nitrogen is supplied with the use of a Agilent 4107 Nitrogen Generator. The generator extracts nitrogen from compressed air. Alternatively, nitrogen can be supplied by Dewar.

The Agilent 4200 MP-AES with MSIS accessory proved to be a reliable, cost-effective instrument for quantifying total As in wine.

<MDL = below Method Detection Limit

| Sample# | Spiked concentration (µgL ⁻¹) | | Measured spiked concentration (µgL-1) | Measured unspiked concentration (µgL-1) | Recovery (%) |
|---------|--|------|---------------------------------------|---|-----------------|
| 8 | 33 | Ave | 32.25 | -1.53 | 102.36 |
| | | %RSD | 0.03 | -0.48 | |
| 8 | 165 | Ave | 153.26 | -1.53 | 93.82 |
| | | %RSD | 0.01 | -0.48 | |
| 23 | 33 | Ave | 42.76 | 10.16 | 98.81 |
| | | %RSD | 0.01 | 0.11 | |
| 23 | 165 | Ave | 164.79 | 10.16 | 93.72 |
| | | %RSD | 0.01 | 0.11 | |
| 28 | 33 | Ave | 39.54 | 3.96 | 107.82 |
| | | %RSD | 0.01 | 0.06 | |
| 28 | 165 | Ave | 161.48 | 3.96 | 95.46 |
| | | %RSD | 0.04 | 0.06 | |
| 32 | 33 | Ave | 57.63 | 27.04 | 92.68 |
| | | %RSD | 0.00 | 0.04 | |
| 32 | 165 | Ave | 211.12 | 27.04 | 111.56 |
| | | %RSD | 0.01 | 0.04 | |

Table 6. The percent recovery results for the four As spiked wine samples at 10 and 50 µgL⁻¹, the results shown represent the concentration in the original sample, calculated accounting for the dilution factor.

Reference

C. K. Tanabe, H. Hopfer, G. Gilleland, A. Liba, S. E. Ebeler and J. Nelson. Total arsenic analysis in Californian wines with hydride generation – microwave plasma – atomic emission spectroscopy (HG-MP-AES). *J. Anal. At. Spectrom*, 2016



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Elemental profiling of Malbec Wines for geographical origin using an Agilent 4200 MP-AES

Application note

Food testing



Introduction

Red wine produced from the Malbec grape is increasing in popularity in the United States. However, the US is a relatively small producer of the wine compared to Argentina, where it is the most extensively planted grape variety in the country. With rising imports into the US from Argentina there are growing concerns relating to the validation of the geographical origin of this wine.

Typically ICP-MS is used to distinguish between wines originating from different regions by comparing the relative concentrations of mineral elements, which are characteristic of the soil composition of the region of production. In this study, a cost-effective approach has been taken using Microwave Plasma-Atomic Emission Spectroscopy (MP-AES) to measure 6 elements (Sr, Rb, Ca, K, Na and Mg). Agilent's Mass Profiler Professional (MPP) integrated chemometrics software and another data analysis package were used to model the MP-AES results to distinguish the geographical origin of 41 Malbec wine samples produced in Argentina and the USA.



Experimental

Samples

Malbec grapes from the 2011 vintage were sourced from 41 different geographical sites; 26 from the Mendoza region of Argentina and 15 from California, USA. In order to reduce the impact of the wine making process on the elemental composition of each wine, and to preserve any elemental differences arising from the geographical origin, two central winemaking facilities only were used to produce the wine. Table 1 lists all the samples, with their detailed geographical origin. Details of the winemaking procedure can be found in the original study [1].

Calibration standards and reagents

Single-element calibration standards (Ca, K, Mg, Na at 10,000 mg/L, and Sr at 1,000 mg/L) were purchased from VHG Labs (Manchester, NH, USA), Rb 1,000 mg/L was from SPEX CertiPrep (Metuchen, NJ, USA), and concentrated nitric acid was obtained from JT Baker (Instra-Analyzed grade, Center Valley, PA, USA). The ionization buffer solution (100,000 mg/L Cs; Agilent, Santa Clara, CA, USA) was diluted to 2,000 mg/L in 1% HNO₃ prior to use. Ultrapure water (18 MΩcm, EMD Millipore Bellerica, MA, USA) and Uvasol spectroscopy grade ethanol from Merck (Whitehouse Station, NJ, USA) were used for the calibration solutions and dilutions.

Instrumentation

An Agilent 4200 MP-AES fitted with a MicroMist concentric nebulizer and baffled cyclonic spray chamber was used throughput the study. An External Gas Control Module (EGCM) was used to inject air into the nitrogen plasma to prevent carbon present in the wine samples from building up on the torch. This ensures stable results over the course of the analysis and reduces the background emissions generated by the organic species present in the sample. A 2,000 mg/L cesium (Cs) Ionization Buffer solution was constantly mixed with the sample stream immediately before entering the spray chamber, using a simple mixing tee.

Each element (Sr, Rb, Ca, K, Na and Mg) was monitored at a specific wavelength to ensure interference-free detection. EGCM and read time settings were optimized for each element. The instrument was calibrated and tuned daily using an Agilent wave calibration solution. All wine samples were analyzed in triplicate after a 1:50 dilution in 5% HNO_3 . A 6-point calibration between 0 and 500 mg/L was carried out for each element in matrix-matched calibration solutions (5% HNO_3 and 0.2% ethanol) to account for matrix interferences of the ethanolic wine solutions.

The sample introduction and calibration parameters used are given in Table 2 and 3 respectively.

Statistical data analysis

Data analysis of the concentrations of the 6 monitored elements monitored in each of the Malbec wines was carried out in RStudio (version 0.98.501, Boston, MA) and Agilent's Mass Profiler Professional (MPP; version 12.61). Multivariate analysis of variance (MANOVA) and individual univariate analysis of variance (ANOVA) for each element were run in RStudio. Elements that differed significantly among the wines were further used in an untargeted Principal Component Analysis (PCA) within the MPP software to visualize the sample differences. As a final analysis, Partial Least Squares – Discriminate Analysis (PLS-DA) was used for the geographical classification of the wines, according to country and to region within a country.

 Table 1. Samples included in the study. For each wine sample, the district, department, and altitude is shown. *Denotes samples from the Yolo region - the only US region outside a recognized American Viticultural Area (AVA). Source: Nelson et al [1].

| Sample code | District | Department / AVA or County | Altitude (meters above sea level) | Sample code | District | Department /AVA or County | Altitude (meters above sea level) |
|----------------|----------------|-------------------------------|---|-------------|--------------------|------------------------------|---|
| M1 | La Consulta | San Carlos | 999 | M22 | El Peral | Tupungato | 1235 |
| M2 | Perdriel | Luján | 964 | M23 | El Peral | Tupungato | 1235 |
| M3 | La Consulta | San Carlos | 999 | M24 | El Peral | Tupungato | 1241 |
| M4 | La Consulta | San Carlos | 999 | M25 | Gualtallary | Tupungato | 1354 |
| M5 | La Consulta | San Carlos | 999 | M26 | Gualtallary | Tupungato | 1353 |
| M6 | Las Compuertas | Luján | 1022 | C1 | Yountville | Napa | Not available |
| M7 | Las Compuertas | Luján | 1022 | C2 | Mount Veeder | Napa | 315 |
| M8 | Las Compuertas | Luján | 1022 | C3 | Mount Veeder | Napa | 510 |
| M9 | Altamira | San Carlos | 1024 | C4 | Mount Veeder | Napa | 497 |
| M10 | Altamira | San Carlos | 1043 | C5 | Oak Knoll District | Napa | 25 |
| M11 | Altamira | San Carlos | 1096 | C6 | Alexander Valley | Sonoma | 58 |
| M12 | Altamira | San Carlos | 1047 | C7 | Alexander Valley | Sonoma | 68 |
| M13 | Altamira | San Carlos | 1043 | C8 | Alexander Valley | Sonoma | 53 |
| M14 | Altamira | San Carlos | 1024 | C9 | Hames Valley | Monterey | 214 |
| M15 | Gualtallary | Tupungato | 1342 | C10 | Monterey County | Monterey | 154 |
| M16 | Altamira | San Carlos | 1052 | C11 | Lodi | San Joaquin | 61 |
| M17 | El Peral | Tupungato | 1235 | C12 | Winters* | Yolo | 88 |
| M18 | Lunlunta | Maipú | 931 | C13 | Winters* | Yolo | 77 |
| M19 | Lunlunta | Maipu | 930 | C14 | Winters* | Yolo | 70 |
| M20 | El Peral | Tupungato | 1235 | C17 | Red Hills | Lake | 648 |
| M21 | El Peral | Tupungato | 1235 | | | | |

 Table 2. 4200 MP-AES operating conditions. Source: Nelson et al [1]

| Parameter | | | Va | lue | | |
|---------------------------------|-----------|---------|---------|---------|---------|---------|
| Element | Sr | Rb | Mg | Ca | Na | K |
| Monitored wavelength (nm) | 407.771 | 780.027 | 279.553 | 396.847 | 589.592 | 769.897 |
| EGCM set- ting | Low | Low | Med | | High | |
| Pump rate (rpm) | | | 1 | 0 | | |
| Sample tubing | Org-Grn | | | | | |
| lonization buffer tubing | Org-Grn | | | | | |
| Waste tubing | Blue-Blue | | | | | |
| Read time (s) | 5 2 | | | | | |
| Number of replicates | 3 | | | | | |
| Sample uptake delay (s) | 50 | | | | | |
| Stabilization delay (s) | 20 | | | | | |
| Fast pump during uptake | Yes | | | | | |
| Background correction | | | Aı | uto | | |

 Table 3. Calibration parameters used for wine sample analysis.

 Source: Nelson et al [1]

| Element | λ (nm) | Calibration Range (mg/L) | Background Correction | Calibration fit | Correlation Coefficient |
|---------|---------|--------------------------------|--------------------------|--------------------|----------------------------|
| Sr | 407.771 | 0-5 | auto | linear | 0.9999 |
| Rb | 780.027 | 0-5 | auto | linear | 0.9997 |
| Mg | 279.553 | 0-5 | auto | linear | 0.9998 |
| Са | 396.847 | 0-5 | auto | linear | 0.9999 |
| Na | 589.592 | 0-5 | auto | linear | 0.9999 |
| К | 769.897 | 0-20 | auto | linear | 0.99999 |

Results and discussion

Three sigma detection limits were determined by analyzing 10 sample blanks. All 6 of the elements monitored were detected in the 41 different wine samples at concentrations above their limits of detection (LODs), as shown in Table 4. All elements also differed significantly among the wine samples in a multi- and univariate analysis of variance at an α level of 5%. Thus, all 6 elements were included in the subsequent PCA and PLS-DA analyses.

Figure 1a shows a clear separation of the wines by country of origin, with only a slight overlap of two US wines. The component loadings plot (Figure 1a) shows that the elemental differences in Na and Sr primarily account for the separation by country of origin.



Figure 1. 2D PCA bi-plots using the 6 elements which differed significantly among the wine samples. (a) Product plot showing the wine samples color coded by geographical origin, (b) Loadings plot with 6 elements (Sr, Rb, Ca, K, Ca, Na and Mg). *Source: Nelson et al.* [1].

However, an almost 100% correct classification of the wines according to their country of origin was obtained using PLS-DA (Figure 2). Using cross-validation, the prediction accuracy for the USA wines was 93.3% and 96.2% for the Argentina wines, leading to an overall accuracy of 95.1% for the PLS-DA model (Table 5). The incorrectly classified wines (M1 for the Argentina wines and C12 for the USA wines), were most likely due to their higher/lower levels in Na, Mg, and K (M1 was low in Na, Mg, and K; C12 was high in Mg) compared to the other wines in the same class. Excellent classification was achieved. If a larger set of samples had been available, we would have tested to see how the model performs with complete unknown samples.



Figure 2. 2D PLS plots using the 6 elements which differed significantly among the wine samples. (a) Sample plot showing the separation of the wines according to geographical origin, with no overlap. (b) Element loadings plot. *Source: Nelson et al [1]*

Table 4. Detection limits (DL) and elemental concentrations for the wines from Argentina and the USA. Shown are mean \bar{x} , standard error of the mean $\sigma_{\bar{x}'}$, and the minimal (min) and maximal (max) concentrations. Concentrations (mg/L) are given for the elements that differed significantly among the five wineries (P \leq 0.05). Source: Nelson et al [1]

| | | Wines from Ca | alifornia, USA | Wines from Men | doza, Argentina |
|----|--------|--------------------------------|-----------------|--------------------------------|-----------------|
| | DL | $\bar{x} \pm \sigma_{\bar{x}}$ | min – max | $\bar{x} \pm \sigma_{\bar{x}}$ | min – max |
| | (mg/L) | (mg/L) | (mg/L) | (mg/L) | (mg/L) |
| Sr | 0.0018 | 0.45 ± 0.02 | 0.24 - 0.83 | 0.77 ± 0.04 | 0.23 - 1.59 |
| Rb | 0.0004 | 3.37 ± 0.03 | 0.57 - 7.83 | 0.99 ± 0.02 | 0.55 - 2.19 |
| Mg | 0.0012 | 80.87 ± 0.42 | 61.75 - 144.86 | 72.87 ± 0.54 | 53.55 - 116.29 |
| Ca | 0.0016 | 51.78 ± 0.19 | 43.26 - 74.01 | 50.46 ± 0.25 | 33.22 - 95.08 |
| Na | 0.0007 | 4.90 ± 0.09 | 3.38 - 8.46 | 37.48 ± 0.48 | 13.71 - 121.87 |
| К | 0.0020 | 1444.42 ± 10.53 | 1120.94-2219.84 | 1181.22 ± 15.45 | 976.85-1989.12 |

 Table 5. Result of the cross validation (leave one out algorithm) for the PLS-DA run on the 2 countries. *Source: Nelson et al* [1]

| | USA (predicted) | Argentina (predicted) | Accuracy |
|------------------|-----------------|--------------------------|----------|
| Argentina (true) | 1 (= M1) | 25 | 96.2% |
| USA (true) | 14 | 1 (= C12) | 93.3 % |
| Overall accuracy | | | 95.1 % |

Conclusions

The Agilent 4200 MP-AES is an easy-to-use, low cost instrument suitable for geographical origin analysis of wine samples when combined with a data analysis package such as Agilent's Mass Profiler Professional (MPP). Six elements, Sr, Rb, Mg, Ca, Na, and K, were useful for broad classification of geographic origin of Malbec wines from Argentina and the US, with 14 out of 15 US samples correctly classified and only 1 out of 26 of the Argentinian wines wrongly classified.

Reference

1. Jenny Nelson, Helene Hopfer, Greg Gilleland, Daniel Cuthbertson, Roger Boulton, Susan E Ebeler. Elemental Profiling of Malbec Wines Made Under Controlled Conditions by Microwave Plasma Atomic Emission Spectroscopy. Am. J. Enol. Vitic. Published ahead of print April 2015



Determination of metals in wine using the Agilent **4100 Microwave Plasma-Atomic Emission Spectrometer**

Application note

Food Testing and Agriculture

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Introduction

The concentrations of certain metals in wine are of great interest because of their influence on the wine-making process. Strict analytical control of the trace element content is required during the entire wine making process. For example, metals such as potassium, calcium, and iron can produce precipitates, cause cloudiness, or affect the taste.

The wine maker needs to properly control the production process so that the quality of the product can be assured. During vintage, when monitoring trace elements is most critical, sample turnaround time (and to a lesser extent sample throughput) becomes important. Most wine labs are small to medium in size, and hence value ease of use and reduced infrastructure requirements.

Metals in wine can be determined by a number of analytical techniques¹⁻¹⁰. The most common technique used is Flame Atomic Absorption (FAA), while ICP-OES is sometimes used in larger central laboratories where extra sample throughput is required, although having elemental analysis capabilities close to the winery during vintage is generally preferred.

This work describes an alternative, safer and cheaper analytical method for the determination of metals in wine using the Agilent 4100 Microwave Plasma-Atomic Emission Spectrometer (MP-AES).

Which measurement technique is right for you?

There are many factors to be taken into account when selecting the right analytical technique. In many cases several techniques will provide adequate detection range, so the technique of choice will depend on factors such as sample throughput requirements, ease-of-use, infrastructure required, and on-going operating costs.

The MP-AES offers significantly reduced on-going operating costs over both FAA and ICP-OES by running on nitrogen that can be supplied via a nitrogen generator. This eliminates the need for on-going gas resupply and avoids flammable gases (required for FAA), enhancing safety and allowing unattended, overnight operation. The reduced infrastructure required for MP-AES also makes it well suited to remote sites where supply of expensive specialty gases can be difficult.

The 4100 MP-AES fits between FAA and ICP-OES in many aspects such as detection power, dynamic range, and speed of analysis. For these key performance metrics, the MP-AES offers a unique alternative to both FAA and ICP-OES.

These features make the MP-AES an attractive technique for many small to medium size laboratories, particularly those at remote locations, and for an increasing number of laboratories requiring the lowest possible on-going operating costs.

Experimental

Instrumentation

The measurements were performed on an Agilent 4100 MP-AES using a dewar nitrogen supply. The 4100 MP-AES is a compact bench-top microwave plasma atomic emission spectrometer that generates a robust, magnetically-excited nitrogen plasma. Operating the instrument with the optional Agilent 4100 Nitrogen Generator further reduces the operating costs.

The sample introduction system used for this application consisted of a standard torch, a double pass glass cyclonic spraychamber and an inert OneNeb nebulizer.

The determination of Ca, K, Na and Mg benefits from the use of an ionization suppressant. The ionization suppressant was mixed with the sample via a T piece placed before the nebulizer. The on-board three channel peristaltic pump was used to deliver the sample through the sample introduction system. A 0.1% w/v Cs (CsCl Analar, Merck) solution was used as an ionization suppressant.

The External Gas Control Module (EGCM) was used to inject air into the plasma when running the diluted wine matrix that contained a small amount of alcohol. The air injection prevents any carbon build up in the torch, ensuring stable results when running these samples over a long time period.

The air injection also reduces the background emissions generated by the organics present in the sample. The EGCM is automatically controlled by the instrument software, and as such requires minimal user interaction.

Because the amount of alcohol in diluted wine samples is low, the air injection rate is selected at a lower rate than the default setting for each wavelength.

The instrument operating conditions are listed in Table 1.

Table 1. Agilent 4100 MP-AES operating conditions

| Parameter | Value | | | | |
|----------------------------|--------------|-------------|---------|---------|---------|
| Element | Са | К | Na | Mg | Fe |
| Wavelength (nm) | 396.847 | 769.897 | 589.592 | 285.213 | 371.993 |
| EGCM setting | Low | Low | Low | Low | Medium |
| Nebulizer | OneNeb | | | | |
| Spraychamber | Double pa | ass glass c | yclonic | | |
| Pump rate | 15 rpm | | | | |
| Sample tubing | Orange/green | | | | |
| Waste tubing | Blue/blue | 9 | | | |
| Read time | 1-10 seco | nds* | | | |
| Number of replicates | 3 | | | | |
| Sample uptake delay | 15 second | ls | | | |
| Stabilization delay | 20 second | ls | | | |
| Fast pump during uptake | On | | | | |
| Background correction | Auto | | | | |
| | | | | | |

*Can be varied based on sample concentrations

For comparison purposes, the samples were also measured on an Agilent 725 radially-viewed ICP-OES instrument and an Agilent 240FS FAA spectrometer.

Standard and Sample Preparation

A variety of wine samples were selected for this study, covering both red and white varieties.

- Wine 1 : Shiraz
- Wine 2 : Cabernet Sauvignon
- Wine 3 : Chardonnay
- Wine 4 : Sauvignon Blanc
- Wine 5 : Viognier

Additionally, two certified reference materials were analyzed to validate the method:

- Red wine: TM-Wine-R1A (Spex CertiPrep)
- White wine: TM-Wine-W1A (Spex CertiPrep)

For MP-AES and ICP-OES analysis, the samples were degassed in an ultrasonic bath, then diluted 1 in 10 (v/v) with 5% HNO_3 (Suprapur, Merck). Standards and blank were prepared in 5% v/v HNO_3 and 2% v/v ethanol (Merck) to matrix match the alcohol content of the wine samples. Care must be taken when adding ethanol into 5% HNO_3 . Ethanol should be added gradually drop-wise with a Pasteur pipette.

For AA analysis, the samples were also degassed and further sample preparation for AA depends on the element of interest.

- For Ca samples were diluted 1 in 10 with 5% HNO₃ and 2000 mg/L Sr (Strontium chloride, Laboratory reagent, BDH).
- For K and Na samples were diluted 1 in 10 with 5% HNO, and 1000 mg/L Cs.
- For Mg and Fe samples were diluted 1 in 10 with 5% $\mathrm{HNO}_{\mathrm{s}}.$

The standards and blanks were matrix matched with the samples, as described above.

Results

Method detection limit

Method detection limit (MDL) is expressed as 3 times the standard deviation of 10 replicate measurements of the blank. Analytical wavelengths used and the MDL by MP-AES are listed in Table 2.

Table 2. Method detection limits (MDL) by MP-AES

| Element | Wavelength (nm) | MDL (µg/L) |
|---------|-----------------|------------|
| Ca | 396.847 | 8 |
| К | 769.897 | 110 |
| Na | 589.592 | 15 |
| Mg | 285.213 | 11 |
| Fe | 371.993 | 15 |

Certified Reference Material and Wine Samples

The accuracy of the measurement of metals in wine samples by MP-AES was verified by the analysis of the certified red and white wine reference material. Good agreement was obtained with certified values, with recoveries between 94% and 110% (see Table 3). Results for the analysis of wine samples by all three techniques can be seen in Table 4. For the five wines analyzed, the MP-AES results are in good agreement with the AA and ICP-OES results.

Table 3. Analysis of CRM samples by MP-AES

| Element | Measured | Certified-TM-Wine-W1A | % Recovery |
|---------|---------------|-----------------------|------------|
| | mg/L | mg/L | |
| Ca | 79 ± 1 | 82.2 ±2 | 96 |
| К | 980 ± 23 | 939 ± 142 | 104 |
| Na | 27.6 ± 0.4 | 25.1 ± 3 | 110 |
| Mg | 119 ± 1 | 123 ± 3 | 97 |
| Fe | 2.03 ± 0.01 | 1.97 ± 0.2 | 103 |

| Element | Measured | Certified-TM-Wine-R1A | % Recovery |
|---------|-------------|-----------------------|------------|
| | mg/L | mg/L | |
| Ca | 47 ± 0.31 | 50 ±2 | 94 |
| К | 1160 ± 32 | 1120 ± 142 | 104 |
| Na | 21.0 ± 0.4 | 22.4 ± 3 | 96 |
| Mg | 127 ± 1 | 123 ± 3 | 103 |
| Fe | 2.43 ± 0.03 | 2.49 ± 0.2 | 98 |

Table 4. Comparison of the analysis of wine sample by three techniques

| Element | Concentration (mg/L) | | |
|---------|----------------------|----------|-------------|
| | 4100 MP-AES | 240FS AA | 725 ICP-0ES |
| Wine 1 | | | |
| Ca | 52 | 52 | 54 |
| К | 1205 | 1116 | 1112 |
| Na | 37 | 37 | 35 |
| Mg | 148 | 149 | 150 |
| Fe | 1.2 | 1.1 | 1.0 |
| Wine 2 | | | |
| Ca | 6.6 | 6.9 | 6.9 |
| К | 1206 | 1197 | 1154 |
| Na | 30 | 34 | 32 |
| Mg | 103 | 100 | 102 |
| Fe | 2.2 | 2.2 | 2.0 |
| Wine 3 | | | |
| Ca | 56 | 59 | 59 |
| К | 900 | 848 | 839 |
| Na | 34 | 33 | 31 |
| Mg | 87 | 86 | 90 |
| Fe | 0.9 | 0.9 | 0.7 |
| Wine 4 | | | |
| Ca | 70 | 70 | 77 |
| К | 756 | 718 | 741 |
| Na | 10 | 11 | 9.0 |
| Mg | 78 | 77 | 83 |
| Fe | 0.4 | 0.4 | 0.3 |
| Wine 5 | | | |
| Ca | 32 | 31 | 34 |
| К | 689 | 627 | 661 |
| Na | 48 | 48 | 45 |
| Mg | 121 | 125 | 134 |
| Fe | 1.8 | 1.7 | 1.7 |

Conclusion

The MP-AES is an accurate and reliable technique for this application and is an ideal alternative to FAA and ICP-OES. Results for certified samples were in good agreement with the CRM reference values and results for various wine samples were in good agreement across all three techniques.

The MP-AES also offers significant benefits over the commonly used FAA, including enhanced productivity through greatly simplified sample preparation and unattended multi-element analysis, higher performance through improved detection limits and greater linear dynamic range, and lower cost of ownership and operating costs by running on nitrogen and eliminating flammable gases such as acetylene and nitrous oxide.

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Food & Agriculture

With high sample throughput and fast sequential measurement, Agilent MP-AES is ideal for food screening laboratories. MP-AES is ideal for contract laboratories where fast turn around is key, including small to mid-sized screening laboratories needing to determine essential nutrients and elements at major levels, and toxic elements at trace levels.

- With the lowest cost of ownership, the MP-AES will give you the edge over your competition by reducing your cost per analysis and improving performance
- Eliminate downtime waiting for gas refills, and achieve fast sample turn around with safe, reliable unattended analysis
- Reduce sample preparation. The axially viewed vertical plasma handles a wide range of samples — from food and soil digests to high salt soil extractions
- Improved long term stability with integrated nebulizer gas humidification as standard on MP-AES
- Increase your sample throughput, compared to conventional flame AA systems, and never have to change burners/gases for different elements again
- Rapid method development and fast start-up means any user can achieve optimum performance
- A MultiCal feature allows the analysis of elements at high and low levels in the same run
- Complete wavelength coverage means you can avoid spectral interferences from majors by simply choosing another wavelength
- Achieve rapid sub-ppb detection of As, Hg and Se with the Multimode Sample Introduction System (MSIS)
- Include P and S in your elemental analysis suite with better detection limits using MP-AES plasma emission technology

Author

Elizabeth Kulikov Agilent Technologies Australia

Determination of available micronutrients in DTPA extracted soils using the Agilent 4210 MP-AES

Application note Food safety and agriculture

Introduction

Micronutrient soil analysis is commonly conducted in agricultural laboratories to assess the quality of soil for plant development and crop yield. Micronutrients such as copper, iron, manganese and zinc can be extracted from soil using solutions containing chelating agents such as diethylenetriaminepentaacetic acid (DTPA).

Typically, the determination of micronutrients in soils is conducted using Flame Atomic Absorption Spectroscopy (FAAS) or Inductively Coupled Plasma Optical Emission Spectroscopy (ICP-OES); however, with agriculture

labs increasingly under pressure to reduce operating costs and improve safety, Microwave Plasma Atomic Emission Spectroscopy (MP-AES) is gaining recognition as a suitable alternative to these techniques.

Why use MP-AES over traditional techniques?

The advantages of MP-AES for the analysis of environmental samples, including soils include:

- Lower running costs and improved safety. MP-AES uses nitrogen gas from either a Dewar or extracted from air using the Agilent 4107 Nitrogen Generator. Eliminating the need for expensive and hazardous gases such as acetylene allows for unattended analysis. It is the ideal instrument for laboratories looking to reduce on-going operating costs or with safety concerns.
- Excellent analytical performance for difficult samples. The stable microwave plasma is capable of analyzing complex matrices such as DTPA soil extracts or soil digests containing high total dissolved solids (TDS), as well as aqueous solutions.
- Multi-elemental analysis.

MP-AES offers improved analytical performance, lower detection limits and a wider calibration range compared to Flame Atomic Absorption Spectroscopy.

• Ease of use.

MP-AES uses intuitive MP Expert software and plug-and-play hardware to simplify instrument setup, method development and analytical performance, with minimal training. Additionally, application specific software applets can be created in MP expert from pre-set templates, further simplifying analysis.

This application note describes the determination of micronutrients Cu, Fe, Mn and Zn in soils following DTPA extraction using the Agilent 4210 MP-AES.

Experimental

Instrumentation

All measurements were performed using the Agilent 4210 MP-AES with it's integrated humidifier accessory and SPS 4 autosampler. The instrument was set up with the standard sample introduction system comprising the Agilent OneNeb Series 2 nebulizer, double-pass glass cyclonic spray chamber and Easyfit torch. Instrument method parameters and analyte settings are listed in Table 1.

 Table 1. Agilent 4210 MP-AES instrument and method parameters.

| Parameter | Value | | | | |
|-----------------------------|------------------------|---------|----------|---------|--|
| Element | Cu | Fe | Mn | Zn | |
| Wavelength (nm) | 324.754 | 259.940 | 257.610 | 213.857 | |
| Nebulizer | | OneNeb | Series 2 | | |
| Nebulizer flow rate (L/min) | | 0.1 | 75 | | |
| Pump rate (rpm) | 15 | | | | |
| Sample pump tubing | Orange/Green Solvaflex | | | | |
| Waste pump tubing | Blue/blue Solvaflex | | | | |
| Read time (s) | 3 | | | | |
| Number of replicates | 3 | | | | |
| Sample uptake delay (s) | 35 | | | | |
| Rinse time (s) | 20 | | | | |
| Stabilization time (s) | 10 | | | | |
| Background correction | Auto | | | | |
| Gas source | Dewar nitrogen | | | | |

Standard and sample preparation

The soil samples were supplied dried and ground. The extraction solution comprised 0.005 M diethylenetriaminepentaacetic acid (DTPA), 0.01 M calcium chloride dihydrate (CaCl₂.2H₂O) and 0.1 M triethanolamine (TEA).

1.97 g of DTPA, 1.47 g $CaCl_22H_20$ and 13.3 mL TEA were dissolved separately in distilled water and combined. The pH was adjusted to 7.3 using conc. HCl and the volume made up to 1 L with distilled water.

10 g of soil was weighed and 20 mL of the DTPA extraction solution was added. After shaking for 120 minutes, the sample was filtered using filter paper.

Multi-element calibration standards were prepared at the following concentrations: 0.5, 2.5 and 5.0 μ g/mL of Cu and Zn, 5.0, 25.0 and 50.0 μ g/mL of Mn and 25.0, 50.0 and 100.0 μ g/mL of Fe. All calibration blanks and standards were prepared in the DTPA extraction solution.

Results and discussion

Working concentration range

Linear calibrations were obtained for all four elements with calibration coefficients greater than 0.999 (Table 2) and less than 10% calibration error for each point. As an example, Figure 1 shows the calibration curve for Cu 324.754 nm and the calibration error for each calibration point (Table 3).

 Table 2. Wavelength and working calibration concentration range.

| Element and line (nm) | Concentration range (µg/mL) | Concentration coefficient |
|-----------------------|--------------------------------|------------------------------|
| Cu 324.754 | 0.5-5 | 1.000 |
| Fe 259.940 | 10-100 | 0.999 |
| Mn 257.610 | 5-50 | 0.999 |
| Zn 213.857 | 0.5-5 | 0.999 |

Figure 1. The calibration curve for Cu 324.754 nm shows excellent linearity across the calibrated range with a correlation coefficient of 1.00000.

 Table 3. Calibration error (%) for each calibration point for Cu 324.754 nm.

| Standards | Calibration error (%) |
|------------|-----------------------|
| Blank | 0.00 |
| Standard 1 | 1.31 |
| Standard 2 | 0.59 |
| Standard 3 | 0.94 |

Method detection limits

Three sigma method detection limits (MDL) were determined from ten replicate measurements of the $0.5 \ \mu g/mL$ spiked blank DTPA extraction solution during the analytical run. The results shown in Table 4 are the average of 3 analytical runs.

Table 4. Agilent 4210 MP-AES element wavelengths used for analysis andMDLs at a sampling weight of 10 g for the DTPA extraction.

| Element | Wavelength (nm) | MDL (mg/kg) |
|---------|-----------------|-------------|
| Cu | 324.754 | 0.06 |
| Fe | 259.940 | 0.03 |
| Mn | 257.610 | 0.03 |
| Zn | 213.857 | 0.05 |

Spike recoveries

To verify the accuracy of the method, a DTPA-extracted soil sample was spiked with Cu, Fe, Mn and Zn at 5, 40, 20 and 5 mg/kg concentration levels respectively. The recoveries for the spiked sample are given in Table 5. The recovery results were within \pm 10% of the expected value for all 4 analytes which highlights the suitability of the method for the application.

 Table 5. Agilent 4210 MP-AES spike recoveries for all elements in the DTPA extracted soil sample.

| Element and line (nm) | DTPA extracted soil sample (mg/kg) | Spiked concentration (mg/kg) | Measured concentration (mg/kg) | Recovery (%) |
|-----------------------------|---|------------------------------------|--------------------------------------|-----------------|
| Cu 324.754 | 0.43 | 5 | 4.58 | 92 |
| Fe 259.940 | 22.81 | 40 | 36.46 | 91 |
| Mn 257.610 | 6.56 | 20 | 18.09 | 90 |
| Zn 213.857 | 0.23 | 5 | 4.62 | 92 |

Long term stability

Long term stability of the Agilent 4210 MP-AES was measured by analyzing a DTPA extracted soil sample approximately every 2 minutes over 3 hours of continuous measurement. Figure 2 shows that excellent stability was achieved, with measurement precision <2% RSD for all elements (see Table 6), over the 3-hour period.

Figure 2. Normalized concentration of Cu, Fe, Mn and Zn in DTPA extracted soil sample, measured over 3 hours.

Table 6. Agilent 4210 MP-AES long term stability results (% RSD) for Cu, Fe, Mn and Zn in DTPA extracted soil sample.

| Element | Wavelength (nm) | %RSD |
|---------|-----------------|------|
| Cu | 324.754 | 1.77 |
| Fe | 259.940 | 1.45 |
| Mn | 257.610 | 1.38 |
| Zn | 213.857 | 1.21 |

Conclusions

The Agilent 4210 MP-AES proved suitable for the costeffective analysis of micronutrients in DTPA extracted soil samples. As the microwave plasma is generated from nitrogen gas, it eliminates the need for expensive and flammable gases, which reduces operational costs and improves lab safety. Compared to FAAS, the high plasma temperature (5000 K) of MP-AES provides a higher sample matrix tolerance, lower detection limits and an expanded working

concentration range.

The method used in this study demonstrated:

- High analytical performance with excellent MDLs and spike recoveries for all elements within \pm 10% of the target values.
- Excellent linearity across a wide concentration range.
- Excellent long term stability, with less than 2% RSD over a 3-hour period.

Determination of major elements in milk using the Agilent 4200 MP-AES

Application note

Food testing & agriculture

Authors

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Introduction

Milk is one of the most important food commodities in the world and its consumption has grown, particularly in developing countries which have experienced strong economic growth and urbanization in recent decades.

As a substantial source of several nutrients such as proteins, enzymes, fats, vitamins and essential elements (also known as minerals), milk plays a key role during all phases of human life. Rapid growth during infancy and early childhood creates high demand for the nutrients that milk provides. This development phase requires a balanced amount of different elements, as mineral deficiencies may impair body development whilst excessive mineral intake may increase the osmotic load and cause complications in the developing kidneys of a child.

Essential elements such as Ca, K, Mg, Na and P have several physiological functions in the tissue structure of humans and other animals, such as maintaining osmotic/electrolyte balance, and acting as a cofactor for many enzymes. Deficiencies in these essential elements causes disturbances

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in the physiological system in any stage of life and, for this reason, such elements must be monitored to ensure the nutritional value of foods. The accurate analysis of essential elements is particularly important in extensively consumed products, such as milk.

Several atomic spectroscopy techniques are routinely used for elemental quantification in milk and dairy products, in particular flame atomic absorption spectrometry (FAAS) and recently microwave plasma atomic emission spectrometry (MP-AES).

The recent introduction of microwave plasma atomic emission spectrometry was a breakthrough revolution in entry-level atomic spectroscopy techniques. The easy to use Agilent 4200 MP-AES has better performance and speed than a FAAS and requires no hazardous and expensive gases. This improves safety and reduces the cost of analysis.

This work shows the performance of the Agilent 4200 MP-AES for quantification of Ca, K, Mg, Na and P in fresh and powdered milk after acid digestion, with quality assurance performed by analysing a Certified Reference Material (CRM) and applying some of the concepts from the US EPA Contract Laboratory Program.

Experimental

Instrumentation

For this study a microwave plasma atomic emission spectrometer, the Agilent 4200 MP-AES (Agilent Technologies, Santa Clara, CA) was used for elemental determination of digested milk samples. Acid digestion was carried out using an UltraWAVE Single Reaction Chamber Microwave Digester (Milestone Inc., Shelton CT).

Standards and reagents

Analytical grade concentrated nitric acid (HNO₃ 67-69%) and hydrochloric acid (HCI 32-35%) were used for sample digestion. The 18.2 MΩ deionized water used was obtained from a Milli-Q[™] Water System (Millipore, Darmstadt, Germany). Calibration and accuracy verification standards were prepared using Agilent (Agilent Technologies, Santa Clara, CA) and Spex (SPEX CertiPrep, Metuchen, NJ) Calibration Standards. Method validation was achieved by analyzing the accuracy verification standards and a milk powder Certified Reference Material (CRM), NIST 1549a (NIST, Gaithersburg, MD).

Microwave sample digestion

Seven different powdered and liquid milk samples were purchased from a supermarket in California, USA and digested before analysis by MP-AES (refer to Table 4).

To prepare the milk samples for microwave digestion approximately 0.25 g of each powdered milk, 0.50 g of the powdered NIST 1549a CRM and 1 g of each fresh milk sample was accurately weighed and transferred to a 15 mL Teflon digestion vial. Before capping the vials, 6 mL of nitric acid and 1 mL of hydrochloric acid was added to each. A blank solution was also prepared, containing 6 mL of nitric acid and 1 mL of hydrochloric acid. Each milk sample and blank solution was prepared in triplicate in accordance with this procedure. Similarly, seven samples of the NIST 1549a CRM were prepared and digested in order to evaluate the accuracy of the analytical procedure.

At least two of the sample vials in each batch of 14 samples digested contained the NIST 1549a CRM. One blank solution was included in each batch.

Microwave digestion of the samples was carried out in accordance with the following procedure: The digestion chamber was initially pressurized to 40-45 mTorr with industrial grade nitrogen gas, before the temperature and pressure were gradually increased to 240 °C and 150 bar respectively over 20 minutes. These values were maintained for a further 15 minutes (the duration of the digestion) to ensure complete digestion.

Upon completion of the program each digested sample was diluted to a final volume of 10 mL with deionized water, before a further 10 times dilution with a solution of 2% nitric acid.

Elemental determination

The Agilent 4200 MP-AES has superior performance compared to FAAS in terms of detection limits, linear range, and sample throughput. The 4200 MP-AES uses magnetically-coupled microwave energy to generate a robust and stable plasma using nitrogen gas. The use of nitrogen improves safety by eliminating expensive, hazardous gases and also results in low operational costs. The nitrogen plasma reaches around 5,000 K and eliminates the chemical interferences that are common in FAAS, such as the formation of refractory CaPO₄. This means that the element-specific sample preparation often required in FAAS can be simplified to a single sample preparation for all elements. The more powerful excitation source also enables phosphorus determinations, which is not possible on FAAS. The instrument features mass flow control of the nebulizer gas, and a torch loader mechanism which automatically connects all gases. Method parameters can be automatically optimized in the MP Expert software, which also features automatic background correction.

Method conditions for digested milk sample analysis in the 4200 series MP-AES are listed in Table 1.

 Table 1. MP-AES 4200 operational conditions for Ca, K, Mg, Na, P

 determination in digested milk

| Common Conditions | | | | | |
|----------------------------|----------------------------|--|--|--|--|
| Background Correction Auto | | | | | |
| Nebulizer | Micromist | | | | |
| Spray Chamber | Double pass glass cyclonic | | | | |
| Pump Speed | 10 rpm | | | | |
| Read Time | 2 s | | | | |
| Replicates | 3 | | | | |
| Stabilization Time | 20 s | | | | |
| Viewing Position | 0 | | | | |

| Elemental Conditions | | | | | | |
|-----------------------|------------------------|------|--|--|--|--|
| Element | Nebulizer Flow (L/min) | | | | | |
| Са | 422.673 | 0.4 | | | | |
| К | 766.491 | 0.8 | | | | |
| Mg | 285.213 | 0.4 | | | | |
| Na | 588.995 | 0.4 | | | | |
| Р | 214.915 | 0.35 | | | | |
| Y (Internal Standard) | 371.029 | 0.4 | | | | |

Results and Discussion

Concentrations working range and method detection limit

Calibrations for all elements were between 5 and 100 ppm, and the correlation coefficient was greater than 0.999 for all wavelengths. Method detection limits (MDL) were calculated as 3 times the standard deviation of 10 consecutive blank readings (3σ). From the MDL, the method quantification limit (MQL) was

calculated as $3.33 \times MDL$. The MDL and MQL are summarized in Table 2.

Table 2. Method Detection Limits (MDL) and Method Quantification Limits (MQL) in mg/L.

| Element/ Wavelength (nm) | MDL | MQL ⁽¹⁾ |
|-----------------------------|-------|---------------------------|
| Ca 422.673 | 0.002 | 0.007 |
| K 766.491 | 0.067 | 0.223 |
| Mg 285.213 | 0.002 | 0.007 |
| Na 588.995 | 0.117 | 0.351 |
| P 214.915 | 0.318 | 1.059 |

(1) Quantification limits in sample must take into account the different dilution factors applied in powdered milk or fresh milk.

Quality Control

Two strategies were adopted to validate the method:

- 1. Analysis of the NIST 1549a milk CRM in seven independent digestions, analyzed among unknown samples.
- Analysis of Initial Calibration Blank and Initial Calibration Verification solutions (ICB & ICV) immediately after the method calibration, followed by Continuing Calibration Blank and Continuing Calibration Verification (CCB & CCV) solutions every 10 samples. The ICB/CCB and ICV/CCV analyses totalled four runs each.

The results from this analysis, shown in Table 3, highlight the ability of the MP-AES to reliably analyze digested milk samples with excellent accuracy, precision and minimal carryover between solutions.

 Table 3. Summarized results and recoveries of NIST 1549a CRM, ICB/CCB and ICV/CCV samples.

| | Ca | К | Mg | Na | Р |
|---|---------------|---------------|---------------|---------------|------------|
| CRM Reference Value (mg/kg) | 8810 ± 240 | 11920 ± 430 | 892 ± 62 | 3176 ± 58 | 7600 ± 500 |
| CRM Measured Conc. (n=7) \pm SD (mg/kg) | 9031 ± 195 | 11683 ± 566 | 928 ± 15 | 3373 ± 108 | 7360 ± 96 |
| CRM Recovery (%) | 102.5 | 98.0 | 104.1 | 106.2 | 96.8 |
| ICB/CCB (n=4) Average \pm SD (mg/kg) | 0.020 ± 0.001 | 0.780 ± 0.155 | 0.004 ± 0.001 | 0.411 ± 0.212 | < MDL |
| ICV/CCV (n=4) Recovery % | 99.9 | 102.6 | 96.7 | 101.9 | 99.7 |

Table 4. Analysis results of powdered and fresh milk digested samples by Agilent 4200 MP-AES.

| Samples | Ca mg/kg (RSD) | K mg/kg (RSD) | Mg mg/kg (RSD) | Na mg/kg (RSD) | P mg/kg (RSD) |
|------------------------------------|-------------------|------------------|-----------------------------------|-------------------|------------------|
| Powdered Instant Nonfat Milk | 11953 (4.5%) | 15296 (3.6%) | 1242 (4.4%) | 4141 (3.5%) | 9611 (1.0%) |
| Powdered Nonfat Milk | 11058 (3.8%) | 16057 (6.6%) | 1176 (1.9%) | 4167 (5.4%) | 9223 (1.1%) |
| Powdered Organic Buttermilk | 9659 (1.5%) | 27253 (1.37%) | 1116 (3.8%) | 4069 (3.7%) | 8489 (2.3%) |
| Powdered Sweet Cream Buttermilk | 8287 (6.6%) | 14421 (8.8%) | 14421 (8.8%) 1053 (5.9%) 4784 (7. | | 7920 (5.9%) |
| Powdered Whole Milk | 8592 (3.1%) | 15157 (3.1%) | 1218 (1.0%) | 3010 (3.3%) | 7750 (0.9%) |
| Fresh Whole Milk | 1150 (2.9%) | 1687 (1.9%) | 109 (0.9%) | 407 (1.9%) | 898 (0.8%) |
| Fresh Nonfat Milk | 1182 (1.6%) | 1726 (0.6%) | 112 (1.3%) | 412 (0.4%) | 904 (0.6%) |

Sample Analysis

To evaluate the performance of the method with real samples, digested powdered and fresh milk samples were also analyzed (shown in Table 4). These results demonstrate the ability of this method to analyze a diverse collection of real samples with good precision, easily covering the vast range of major element concentrations determined (e.g. from 1150 to 11953 mg/ kg for Ca).

Conclusion

An accurate and robust method has been developed for the determination of major elements in digested milk samples on the 4200 MP-AES. The detection limits achieved were found to be well below those required for milk analysis, and excellent recoveries were obtained for the CRM (between 110—90%) and ICV/CCV (between 105—95%).

The 4200 MP-AES is the ideal instrument for those looking to move away from FAAS and extend their laboratory's analytical capabilities. Recognized benefits of the MP-AES include reduced running costs, enhanced productivity through numerous ease-of-use features and simplified sample preparation, improved safety, and higher analytical performance such as better detection limits and greater linear dynamic range.

References

[1] Khan, N.; Jeong, I. S.; Hwang, I. M.; Kim, J. S.; Choi, S. H.; Nho, E. Y.; Choi, J. Y.; Park, K. S.; Kim, K. S; Analysis of minor and trace elements in milk and yogurts by inductively coupled plasma-mass spectrometry (ICP-MS), *Food Chemistry* 147 (2014) 220–224.

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Determination of major, minor and trace elements in rice flour using the 4200 Microwave Plasma-Atomic Emission Spectrometer (MP-AES)

Application note

Food testing

Introduction

The analysis of foodstuffs, such as rice, is of particular interest for nutrient elements present at high concentrations, and also for toxic elements such as cadmium that can be present at trace levels. The analysis is important to ensure product quality and safety, as well as determining product origin. Food scares related to contamination not only constitute a health risk but also undermine consumer confidence. This can lead to lost earnings through reduced sales and loss of credibility through adverse publicity.

Flame Atomic Absorption Spectroscopy (FAAS) is well established for the analysis of foods, but with lab budgets coming under increasing pressure and current market trends for lower cost of ownership, improved

Authors

John Cauduro Agilent Technologies, Australia performance, ease of use, and safety, many FAAS users are looking to transition to another technique to expand their analytical capabilities.

Agilent has expanded its atomic spectroscopy portfolio to include the Microwave Plasma-Atomic Emission Spectrometer. The Agilent 4200 MP-AES is the second generation microwave plasma instrument that features an improved waveguide design that is capable of running samples with high total dissolved solids without compromising detection limits. The 4200 MP-AES significantly reduces running costs through the use of nitrogen as its plasma gas. The use of nitrogen also increases safety, by removal of flammable gases, and allows unattended operation of the instrument. The 4200 MP-AES is easy to use, and is able to achieve lower detection limits than a standard FAAS, as well as being able to determine additional elements such as phosphorus.

This application describes the analysis of rice flour for cadmium and other major, minor and trace elements on the 4200 MP-AES.

Experimental

Instrumentation

The innovative 4200 MP-AES features a second generation waveguide and torch, with mass flow controlled nebulizer gas flow. The 4200 MP-AES has a robust toroidal plasma with a central channel temperature of ~5,000 K which eliminates many of the chemical interferences that are present in FAAS and also expands the concentration working range of the 4200 MP-AES when compared the FAAS. This means that the element specific sample preparation that is commonplace when using FAAS is not necessary when using the 4200 MP-AES, improving ease of use and reducing cost. The 4200 MP-AES also achieves lower detection limits than FAAS, particularly for phosphorus, which enables the analysis of extra elements. By running on nitrogen, the 4200 MP-AES offers reduced operating costs and increased lab safety compared to flame AA, through the avoidance of flammable and costly gases such as acetylene, and nitrous oxide.

The analysis was carried out using a 4200 MP-AES equipped with the standard sample introduction setup consisting of the OneNeb nebulizer and a double pass spray chamber. An SPS 3 autosampler was used to deliver samples to the instrument, allowing the system to be operated unattended.

The MP-AES features continuous wavelength coverage which allows the analyst to select wavelengths that are appropriate for the expected concentration range, and free from spectral interferences. Method conditions for the selected wavelengths are shown in Table 1 and common method conditions are shown in Table 2.

Table 1. Agilent 4200 MP-AES operating parameters

| Element | Wavelength | Read time (s) | Nebulizer Flow (L/min) |
|---------|------------|---------------|------------------------|
| Р | 214.915 nm | 2 | 0.55 |
| Cd | 228.802 nm | 10 | 0.55 |
| Mg | 280.271 nm | 1 | 0.55 |
| Zn | 213.857 nm | 5 | 0.55 |
| Mn | 403.076 nm | 3 | 0.55 |
| К | 766.491 nm | 1 | 0.55 |
| Cu | 324.754 nm | 2 | 0.75 |
| Fe | 438.354 nm | 5 | 0.75 |
| Са | 422.673 nm | 1 | 1.00 |

Sample preparation

NIES CRM No.10c Rice Flour (NIES, Japan) was analyzed to validate the analytical method. The rice flour samples were digested using a Milestone Ethos microwave digestion system¹. Samples were prepared in duplicate with approximately 0.5 g of rice flour CRM accurately weighed into separate TFM vessels. This was followed by the addition of 7 mL of HNO₃ and 1 mL of H₂O₂ and placed in the microwave digestion unit. The samples were digested using the preloaded digestion methods, allowed to cool, and then made up to 25 mL with deionized water. The final solution contained 2% total dissolved solids. No ionization suppressants or matrix modifiers were required for the analysis.

Calibration range

The calibration concentration range of the standard solutions are summarized in Table 3. As the working range of 4200 MP-AES far exceeds that of FAAS (by up to 20x in some instances), only one dilution of the sample is required to measure the complete set of elements. The calibration fit for all wavelengths used was linear.

Results and Discussion

Method detection limits (MDLs)

MDLs were determined from the analysis of 10 digested blank samples. The MDLs (3o) for the selected analytical wavelengths are listed in Table 4.

| Table 2. Common method conditions | | Table 3. Workir | ng concentration rang | e of the 4200 MP | AES standard solutions |
|-------------------------------------|------------------------------------|-----------------|-----------------------|------------------|------------------------|
| Parameter | Value | Element | Wavelength | Concentration | n range |
| Replicates | 3 | Р | 214.915 nm | 0–100 | ppm |
| Pump rate | 15 rpm | Cd | 228.802 nm | 0–1.0 | ppm |
| Sample uptake delay | 30 seconds | Mg | 280.271 nm | 0—40 | ppm |
| Rinse time | 60 seconds | Zn | 213.857 nm | 0-4.0 | ppm |
| Stabilization time | 10 seconds | Mn | 403.076 nm | 0–1.0 | ppm |
| Fast Pump during Uptake and Rinse | On (80 rpm) | К | 766.491 nm | 0–100 | ppm |
| Nebulizer | OneNeb | Cu | 324.754 nm | 0–1.0 | ppm |
| Spray chamber | Double pass cyclonic | Fe | 438.354 nm | 0–1.0 | ppm |
| Autosampler | Agilent SPS 3 | Са | 422.673 nm | 0-4.0 | ppm |
| Sample pump tubing | Orange/green | | | | |
| Waste pump tubing | Blue/blue | | | | |
| Table 4 Agilant 4200 MP AES alamant | wavelength and MDL (mg/kg in compl | o) | | | |

Table 4. Agilent 4200 MP-AES element wavelength and MDL (mg/kg in sample)

| Element/ Wavelength (nm) | Ca 422.673 | Cd 228.802 | Cu 324.754 | Fe 438.354 | K 766.491 | Mg 280.271 | Mn 403.076 | P 214.915 | Zn 213.857 |
|--------------------------------|------------|------------|------------|------------|-----------|------------|------------|-----------|------------|
| MDL (mg/kg) | 0.10 | 0.16 | 0.05 | 0.44 | 3.0 | 0.06 | 0.05 | 13 | 0.15 |

Analysis of certified reference material

Results of the analysis of major, minor and trace elements in rice is listed in Table 5. The measured values (average result on two different 4200 MP-AES instruments carried out in duplicate) are in good agreement with the certified values for all CRM samples. The results demonstrate the capability of the 4200 MP-AES to achieve excellent results across a wide concentration range in a sample with 2% dissolved solids.

Long term stability

A digested rice flour sample was repeatedly analyzed under method conditions over 8 hours to test the long term stability of the method. The test was performed under controlled laboratory environmental conditions within the instrument operating specification, with a recalibration every 2 hours. The resulting stability plot is show in Figure 1. Excellent stability of < 3 % RSD for all elements was achieved, demonstrating the capability of the 4200 MP-AES, OneNeb nebulizer and mass flow controlled nebulizer gas flow to handle 2% total dissolved solids.

Table 5. Results of NIES No.10c Rice Flour. All results in mg/kg in the solid sample.

| Element/ Wavelength (nm) | Ca 422.673 | Cd 228.802 | Cu 324.754 | Fe 438.354 | K 766.491 | Mg 280.271 | Mn 403.076 | P 214.915 | Zn 213.857 |
|--------------------------------|------------|------------|------------|------------|-----------|------------|------------|-----------|------------|
| Mean | 96.0 | 1.96 | 4.13 | 11.50 | 2700 | 1174 | 37.35 | 3139 | 22.02 |
| SD | 2.5 | 0.11 | 0.29 | 1.03 | 105 | 23 | 1.04 | 92 | 0.48 |
| Certified value | 95 | 1.82 | 4.1 | 11.4 | 2750 | 1250 | 40.1 | 3350 | 23.1 |
| 2SD certified | 2 | 0.06 | 0.3 | 0.8 | 100 | 80 | 2.0 | 80 | 0.9 |
| % difference | 101.0 | 107.7 | 100.8 | 100.9 | 98.2 | 93.9 | 93.1 | 93.7 | 95.3 |

Figure 1. Normalized concentration of elements in a rice flour digest analyzed over an 8 hour time period, with recalibration every 2 hours.

Conclusion

A method for the determination of major, minor and trace elements in rice flour has been described. The next generation 4200 MP-AES achieved recoveries in a rice CRM of +/- 10 % of the assigned value, with MDLs sufficient for the analysis and excellent long term stability.

The excellent analytical performance, including phosphorus which is not practical by FAAS, multielement unattended operation, improved safety and ease of use make the 4200 MP-AES the ideal alternative for FAAS users looking to transition to a new technique. Furthermore, the sample preparation process can be simplified, with no modifiers or ionization suppressants required due to the higher temperature excitation source of the MP-AES.

Reference

1. Milestone Application Note. Food/Feed. Rice Flour. ID HPR-FO-39. Milestone Ethos with internal temperature sensor, HPR1000/10S high pressure segmented rotor.

Determination of major elements in fruit juices using the Agilent 4200 MP-AES with the Agilent 4107 Nitrogen Generator

Application note

Food testing

Authors

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Introduction

Major elements such as calcium, magnesium, sodium and potassium are essential nutrients in food and the routine monitoring of the levels of these elements in fruit juices is a common quality control process. Flame Atomic Absorption Spectroscopy (FAAS) is well suited to this application as it delivers the performance required for the analysis at a reasonable price. However, with the introduction of the Agilent Microwave Plasma-Atomic Emission Spectrometer (MP-AES), several of the analytical challenges of using FAAS for this application have been overcome, making it the ideal instrument for laboratories looking to transition away from FAAS to a more powerful and safer technique. The 4200 MP-AES main operating gas, nitrogen, is supplied from an Agilent 4107 Nitrogen Generator (with air supplied from an air compressor). This greatly reduces the running costs and eliminates the safety concerns associated with specialty gases required by FAAS such as acetylene and nitrous oxide. The nitrogen-based plasma source of the 4200 MP-AES operates at a higher temperature than the flame source of a FAAS, avoiding the chemical interferences present in FAAS (especially for elements such as Ca). This eliminates the time consuming, element specific sample preparation and burner head changeover that is required when analyzing Ca, Na, K and Mg in the same sample by FAAS. The analysis of this application by MP-AES also removes the need for costly and time consuming modifiers and ionization suppressants. The plasma source in the MP-AES also leads to improved performance with respect to detection limits and linear dynamic range when compared to FAAS, which is important in an analysis where the elements can be present over a wide range of concentrations. With no flammable gases required, the MP-AES is able to operate unattended which increases sample throughput.

This application note describes the analysis of fruit juice samples using an Agilent 4200 MP-AES running with an Agilent 4107 Nitrogen Generator.

Experimental

Instrumentation

All measurements were performed using an Agilent 4200 MP-AES, with nitrogen supplied from an Agilent 4107 Nitrogen Generator. The sample introduction system consisted of a double pass spray chamber and OneNeb nebulizer. The instrument was controlled by the powerful and easy-to-use MP Expert software. The MP-AES features continuous wavelength coverage and MP Expert features an extensive wavelength database that allows the selection of wavelengths that are appropriate for the concentration range required for the analysis. For instance, in this application, the less sensitive Mg 518.360 nm line was preferred over the more sensitive Mg 285.213 line.

| 5 | | | | |
|-------------------------|-------------|---------------|-----------|---------|
| Parameter | Value | | | |
| Element | Са | Mg | Na | К |
| Wavelength | 422.673 | 518.360 | 589.592 | 769.897 |
| Nebulizer | OneNeb | | | |
| Nebulizer flow rate | Default (0. | 75 L/min) | | |
| Spray chamber | Double pas | ss glass cycl | onic | |
| Pump rate | 15 rpm | | | |
| Sample pump tubing | Orange/gr | een | | |
| Waste pump tubing | Blue/blue | | | |
| Autosampler | Agilent SP | S 3 | | |
| Read time | 1 second | | | |
| Number of replicates | 3 | | | |
| Fast pump during uptake | On | | | |
| Sample uptake delay | 30 seconds | 6 | | |
| Rinse time | 40 seconds | 6 | | |
| Stabilization time | 20 seconds | 6 | | |
| Background correction | Auto | | | |
| Gas source | Agilent 410 |)7 Nitrogen | Generator | |

| | Table | 1. Agilent | 4200 I | MP-AES | operating | conditions |
|--|-------|------------|--------|--------|-----------|------------|
|--|-------|------------|--------|--------|-----------|------------|

Standard and sample preparation

Two quality control (ΩC) test materials were analyzed to validate the method:

- Apple Juice T1650QC (certified by FAPAS*)
- Grapefruit Juice T0842QC (certified by FAPAS*)

*FAPAS – The Food and Environmental Research Agency, York, UK.

Materials were purchased from Graham B Jackson (Aust) P/L.

Additionally, a commercially available apple juice was analyzed in the long term stability studies.

All fruit juice samples were diluted 20x with 5% HNO₃ (ACS Grade, Merck). No other modifiers or ionization suppressants were required.

Standards were prepared from a 10,000 mg/L multi element standard (Inorganic Ventures). All calibration blanks and standards were prepared in 5% HNO₃.

Results

Working range

The working concentration range of the standard solutions are summarized in Table 2. As the working range of MP-AES far exceeds that of FAAS (by up to 20 times in some instances), only one dilution of the sample is required to measure the complete set of elements.

| Table 2. Working concentration range of the 4200 MP-AES standard soluti | ions |
|---|------|
|---|------|

| Element | 4200 MP-AES concentration range (mg/L) | Correlation coefficient |
|------------|--|-------------------------|
| Ca 422.673 | 0–20 | 0.99990 |
| Mg 518.360 | 0–100 | 0.99988 |
| Na 589.592 | 0–20 | 0.99996 |
| K 769.897 | 0–100 | 0.99968 |

Recoveries

Table 3 shows the concentration and recovery results of the four elements in the two fruit juices. The recovery results for Ca, Mg, Na, K in the fruit juices using this method were within +/- 10% of the assigned value. All results measured in this study were within the certified ranges of the two quality control test materials.

 Table 3. Recovery results of 4 elements in the fruit juices using the 4200

 MP-AES with the nitrogen generator

| Apple Juice | Certified value (n | ng∕L) | Found | % Recovery | |
|---|---|---|---|-------------------------|--|
| T0840QC | Assigned value | Range | (mg/L) | | |
| Magnesium | 49.0 | 40.3–57.8 | 49.9 ± 0.6 | 102 | |
| Sodium | 21.2 | 16.9–25.4 | 22.2 ± 0.5 | 105 | |
| Potassium | 1044 | 926–1161 | 1039 ± 29.7 | 100 | |
| | | | | | |
| Grapefruit | Certified value (n | ng/L) | Found | % Recovery | |
| Grapefruit Juice T0842QC | Certified value (n Assigned value | ng/L) Range | Found (mg/L) | % Recovery | |
| Grapefruit Juice T0842QC Calcium | Certified value (n Assigned value 145.6 | ng/L) Range 123.6–167.6 | Found (mg/L) 158.3 ± 3.2 | % Recovery | |
| Grapefruit Juice T0842QC Calcium Magnesium | Certified value (n Assigned value 145.6 92.5 | ng/L) Range 123.6–167.6 77.5–107.4 | Found (mg/L) 158.3 ± 3.2 91.1 ± 0.6 | % Recovery 109 99 | |

Long term stability

A commercial apple juice solution (diluted 20x with 5% HNO₃) was repeatedly analyzed over a period of 6 hours. The resulting stability plot is shown in Figure 1. All elements have an RSD of less than 4% over 6 hours. With the OneNeb nebulizer and mass flow controlled nebulizer gas flow, excellent stability results were obtained for a sample with a complex high sugar matrix.

Figure 1. Normalized concentration of potassium in an apple juice sample over 6 hours

Cost savings with the 4200 MP-AES

The potential cost saving of using the 4200 MP-AES for this application was estimated by comparing an FAAS purchased with an air compressor and 1 year of consumables to an MP-AES purchased with air compressor, nitrogen generator, SPS 3, and 1 year of consumables. The analysis requirements were assumed to be 500 samples per week and 4 elements per sample. The calculation assumes that the FAAS is run without an autosampler and that 3 elements are analyzed with air/acetylene and 1 element with nitrous oxide/acetylene. In this example the results show an estimated cost saving of greater than US \$220,000 over a 7 year evaluation period¹. A global average gas cost was used in this calculation and results will vary from country to country.

¹This example is intended to help you compare the running costs and savings of the MP-AES vs. flame AA. The applied formulas and parameters are correct to the best of our knowledge, but we cannot guarantee the results. Savings may vary depending on factors such as local gas and electricity costs, operator costs, number and types of elements. For this calculation operator labor costs were set to USD \$25/hour and electricity costs were set to USD \$0.18 per kW.

Conclusion

A simple and rapid method using MP-AES has been developed to analyze Ca, Mg, Na and K in fruit juice. The recoveries obtained from the analysis of the two QC test materials were within +/- 10% of the assigned values and within the certified concentration range. Using the standard sample introduction system supplied with the 4200 MP-AES, excellent long term stability was obtained over a 6 hour period.

The 4200 MP-AES is the ideal instrument for those customers who are looking to transition away from FAAS and extend their laboratory's analytical capabilities. Recognized benefits of MP-AES include reduced running costs, enhanced productivity through numerous ease-of-use features and simplified sample preparation, improved safety, and higher analytical performance such as better detection limits and greater linear dynamic range.

Analysis of aluminum in beverages using the Agilent 4100 Microwave Plasma-Atomic Emission Spectrometer (MP-AES)

Application note

Food Testing

Authors

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Abstract

Aluminum present in a beverages can affect the taste. This application note describes the determination of aluminum in beverages using an Agilent 4100 MP-AES. Beverages contain a variety of matrix constituents, including salt, sugar, and alcohol. A study was performed to determine how these matrix constituents affect aluminum measurements, and what could be done to reduce such effects. It was determined that sufficient analysis is possible so long as the matrix concentration and alcohol concentration are known to some extent.

Introduction

At present, absorption spectrophotometry, atomic absorption spectrophotometry and inductively coupled plasma-atomic emission spectroscopy have been adopted for performing elemental analyses of food. Aluminum content in food and beverages is restricted by municipal water supply quality standards to a maximum of 0.2 mg/L on the basis of the Japanese Waterworks Law. When flame atomic absorption spectroscopy (FAAS) is utilized, low sensitivity for aluminum and high matrix constituents in the beverage can cause problems with burners getting clogged. This study was conducted to see if an MP-AES, instead of an FAAS, could be used for beverage analysis.

Experimental

Instrumentation

The measurements were performed on an Agilent 4100 MP-AES. The 4100 MP-AES is a compact bench-top spectrometer that generates a robust, magnetically-excited nitrogen plasma.

A 2.45-GHz air-cooled magnetron is used to generate a magnetic field around a torch. The skin effect of that magnetic field causes plasma to form in the shape of a donut, just as with inductively coupled plasma, and it becomes possible to introduce liquid samples at a steady rate (see Figure 1). The nitrogen used to generate the plasma can be supplied via a simple air compressor and the Agilent 4107 Nitrogen Generator. A clear advantage of in-house gas generation is the reduced costs of operation and maintenance compared to conventional gas resupply.

The sample introduction system used for this application consisted of a standard torch, a single pass cyclonic spray chamber and a glass concentric nebulizer.

Table 1 lists the instrument operating conditions.

 Table 1. Agilent 4100 MP-AES operating conditions

| Parameter | Value |
|------------------|-----------|
| Microwave power | 1.0 kW |
| Pump speed | 15 rpm |
| Integration time | 3 seconds |

Standard and Sample Preparation

Samples included:

- Barley tea
- Green tea
- Black tea
- Coffee
- Cola
- Sports drink
- Beer
- Shochu highball

Figure 1. Schematic and basic principles behind the Agilent 4100 MP-AES

Results

Quantitative lower limits and stability

The quantitative lower limits and stability in aqueous solutions and in ethanol were measured. A 0.1% nitric acid solution and ethanol diluted at 0.2 mg/L was prepared using standard solutions. The quantitative lower limit was assumed to be a concentration ten times the standard deviation (σ) obtained from repeatedly measuring the blank ten times. The stability was calculated by repeatedly measuring each of the 0.2 mg/L solution ten times (see Table 2).

Table 2. Method detection limits (MDL) by MP-AES

| Aluminum | Limit of quantification (µg/L) | Stability at 0.2 mg/L (%RSD) |
|------------------|-----------------------------------|---------------------------------|
| Aqueous solution | 1.9 | 1.4 |
| Ethanol (100%) | 7.9 | 0.7 |

The results for limit of quantification and stability confirm that microwave plasma atomic emission spectroscopy is sufficiently applicable for the analysis of aluminum in beverages.

The effects of sugar

With the emission intensity of aluminum at 0.2 mg/L with a sugar concentration of 0 g/100 mL given a value of 1, the effects of varying the sugar concentration between 0 and 50 g/100 mL were measured. The sugar concentrations of the samples were: approx. 2-5 g/100 mL for black tea (with sugar), and 11 g/100 mL for cola (see Figure 2).

Figure 2. Variation in emission intensity due to differences in sugar concentration

The measurements show that sugar concentrations up to approximately 10 g/100 mL do not have a significant effect. Additional standards and matrix matching are needed for concentrations above 10 g/100 mL. The samples examined had a sugar concentration of about 11 g/100 mL, so the analysis was performed using the absolute calibration method.

The effects of ethanol

An additional study was conducted to see if the 4100 MP-AES could be used to determine aluminum content in alcoholic beverages sold in aluminum cans. Subjecting alcohol to plasma produced a relatively large amount of carbon in relation to the concentration of alcohol. Carbon can cause the torch injector to become blocked. To prevent this, air was mixed with the support gas before subjecting it to the plasma.

With the emission intensity of aluminum in a 0.2 mg/L solution set to 1, the effects of varying the ethanol content from 0 to 10% in the solution were examined (see Figure 3).

There were no significant variations either with or without air at ethanol concentrations of about 5%, but emission intensity declined at concentrations above 5% with no air added. Beer is approximately 5% alcohol, while some shochu highballs are higher, at about 8%. For that reason, the sample analysis was performed with air added.

Aluminum was added to each sample, and the results of the recovery tests are given in Table 3.

Figure 3. Variation in emission intensity due to differences in ethanol concentration, with and without adding air. Red squares = with air, blue diamonds = without air

Table 3. Aluminum addition recovery tests

| AI | Unspiked (mg/L) | 0.2 mg/L added (mg/L) | Recovery rate (%) |
|-----------------|--------------------|--------------------------|----------------------|
| Barley tea | 0.00 | 0.22 | 110 |
| Coffee | 0.01 | 0.23 | 109 |
| Sports drink | 0.01 | 0.22 | 105 |
| Cola | 0.05 | 0.24 | 96 |
| Beer | 0.04 | 0.23 | 96 |
| Shochu highball | 0.01 | 0.22 | 105 |
| AI | Unspiked (mg/L) | 1.0 mg/L added (mg/L) | Recovery rate (%) |
| Green tea | 1.14 | 2.12 | 99 |
| Black tea | 2.45 | 3.38 | 98 |

Conclusion

This study has shown that the limit of quantification for aluminum is $1.9 \ \mu g/L$ in aqueous solutions and $7.9 \ \mu g/L$ in ethanol, which adequately meets municipal water supply quality standards as stated in the Japanese Waterworks Law. Favorable results with regard to stability were also obtained. The results of the examination for the effects of beverage matrices (sugar and alcohol) showed that direct measurements without matrix matching can be done for concentrations of about 10 g/100 mL of sugar in beverages, and that the Agilent 4100 MP-AES can also easily and rapidly analyze samples with differing alcohol concentrations if air is mixed in. Thus, it has been demonstrated that the MP-AES has low running costs, is easy to operate, and can perform analyses of aluminum in beverages.

Cost-effective analysis of major, minor and trace elements in foodstuffs using the 4100 MP-AES

Application note

Foods and beverages

Author

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Introduction

Whether the goal is food safety, ensuring quality or establishing provenance, measuring the trace element content of foods and beverages that we all consume is of paramount importance. While some elements are essential for our well being at low concentrations, others like lead and chromium are highly toxic and more still are being linked to viral, neurological and other diseases. Food scares related to contamination or poor quality not only constitute a health risk, they also undermine consumer confidence. This can lead to lost earnings through reduced sales and loss of credibility through adverse publicity.

Atomic spectroscopy is well established for the analysis of metals in foods and the technique employed often depends on the requirements of the application in terms of elements of interest, expected concentrations, and number and type of samples. Other important procurement factors that influence instrument selection include purchase and operational budget for consumables, gases, power and labor, as well as service and maintenance costs.

Agilent Technologies

With lab budgets coming under increasing pressure, Agilent has expanded its atomic spectroscopy portfolio to include the 4100 Microwave Plasma-Atomic Emission Spectrometer (MP-AES). MP-AES is a new analytical technique that uses a microwave-induced nitrogen plasma to provide elemental analysis, with significantly reduced running costs through the use of nitrogen as its plasma gas.

Experimental

This work describes the analysis of various certified and standard reference materials per the sample descriptions below:

- NIES CRM No.7 Tea Leaves: from National Institute of Environmental Studies (NIES), Japan.
- NIES CRM No.10c Rice Flour: from National Institute of Environmental Studies (NIES), Japan.
- NIST SRM 1577 Bovine Liver: from National Institute of Standards and Testing, USA.
- CRM-Wheat Flour: from High Purity Standards, USA
- CRM-Milk Powder: from High Purity Standards, USA
- CRM-Oyster Tissue: from High Purity Standards, USA

Sample preparation

A simple acid digestion method was used to prepare three of the samples. Initially, 0.25 g of the tea leaves CRM, 0.5 g of bovine liver SRM and 1 g of rice flour CRM were weighed into separate 250 mL beakers. This was then followed by the addition of 10 mL of HNO₃ and each beaker was covered with a watch glass. The samples were heated on a hot plate until completely dissolved. After cooling to room temperature, each digest was transferred to a 100 mL volumetric flask and made up to the required volume by adding Milli-Q water.

Pre-prepared sample solutions of CRM-Wheat Flour, CRM-Milk Powder and CRM-Oyster Tissue in 4% HNO₃ were purchased from High Purity Standards, USA.

Working standards and a blank were matrix-matched with the samples.

Instrumentation

The innovative 4100 MP-AES with its proprietary Microwave Excitation Assembly is a sequential atomic emission spectroscopic technique capable of fast, unattended multi-element analysis at varying concentration levels using a nitrogen plasma. The unique Microwave Excitation Assembly focuses and contains the microwave energy that is created via a concentrated axial magnetic field around the torch. This creates a robust toroidal plasma that allows the stable introduction of liquid samples. With a central channel temperature of ~5,000 K, MP-AES is highly suited to spectroscopic analysis, as it creates high intensity atomization emission lines. In addition to simplified spectra, nitrogen-MP-AES offers reduced operating costs and increased lab safety compared to flame AA, through the avoidance of costly and highly flammable gases such as acetylene.

The analysis was carried out using an Agilent 4100 MP-AES equipped with a standard MP-AES torch, concentric nebulizer, and glass cyclonic spray chamber.

Operating parameters are shown in Table 1.

 Table 1. Agilent 4100 MP-AES operating parameters

| Instrument parameter | Setting |
|-----------------------|--------------------|
| Nebulizer pressure | 160–180 kPa |
| Read time | 3 s (10 s for MDL) |
| Number of replicates | 3 (10 for MDL) |
| Stabilization time | 15 s |
| Background correction | Auto |

Results

Method detection limits

The Method Detection Limits were determined from the analysis of digested blank samples. The selected analytical wavelengths and method detection limits (3σ) are listed in Table 2.

| Table 2. Agilent 4100 MP-AES element wavelength and method detectio | n |
|---|---|
| imits (ppb) | |

| Element | Wavelength (nm) | MDL (ppb) |
|---------|-----------------|-----------|
| AI | 396.152 | 0.5 |
| Ва | 455.403 | 0.02 |
| Са | 445.478 | 14 |
| Cd | 228.802 | 1.2 |
| Со | 340.511 | 4 |
| Cr | 425.433 | 0.5 |
| Cu | 327.396 | 0.4 |
| Fe | 371.993 | 3 |
| К | 769.897 | 3 |
| К | 404.414 | 280 |
| Р | 213.618 | 100 |
| Pb | 405.781 | 5 |
| Pb | 368.343 | 12 |
| Mg | 518.361 | 4 |
| Mn | 403.076 | 0.5 |
| Мо | 379.825 | 1.5 |
| Na | 589.592 | 3 |
| Na | 568.821 | 140 |
| Ni | 341.476 | 2 |
| Ni | 352.453 | 2 |
| Sr | 407.771 | 0.01 |
| Zn | 213.857 | 4 |

Analysis of foodstuffs

Results of the analysis of major, minor and trace extractable elements in six different foodstuffs are listed in Tables 3 to 8. The measured values (carried out in triplicate) are in good agreement with the certified values for all CRM and SRM samples. Table 3. Results of NIES No.7 Tea Leaves

| Element | Measured values | Certified values |
|---------|-----------------|------------------|
| | wt% | wt% |
| Са | 0.314 ± 0.013 | 0.320 ± 0.012 |
| Mg | 0.150 ± 0.004 | 0.153 ± 0.006 |
| К | 1.861 ± 0.074 | 1.86 ± 0.07 |
| | mg/kg | mg/kg |
| Ва | 5.76 ± 0.57 | 5.7* |
| Cd | nd | 0.03 ± 0.03 |
| Со | nd | 0.12* |
| Cr | nd | 0.15* |
| Cu | 7.13 ± 0.81 | 7 ± 0.3 |
| Pb | nd | 0.8 ± 0.03 |
| Ni | 6.03 ± 0.63 | 6.5 ± 0.3 |
| Sr | 3.63 ± 0.43 | 3.7* |
| Zn | 34 ± 3 | 33 ± 3 |

* Reference values only

Table 4. Results of NIES No.10c Rice Flour

| Element | Measured values | Certified values |
|---------|-----------------|-------------------------|
| | wt% | wt% |
| Mg | 0.127 ± 0.006 | 0.125 ± 0.008 |
| К | 0.279 ± 0.012 | 0.275 ± 0.010 |
| Р | 0.300 ± 0.010 | 0.335 ± 0.008 |
| | mg/kg | mg/kg |
| AI | 1.49 ± 0.13 | 1.5* |
| Са | 95.4 ± 7.0 | 95 ± 2 |
| Cd | 1.83 ± 0.14 | 1.82 ± 0.06 |
| Со | nd | 0.007* |
| Cr | nd | 0.08* |
| Cu | 4.03 ± 0.32 | 4.1 ± 0.3 |
| Fe | 106 ± 0.15 | 11.4 ± 0.8 |
| Мо | nd | 1.6 ± 0.1 |
| Ni | nd | 0.30 ± 0.03 |
| Sr | 0.2 | 0.2* |
| Zn | 21.8 ± 1.0 | 23.1 ± 0.8 |

* Reference values only

Table 5. Results of NIST 1577 Bovine Liver

| Element | Measured values | Certified values |
|---------|-----------------|------------------|
| | wt% | wt% |
| Na | 0.247 ± 0.006 | 0.243 ± 0.013 |
| К | 1.00 ± 0.08 | 0.97 ± 0.06 |
| | mg/kg | mg/kg |
| Са | 131 | 123* |
| Cd | nd | 0.27 ± 0.04 |
| Co | nd | 0.18* |
| Cu | 185 ± 6 | 193 ± 10 |
| Fe | 266 ± 5 | 270 ± 20 |
| Pb | nd | 0.34 ± 0.08 |
| Mg | 625 ± 45 | 605* |
| Mn | 10.4 ± 1.41 | 10.3 ± 1 |
| Мо | nd | 3.2* |
| Sr | 0.15 ± 0.07 | 0.14* |
| Zn | 125 ± 4 | 130 ± 10 |

Table 7. Results of CRM-Milk Powder

| Element | Measured values (mg/kg) | Certified values (mg/kg) |
|---------|-------------------------|--------------------------|
| AI | nd | 0.020 ± 0.002 |
| Са | 131 ± 9 | 130 ± 1 |
| Со | nd | 0.0004* |
| Cr | nd | 0.0003* |
| Cu | 0.006 ± 0.001 | 0.007 ± 0.001 |
| Fe | 0.018 ± 0.002 | 0.020 ± 0.001 |
| К | 178 ± 6 | 170 ± 2 |
| Р | 98.7 ± 1.3 | 100 ± 1 |
| Pb | nd | 0.002* |
| Mg | 11.9 ± 0.2 | 12 ± 0.1 |
| Mn | 0.003 ± 0.002 | 0.003* |
| Na | 48.7 ± 2.6 | 50 ± 1 |
| Zn | 0.48 ± 0.05 | 0.50 ± 0.01 |

* Reference values only

Table 8. Results of CRM-Oyster Tissue

* Reference values only

Table 6. Results of CRM-Wheat Flour

| Element | Measured values (mg/kg) | Certified values (mg/kg) |
|---------|-------------------------|--------------------------|
| AI | 0.83 ± 0.02 | 0.85 ± 0.01 |
| Са | 9.64 ± 0.97 | 9.5 ± 0.1 |
| Cd | nd | 0.0015* |
| Со | nd | 0.001* |
| Cr | 0.013 ± 0.001 | 0.014* |
| Cu | 0.09 ± 0.008 | 0.1 ± 0.002 |
| Fe | 0.81 ± 0.04 | 0.90 ± 0.01 |
| К | 62.5 ± 0.5 | 65 ± 0.7 |
| Р | 61.1 ± 1.7 | 65 ± 0.7 |
| Pb | 0.05 ± 0.001 | 0.050 ± 0.003 |
| Mg | 20.8 ± 0.1 | 20.0 ± 0.2 |
| Mn | 0.36 ± 0.02 | 0.4 ± 0.008 |
| Ni | nd | 0.009 ± 0.001 |
| Zn | 0.47 ± 0.05 | 0.50 ± 0.01 |

| Element | Measured values (mg/kg) | Certified values (mg/kg) |
|---------|-------------------------|--------------------------|
| AI | 2.92 ± 0.07 | 3* |
| Са | 15.0 ± 0.49 | 15* |
| Cd | nd | 0.03* |
| Со | nd | 0.004* |
| Cr | nd | 0.007* |
| Cu | 0.56 ± 0.05 | 0.6* |
| К | 100 ± 0.96 | 100* |
| Р | 79.1 ± 0.9 | 80* |
| Pb | nd | 0.005* |
| Mg | 12.1 ± 0.2 | 12* |
| Mn | 0.18 ± 0.01 | 0.2* |
| Na | 48.9 ± 0.8 | 50* |
| Ni | nd | 0.01* |
| Zn | 8.3 ± 0.4 | 9* |

* Reference values only

* Reference values only

Conclusions

MP-AES offers any food testing facilities dependant on acetylene-based instrumentation a real alternative in terms of sensitivity, multi-element capability and speed of analysis, while cutting operating costs and improving the safety of the lab environment through the use of non-flammable nitrogen.

This study shows that following a quick and simple acid digestion sample preparation procedure (required for three of the six diverse food samples), all six certified and standard reference materials can be analyzed for trace and major element concentrations with good accuracy by MP-AES. The addition of the Agilent 4107 Nitrogen Generator is also possible in order to perform this analysis with significantly lower gas costs or for analysis in remote locations where sourcing of gases is costly or difficult.

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Total metals analysis of digested plant tissue using an Agilent 4200 Microwave Plasma-AES

Application note

Agriculture

Introduction

Plant growth and development largely depends on the composition and concentration of mineral nutrients available in the plant leaves and other tissues. These essential nutrients are divided into macronutrients (required in larger quantities because of their structural roles in the plant) and micronutrients (required in smaller quantities because they tend to be involved in regulatory roles in the plant). A deficiency or enrichment of nutrients may result in decreased plant productivity, crop yield or plant quality.

Analysis of the total metal content in plants is often carried out by Flame Atomic Absorption Spectrometry (FAAS) or Inductively Coupled Plasma Optical Emission Spectroscopy (ICP-OES). More recently, agricultural testing labs looking to upgrade or replace their FAAS with a more powerful technique are looking to Microwave Plasma—Atomic Emission Spectrometry (MP-AES) with its many advantages. MP-AES is a multi-element technique that offers better detection limits over a wider working analytical range than FAAS and more elements are available for analysis by MP-AES, including

Agilent Technologies

phosphorus, an expensive and widely used major nutrient in soil fertilization.

For laboratories that have difficulty in sourcing gases, are looking to reduce operating costs or are under pressure to improve safety by removing flammable gases, the MP-AES is ideal as it uses nitrogen gas, that can be generated from air.

This application note describes the sample preparation procedure and analytical method used to determine Cu, Fe, Mn, Zn, Na, K, Ca, Mg, B and P in a plant reference material using the Agilent 4200 MP-AES.

Experimental

Instrumentation

All measurements were performed using an Agilent 4200 MP-AES with nitrogen plasma gas supplied via an Agilent 4107 Nitrogen Generator. The generator alleviates the need and expense of sourcing analytical grade gases. The sample introduction system comprised a double-pass cyclonic spray chamber and the OneNeb nebulizer.

An Agilent SPS 3 autosampler was used to deliver samples to the instrument, allowing the system to be operated unattended. The instrument operated in a fast sequential mode and featured a Peltier-cooled CCD detector. Background and spectral interferences could be simutaneously corrected easily and accurately using Agilent's MP Expert software. Method parameters are given in Table 1.

Table 1. MP-AES method parameters

| Parameter | Value |
|-----------------------------------|---------------|
| Replicates | 3 |
| Pump rate | 15 rpm |
| Sample uptake delay | 35 seconds |
| Rinse time | 30 seconds |
| Stabilization time | 15 seconds |
| Fast Pump during uptake and rinse | On (80 rpm) |
| Autosampler | Agilent SPS 3 |
| Sample pump tubing | Orange/green |
| Waste pump tubing | Blue/blue |

Samples

Botanical reference material (RM) ASPAC 80 Pasture was obtained from the Australasian Soil and Plant Analysis Council (ASPAC, Carapook, VIC, Australia).

Sample preparation

Microwave digestion was used to prepare the ASPAC 80 RM for total metals analysis of Cu, Fe, Mn, Zn, Na, K, Ca, Mg, B and P by MP-AES. 7 mL of HNO_3 and 1 mL H_2O_2 was added to 0.18 g of the sample. A preloaded method for the MARS (CEM, Corporation, USA) microwave was used to digest the sample. Once cooled, the solution was diluted to 50 mL using ultrapure water. No further sample preparation was required and no modifiers or ionization buffers were added.

Wavelength selection and calibration range

Details of wavelength selection and calibration range are given in Table 2. Continuous wavelength coverage allows lines to be chosen that have appropriate sensitivity for the concentration range, and avoid spectral interferences.

Table 2. Wavelength and working calibration concentration range

| Element and wavelength (nm) | Calibration range (ppm) |
|-----------------------------|-------------------------|
| Cu 324.754 | 1–5 |
| Fe 259.940 | 5–25 |
| Mn 257.610 | 5–25 |
| Zn 213.857 | 1–5 |
| Na 568.820 | 2–100 |
| K 766.491 | 1–100 |
| Ca 445.478 | 20–100 |
| Mg 383.829 | 1–100 |
| B 249.772 | 0.25–1.0 |
| P 213.618 | 10–80 |

Results and discussion

Calibration

A typical calibration curve for phosphorus is displayed in Figure 1. The curve shows excellent linearity across the concentration range. The large linear dynamic range means that less sample dilutions are needed which improves productivity and reduces the risk of sample contamination.

Figure 1. Calibration curve for phosphorus

Sample analysis

The plant RM sample was analyzed for all elements in a single measurement. The quality of the MP-AES results was evaluated by comparing them with the reference values for ASPAC 80. Table 3 shows good accuracy was achieved for all elements over a wide concentration range. The ability of the MP-AES to determine all elements in a single sample measurement greatly simplifies the workflow when compared to an FAAS, and eliminates the need for lamp changes, measurements in absorption and emission, and in the case of B and P, analysis of the samples by other techniques.

 $\label{eq:table_state} \textbf{Table 3.} \ \text{MP-AES results for total metals content of plant reference} \\ \text{material ASPAC 80.} \\$

| Element | Wavelength nm | Measured value µg∕g | Reference value µg⁄g | Accuracy % |
|---------|------------------|---------------------------|----------------------------|---------------|
| Cu | 324.754 | 13.6 | 14.7 ± 1.2 | 93 |
| Fe | 259.940 | 316.13 | 324 ± 32 | 98 |
| Mn | 257.610 | 125.8 | 138 ± 10 | 91 |
| Zn | 213.857 | 54.6 | 58.1 ± 5.3 | 94 |
| Na | 568.263 | 2512 | 2460 ± 210 | 102 |
| K | 766.491 | 27302 | 26700 ± 1850 | 102 |
| Са | 445.478 | 10563 | 11100 ± 600 | 95 |
| Mg | 383.829 | 3239 | 3350 ± 220 | 97 |
| В | 249.772 | 21.65 | 23.7 ± 3.4 | 91 |
| Р | 213.618 | 3223.35 | 2970 ± 250 | 109 |

Conclusions

The study shows the effectiveness of the Agilent 4200 MP-AES for the analysis of total metal content of a plant-based reference material following microwave digestion. Elements that are difficult to analyze by FAAS such as B and P were included, with all data acquired in a single run. Accurate determinations over a wide concentration range were obtained showing the suitability of MP-AES for the application. When compared to an FAAS, the workflow on the MP-AES is also simplified by eliminating the need for multiple sample preparations, lamp changes and measurements in absorption and emission modes.

Current trends in the market for lower detection limits, lower cost of analysis, improved ease of use and improved safety, are all met by the Agilent 4200 MP-AES. The instrument uses nitrogen, eliminating expensive and hazardous gases such as acetylene, increasing safety, and allowing for unattended operation of the instrument, even in remote locations. When the N₂ is supplied using the Agilent 4107 Nitrogen Generator that extracts N₂ from air, running costs are greatly reduced compared to an FAAS or ICP-OES that rely on a constant supply of analytical grade gases.

With greater sensitivity, linear dynamic range, and sample throughput compared to FAAS, the Agilent 4200 MP-AES is the ideal replacement for labs looking to extend their analytical capabilities.

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Determination of exchangeable cations in soil extracts using the Agilent 4100 Microwave Plasma-Atomic Emission Spectrometer

Application note

Agriculture

Introduction

Accurate, routine testing of nutrients in soil samples is critical to understanding its potential fertility. Many of the nutrients that are vital to plants are exchangeable cations. These are ions loosely attached to and/or adsorbed onto clay particles and organic matter in soil that may become available to plants. Determination of these cations is of great interest for agronomic diagnostic and soil sustainability, enabling more accurate assessment and management of nutrient requirements [1, 2]. If the results indicate there is a nutrient imbalance, then this can be corrected for by the application of a suitably formulated fertilizer.

This application note describes an analytical method for the determination of Ca, K, Mg, Mn and Na in soils using the Agilent 4100 Microwave Plasma-Atomic Emission Spectrometer (MP-AES). A chemical extraction with 1 M ammonium acetate is recommended (standard NF X 31-108) [1, 2, 3]. In this work, results obtained with MP-AES are compared to those obtained by other wellproven, validated techniques flame atomic absorption spectrometry (FAAS) and inductively coupled plasmaoptical emission spectrometry (ICP-OES), and with inter-laboratory results to demonstrate the reliability and accuracy of MP-AES data.

Which measurement technique is right for you?

There are many factors to be taken into account when selecting the right analytical technique. In many cases several techniques will provide adequate detection range, so the technique of choice will depend on factors such as sample throughput requirements, ease-of-use, infrastructure required, and on-going operating costs. In the case of this application, it has been more common for smaller laboratories with low sample throughput requirements to use FAAS, while some larger laboratories (with higher sample throughput requirements) may use ICP-OES.

The 4100 MP-AES fits between FAAS and ICP-OES in many aspects such as detection power, dynamic range, and speed of analysis. For these key performance metrics, the MP-AES offers a unique alternative to both FAAS and ICP-OES.

There are also some clear differentiating benefits of the MP-AES technology over these more traditional options. By eliminating the need for on-going gas resupply, the MP-AES offers significantly reduced on-going operating costs over both FAAS and ICP-OES — and avoids flammable gases (required for FAAS), hence enhancing safety and allowing unattended, overnight operation. The reduced infrastructure required for MP-AES also makes it well-suited to remote sites where supply of gases can be difficult and/or expensive. These features make the MP-AES an attractive technique for many small to medium size agricultural laboratories, particularly those at remote locations, and for an

increasing number of laboratories requiring the lowest possible on-going operating costs.

Experimental

Instrumentation

The 4100 MP-AES revolutionizes the way analysts conduct multi-elemental analysis. Using a microwave plasma that is based on nitrogen, supplied from a compressed air supply and the Agilent 4107 Nitrogen Generator, the 4100 MP-AES does not require flammable or expensive gases such as acetylene, nitrous oxide or argon. This improves lab safety, results in a significant reduction in operating costs and allows installation in mobile labs or remote locations where gas supplies may not be available.

Additionally, the 4100 MP-AES has been designed to improve the analytical performance and productivity when compared with FAAS, with good sensitivity and detection limits down to sub ppb levels over a wide linear range.

Instrument operating conditions are listed in Table 1.

Table 1. Agilent 4100 MP-AES operating conditions

| Instrument parameter | Setting |
|--------------------------------|----------------------------|
| Nebulizer | OneNeb |
| Spray chamber | Glass cyclonic single-pass |
| Sample tubing | White-white |
| CsCl tubing | Orange-yellow |
| Waste tubing | Blue-blue |
| Read time | 3 s |
| Number of replicates | 3 |
| Stabilization time | 15 s |
| Fast pump during sample uptake | Yes |
| Pump speed | 15 rpm |

The analysis of soil samples was also carried out by ICP-OES and FAAS. Conditions of analysis are as described in Table 2. Table 2. Global conditions of analysis

| Instrument | Dilution | Comments |
|---------------------|---|--|
| Agilent 4100 MP-AES | On-line with CsCl 1.5% | OneNeb nebulizer |
| Agilent 280 FS AAS | SIPS 20. Dilution factor set by element | CsCl as modifier for Na and K, La/CsCl as modifier for Mg |
| Agilent 725 ICP-OES | No dilution | OneNeb nebulizer |

Material

Soil samples were provided (air-dried and sieved <2 mm) by the French inter-laboratory comparisons organization BIPEA (Inter-professional Bureau of Study and Analysis, France). All samples have been recently analyzed in proficiency testing, so that the reference values and standard deviations for CaO, K₂O, MgO and Na₂O concentrations in ammonium acetate were known. Note: manganese was not included in the testing scheme but has been included in this investigation.

The reference numbers of the soil samples used were 403, 418, 421 and 423.

Sample preparation

As described in the standard NF X 31-108, 1 M ammonium acetate adjusted to pH 7, was used as extractant. The total concentration of dissolved salt was 77 g/L.

For each soil, 50 mL of 1 M ammonium acetate was added to 2.5 g of soil. The mixture was shaken by rotation in a room at 20 °C ±2 °C over 1 hour. After extraction, the samples were filtered using a filter paper and the clear solutions were analyzed immediately.

Standard preparation

Four multi-element standard solutions were prepared in 1 M ammonium acetate. Table 3 provides details of the calibration concentration range for each analyte. Single element stock solutions from Merck Germany were used. Table 4 displays selected wavelengths and calibration parameters used for analysis.

Cesium chloride was used as an ionization buffer. This was added on-line via a 'Y' piece to avoid manual spiking of standards and samples. Table 3. Calibration standards used for soil extraction analysis (mg/L)

| | Ca | К | Mg | Mn | Na |
|------------|-----|----|----|-----|-----|
| Blank | 0 | 0 | 0 | 0 | 0 |
| Standard 1 | 100 | 5 | 5 | 1.0 | 0.5 |
| Standard 2 | 200 | 10 | 10 | 2.0 | 1.0 |
| Standard 3 | 300 | 15 | 15 | 3.0 | 2.0 |
| Standard 4 | 600 | 30 | 30 | 6.0 | 4.0 |

 Table 4. Agilent 4100 MP-AES wavelengths and calibration parameters selected for analysis

| Element | Wavelength (nm) | Read time (s) | Nebulizer pressure (kPa) | Background correction |
|---------|--------------------|------------------|-----------------------------|-----------------------|
| Са | 430.253 | 3 | 240 | Auto |
| К | 769.897 | 3 | 240 | Auto |
| Mg | 383.829 | 3 | 240 | Auto |
| Mn | 403.076 | 3 | 240 | Auto |
| Na | 588.995 | 3 | 240 | Auto |

Results

Calibration

The calibration curves for Ca, K, Mg, Mn and Na on the MP-AES are displayed in Figure 1. Results show good linearity. This highlights the better linear dynamic range achieved with the 4100 MP-AES as compared to FAAS. Less sample dilutions are then needed when using MP-AES avoiding sample contamination and enhancing productivity.

Sample analysis

The accuracy of the results obtained by MP-AES was evaluated by two methods: i) comparison of the MP-AES results with results obtained with another analytical technique (FAAS and/or ICP-OES) and ii) calculation of z-scores for the MP-AES results with data from the inter-laboratory test (reference results and corresponding standard deviations).

Figure 1. Typical MP-AES calibration curves for Ca, Mn, K, Na and Mg

Mn (403.076 nm) Calibration

Comparison of MP-AES with ICP-OES and/or FAAS Figures 2 to 6 show for all cations an excellent agreement between the 4100 MP-AES results and those from other spectrometric techniques. The correlation of the analytical results between the different techniques was made and linear regressions are observed: coefficients of determination are $0.995 < R^2 < 0.999$ and the slopes are between 0.969 and 1.043 (Table 5).

Figure 2. Reference results and Ca concentrations measured in soil extracts by MP-AES and ICP-OES (FAAS not determined)

Figure 3. Reference results and K concentrations measured in soil extracts by MP-AES, ICP-OES and FAAS

Figure 4. Reference results and Mg concentrations measured in soil extracts by MP-AES, ICP-OES and FAAS

Figure 5. Mn concentrations measured in soil extracts by MP-AES and ICP-OES (FAAS not determined and no reference results available)

Figure 6. Reference results and Na concentrations measured in soil extracts by MP-AES, ICP-OES and FAAS

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