

Determination of chromium content in pyrotechnics used for fireworks and firecrackers based on inductively coupled plasma optical emission spectrometric approach (ICP-OES)

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Abstract. Inductively coupled plasma optical emission spectrometric approach(ICP-OES) is used to determine the chromium content in pyrotechnics used for fireworks and firecrackers. Element of chromium is commonly found as impurity in chemical materials used for pyrotechnics in fireworks and firecrackers. Statistical analysis shows that chromium content in pyrotechnics is below 5%.Concept of this method: considering the weight of the sample is 400mg,constant volume is 0.5L and the concentration of chromium is below 40mg/L in sample solution, the determination scope of the method for the chromium content would be between 5%.Further experiments proved that the fitting correlation coefficient of chromium calibration curve is 0.9995 or higher, recovery is 93.96%-100.49%.The allowable differential value is 0.4% between two single tests under repeatable conditions. This method can completely satisfy the requirements of the fireworks and firecrackers industry with high accuracy and good precision.

Introduction

In China, fireworks and firecrackers are very important consumer recreational products in people's everyday life since ancient times. Gorgeous colors produced by fireworks and firecrackers are even the leading role of foiling festal atmosphere in every grand holiday celebrations. In recent years, with the rapid development of global trade, fireworks and firecrackers are becoming more and more popular all over the world, more and more consumers are fascinated by different kinds of patterns, pictures, and sound effects of fireworks and firecrackers. Pyrotechnics used to produce various kinds of pyrotechnic effects in fireworks and firecrackers are consists of many different solid chemicals. Chromium element is commonly found in pyrotechnics as impurity in chemical materials used for pyrotechnics in fireworks and firecrackers, and which would seriously affect the environment. Quantitative analysis of chemical compositions in pyrotechnics such as chromium content is required under the Globally Harmonized System of Classification and Labeling of Chemicals(GHS)to be complemented in the fireworks and firecrackers industry. Meanwhile it will also provide a scientific and effective technical support to the management and supervision of safety production for the government, and improve products' quality level by the manufacturers. It can also be utilized as a tool in providing valuable data in the judgment in some major arbitration and security incident analysis. Quantitative analysis method of the chromium content reported in current literature is limited to traditional chemical analysis, such methods have the following disadvantages:(1) Long detecting period. Generally, it will take a skilled technician two whole days or so to complete the detection.(2) The operation is more complicated. It needs to go through many steps such as dissolving sample, filtration, precipitation collection, drying and weighing precipitation and ect. Comparing with traditional chemical analysis methods, this method based on inductively coupled plasma optical emission spectrometric approach has the advantages of simple operation steps, short period of detection, high accuracy and good precision.



Theory

Chromium element is commonly found as impurity in chemical materials such as potassium dichromate and potassium chromate in pyrotechnics in fireworks and firecrackers. Statistical analysis shows that potassium dichromate or potassium chromate in pyrotechnics is below 10%, which concludes that the chromium content in pyrotechnics would not be above 5% as mass fraction. Concept of the method: considering the weight of the sample is 400mg, constant volume is 0.5L and the concentrations of the chromium is below 40mg/L in sample solutions, the determination scope of the method for the chromium content is below 5%. In accordance with the relevant safety regulations, the sample was ground into powder of less than 180 micron. Then the sample powder is placed in an explosive–proof oven at 50°C–55°C and dried for 4 hours, and then placed into a dryer for cooling down to room temperature. Pretreated sample is fully dissolved in 100mL pure water and then filtered into volumetric flask as sample solution. The sample solution is injected into the plasma as a light source, and we can measure the spectral intensity corresponding to the element wavelength in the plasma spectrometer. The actual content of chromium element in the sample can be calculated from the concentrations of the chromium reading by the calibration curve.

Experiment section

Reagents

Unless otherwise stated, all the reagents should be guaranteed reagents and pure water is secondary grade water as described in ISO 3696(1987). Hydrochloric acid (1+19): mix 10 mL hydrochloric acid and 190 mL pure water thoroughly. Standard solution of the chromium: mass concentration is 1,000mg/L. Standard working solution of the chromium: separately pipette the standard solution of chromium with volume 0 mL,0.1mL, 0.5mL, 1mL, 2mL and 5mL into four 100mLvolumetric flasks, and add hydrochloric acid solution (1+19) to reach 100 mL in each volumetric flask, mix thoroughly for later use. Concentrations of the standard working solution in different flasks are shown in Table 1.

Table 1 Concentrations of series standard working solutions of chromium mg/L

element	concentrations of series standard working solutions							
	N1	N2	N3	N4	N5	N6		
Cr	0	1	5	10	20	50		

Instrument and apparatus

Explosive-proof oven with accuracy to $\pm 2^{\circ}$ C. Analytical balance with accuracy to 0.1 mg. Inductively coupled plasma atomic emission spectrometer (ICP-OES): United States Thermo Fisher (former Thermo Electron Corporation) Company iCAP6300 series.

Operation step

- (1) Weigh the sample of about 400mg, accuracy to 0.1 mg.
- (2) Place the sample into a clean 300 mL beaker, add 100mL pure water into the beaker, then place the beaker on an electric stove to make the solution boiled for about 20 min. Filter the solution through filter paper to an 0.5L volumetric flask, wash the beaker and the filter paper several times with hydrochloric acid (1+19), and make a constant volume with pure water after the filtered solution



cooling down to room temperature.

(3) Parameters of the ICP-OES instrument parameters are shown in Table 2.

Table 2Parameters of the ICP-OES instrument

parameters\element	Cr
Flush pump speed	100rpm
Analysis pump speed	75rpm
Pump stable time	5s
Tube Type	Polyethylene Orange / White
RF power	1150w
Auxiliary gas flow	0.5 L/min
Nebulizer gas flow	0.70 L/min
Vertical viewing height	12mm
Spectral line	Cr2835,Cr2843
Repeat times	3

- (4) Calibration (working) curve: according to the requirements of the method and the instrument criteria, we set the instrument to optimum analysis conditions, and adjust it to the best working condition, and determine spectral intensity of the series standard solution from N_1 to N_6 to establish the calibration (working) curve with the elemental concentrations as independent variable and the spectral intensity as the dependent variable. The linear correlation coefficient of the regression curve should be 0.995 or higher.
- (5) Sample determination: determine the spectral intensity of the chromium in blank solution and every sample solution under the best analysis condition and read the concentrations from the calibration curve according to the spectral intensity.

Results calculation

Content of the chromium element in the sample can be calculated as mass fraction W and its value shown in% according to the following formula.

$$W = \frac{cV}{m} \times 100\%$$

Where: c— concentration of the chromium element in the sample solution read by ICP-OES in mg/L; V—constant volume of the volumetric flask used for the sample solution, expressed in liters(L); m—quantity of the sample, expressed in milligrams (mg).

W— the content of the chromium in the sample, expressed in %.

Experimental data and analysis

Recovery test

To assess the accuracy of the method, we used the standard substances addition recovery test. We added the chromium standard substance to some actual pyrotechnic samples, dissolved the samples and determined the contents of chromium in the sample solution. The values of the chromium contents we measured are compared with the theoretical ones, and the experimental data is shown in Table 3 below.



Table 3 Recovery test results

Elements line		Measured concentrati ons of the sample solution (mg/L)	quality contents of	Measured quality contents of the sample (%)		Recovery (%)	Concentration range of the calibration curve (mg/L)
	1.82	1.71	0.11	0.11	0.01	93.96	
	6.84	6.65	0.43	0.42	0.01	97.22	
Cr2835	13.64	12.99	0.85	0.81	0.04	95.21	1–50
Cr2855	26.07	26.20	1.63	1.64	0.01	100.49	1–30
	38.50	37.95	2.41	2.37	0.03	98.56	
	42.15	41.60	2.63	2.60	0.03	98.70	

Precision test

The precision of the method is expressed in the allowable differential value between two single tests under repeatable conditions. The content of the chromium in actual pyrotechnics is below 5%. Three samples of different chromium contents to be determined under the same condition over a period of time(almost two months). Each sample was repeatedly measured in 4 rounds with 11 measurements in each round. We calculated the allowable differential values under repeatable conditions, and the results are listed in the Table 4 below.

Table 4 Precision test results

No.	rounds			Value	es (%))		Standard deviation of each round Si (%)	Standard deviation of the repeatable measurements Sr (%)	Allowa ble differen tial values Ir (%)
	1	1.78	1.64	1.83	1.86	1.85	1.68	0.1087		0.3189
		1.79	1.86	1.81	1.52	1.82		0.1067		
	2	1.75	1.84	1.83	1.86	1.85	1.89	0.1067		
01	2	1.84	1.72	1.81	1.52	1.89		0.1007	0.1121	
01	3	1.81	1.84	1.83	1.78	1.55	1.89	0.1167	0.1121	
	3	1.84	1.75	1.54	1.82	1.82		0.1107		
	4	1.86	1.84	1.53	1.73	1.85	1.89	0.1140		
		1.84	1.73	1.61	1.82	1.82		0.1140		
	1	3.23	3.36	3.21	3.24	3.51	3.34	0.1081	0.1106	0.3131
		3.27	3.24	3.19	3.11	3.36				
	2	3.06	3.36	3.15	3.24	3.17	3.18			
02		3.27	3.36	3.01	3.11	3.31		0.1172		
02	3	3.16	3.12	3.13	3.24	3.31	3.12	0.1013		
		3.12	3.26	3.42	3.14	3.11		0.1013		
	4	3.31	3.27	3.31	3.24	3.34	3.34	0.1150		
		3.43	3.26	3.01	3.31	3.11		0.1159		
	1	4.23	4.36	4.21	4.24	4.51	4.34	0.1047	0.1021	0.2932
		4.27	4.24	4.19	4.11	4.26				
03	2	4.06	4.36	4.15	4.24	4.17	4.18	0.1136		
		4.27	4.36	4.01	4.18	4.31				
	3	4.16	4.12	4.13	4.24	4.31	4.12	0.1005		



	4.12	4.26	4.42	4.24	4.11		
4	4.31	4.21	4.31	4.24	4.34	4.34	0.0005
4	4.43	4.26	4.11	4.21	4.11		0.0985

Explanation of the results calculation in table 4:

Standard deviation of each round (Si) refers to the standard deviation of a sample which has been measured parallel for n times under the same condition in the same time span and calculated as follow:

$$S_{i} = \sqrt{\frac{\sum_{i=1}^{n} (X_{i} - \overline{X})^{2}}{n-1}}$$

X_i— value of the i-th parallel measurement;

 \overline{X} — average value of n times parallel measurements;

n — number times of parallel measurements.

Standard deviation of the repeatable measurements Sr(%) refer to the standard deviation of a sample which has been measured parallel for m times under the same condition in different time. Different time interval refers to the period of time interval in which we can maintain the same determined conditions, so the precision of m rounds repeatable measurements are equivalent. Standard deviation(Sr) is calculated as follow:

$$S_r = \sqrt{\frac{\sum S_i^2}{m}}$$

S_i— standard deviation of the i-th round of n times parallel measurements.

M—number of times of parallel measurements.

The allowable differential value(Ir) refers to the possibility of the absolute difference exceeding Ir between any two results is only 5% (when the probability with 95% confidence) which are measured with the same sample by the same method in the same condition. The allowable differential value(Ir) is calculated as follow:

$$I_r = 2.83S_r$$

As we can see from Table 4,the allowable differential values of chromium in this method is 0.4%.

Conclusions

Inductively coupled plasma optical emission spectrometric approach(ICP-OES) is used to determine the chromium content in pyrotechnics used for fireworks and firecrackers, this method is accurate and quick with high accuracy and good precision. When the chromium content in the sample is below 5% as mass fraction, the recovery is 93.96%-100.49%. The allowable differential value was 0.4% between two single tests under repeatable conditions. In other word, this method can completely satisfy the requirements of the fireworks and firecrackers industry.

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