

Measurement of Chromium in Environmental Waters by Zeeman Corrected Graphite Tube Atomization

Application Note

Atomic Absorption

Introduction

The measurement of trace metal levels in environmental waters is ideally performed by atomic absorption spectroscopy, and the literature on the techniques used is very extensive [1–7]. Chromium, along with other elements such as cobalt, copper, lead, manganese and nickel, requires monitoring in the environment.

The levels encountered are often too low to be measured by direct aspiration into the flame of an atomic absorption spectrometer, thus two methods are commonly employed to overcome this problem. The first is to form a metal complex using various ligands (pentan-2, 4-dione, acac; ammonium 1-pyrrolidinecarbodithioic acid, APDC) and extracting the complex into an organic solvent (4-methylpentan-2-one, MIBK; 2,5-methylheptan-4-one, DISK) [2]. By careful selection of the relative volumes, the analyte can be preconcentrated before aspiration into a flame. This method is relatively inexpensive but tedious and prone to contamination. Solvent extraction can also be used for furnace work [1–6].

The other method is by direct injection of the water sample into a graphite furnace [1,3–7]. Sample preparation and the possibility of contamination are therefore minimized [4]. The use of a background corrector is strongly recommended so that the effect of background interferences is reduced [7].



Author

Jonathan H. Moffett

Practical

Water samples collected under Environmental Protection Agency recommended conditions were used for method development. The method was then checked by measuring the amount of chromium in certified water samples. The first water sample was supplied by the US Environmental Protection Agency [8]. The other sample was a standard reference material obtained from the National Bureau of Standards [9].

Experimental

An Agilent SpectrAA-30 Zeeman atomic absorption spectrometer with a programmable sample dispenser and a chromium hollow cathode lamp-were used for this study. An Epson MX-80 printer was used for all printouts and signal graphics traces. A partitioned graphite tube was used in the furnace. The instrument parameters used for the final studies are reproduced in Table 1.

The chromium standard used in this study was prepared from BDH Spectrosol stock solution of 1000 mg Cr/L in 1 M nitric acid and diluted using deionized distilled water to give a final working concentration of 20 μ g Cr/L.

workin	g concentratio	n of 20 µ	g Cr/L.		
Table 1.	Instrument Para	ameters			
	PROGRAM 7 Cr	EFFLUEN	WATER		
	CALIBRA MEASURE LAMP PC BLIT WI SLIT WI WAVELEN SANPLE TIME CC MEASURE REPLICA BACKGRC	DBITION JRRENT (mm) EIGHT (nm) INTRODUC INTRODUC INTRODUC INTRODUC INTRODUC INTES JUND CORR	E E TION E (sec)	7 0.2 REDUCED 337.9 SAMPLER AUTOMIX 0.05 2.0 2 2 ON	ING
STEP NO.	TEMPERATURE		GAS FLO (L/min)	•	RI
1 2		5.0	3.0	NORMAL	N
3	120	10.0	3.0	NORMAL	N
4	1100	20.0	3.0		N
5	1100		3.0		N
6			0.0		
7				NORMAL	Y
8				NORMAL	Y
4	2500	2.0	3.0	NORMAL	N

		SAMPLER PARAMETERS VOLUMES (uL)							
		SOLUT	ION	1	BLANK		MOI	DIFIER	
BLANK	. حتير پيرو ڪند سيم جينة گند ه			~	15				
STANDARD	1	2			13				
STANDARD	2	5			10				
STANDARD	3	10			5				
SAMPLE		10			5				
		RECALIB	RATIO	N RATE		0			
		RESLOPE	RATE			0			
MULTIPLE	INJECT	ND	нот	INJECT		NO	PRE	INJECT	NO

Results and Discussions

The collection of the environmental water samples and the preparation of all the standards meant that the chromium ions were present in a dilute nitric acid matrix. This particular matrix stabilizes the analyte in solution and allows the use of an ashing temperature which is high enough to remove all organic material. Signal graphics traces of a standard and a sample are reproduced together in Figure 1. Close examination reveals that there is very little difference in the respective background signals.

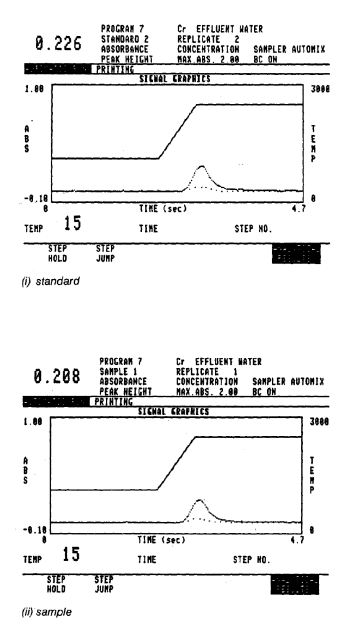


Figure 1. Signal graphics traces.

READ

NO NO NO NO YES YES

NΩ

COMMAND

A study was performed to see if there are any matrix effects. Because all organic matter is apparently destroyed, other metal ions present in the matrix could be a potential source of interferences. A standard additions calibration slope was compared to a normal concentration calibration slope. The data points and slopes are shown in Figure 2. As can be seen, the slopes are virtually identical. The concentration calibration mode was used for convenience in subsequent studies. The validity of the method was checked by analyzing samples of known concentration. A concentration calibration curve was determined followed by analysis of samples of USEPA sample 4 and NBS SRM 1643b respectively. The autorun report is reproduced in Figure 3. The solutions being dilute aqueous solutions, their density would be the same as pure water to a very good approximation. Hence the numerical value for ng/L would be equal to the value in ng/g. The results obtained are compared in Table 2, and are seen to be in very good agreement.

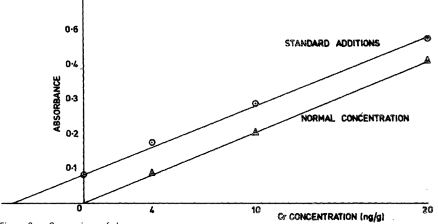


Figure 2. Comparison of slopes.

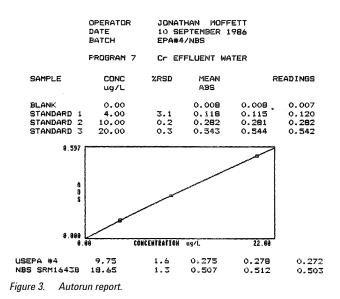


Table 2. Results Comparison

Comparison of experimentally determined results with quoted results					
Standard		Quoted (ng/g)	Found (ng/g)		
US EPA	Sample 4	10.2 ± 1.1	9.75 ± 0.16		
NBS SRM	1643b	18.6 ± 0.4	18.65 ± 0.24		

Conclusion

The determination of trace levels of chromium in waters and effluents can be done quickly and accurately and with good precision using a graphite furnace with an effective Zeeman background corrector.

References

- 1. Trevor McKenzie, Varian Instruments At Work No AA-8, October 1980.
- 2. Lucinda Voth, Varian Instruments At Work No AA-17, August 1981.
- 3. Douglas Shrader, Lucinda Voth, Lawrence Covick, Varian Instruments At Work No AA-30, June 1983.
- 4. Douglas Shrader, Lucinda Voth, Lawrence Covick, Varian Instruments At Work No AA-31, June 1983.
- Douglas Shrader, Lucinda Voth, Lawrence Covick, Varian 5. Instruments At Work No AA-33, September 1983.
- 6. Lucinda Voth, Varian Instruments At Work No AA-34, September 1983.
- 7. Lucinda Voth, Varian Instruments At Work No AA-35, September 1983.
- 8. Environmental Monitoring and Support Laboratory, US Environmental Protection Agency., Cincinnati, USA.
- 9. National Bureau of Standards, US Department of Commerce, Washington DC, USA.

For More Information

For more information on our products and services, visit our Web site at www.agilent.com/chem

www.agilent.com/chem

Agilent shall not be liable for errors contained herein or for incidental or consequential damages in connection with the furnishing, performance, or use of this material.

Information, descriptions, and specifications in this publication are subject to change without notice.

© Agilent Technologies, Inc., 1987 Printed in the USA November 1, 2010 AA069



Agilent Technologies