

L mol<sup>1</sup> cm<sup>-1</sup> and 0.0018-µg.cm<sup>-2</sup> respectively. The interference effect of various diverse ions has been studied to be observed shows 1:1 [Mo (VI): CMTSC] stoichiometry with stability constant 3.56x10<sup>6</sup>. The standard deviation of the method in the determination of 0.383-µg ml<sup>-1</sup> of Molybdenum (VI) was 0.001. First and second order derivative spectrophotometric methods are developed at  $\lambda$ max 420 nm and 440 nm respectively for the determination of Molybdenum (VI) which was more sensitive than the zero order method. The developed method has been used for the determination of Molybdenum (VI) in foodstuffs, pharmaceutical samples and in alloys.

**Introduction:** The potential analytical applications of hydrazone derivatives have been reviewed by Singh et al<sup>1</sup>. Hydrazones are important class of known analytical reagents. Derivative spectrophotometry was a very useful approach for determining the concentration of single component in mixtures with overlapping spectra as it may eliminate interferences. Hydrazones and its derivatives reacts with many metal ions forming colour complexes and act as chelating agents. A micellar solution has the ability to enhance the stability of metal complex and has been utilized as a medium for the spectrophotometric determination of the metal chelate <sup>2-4</sup>.

Molybdenum has been shown to be essential trace element in animal physiology<sup>5</sup>. Analytical methods have been described for the determination of molybdenum in plant material, but application to the low levels in a range of real foodstuffs for human consumption is rare.

The measurement of Molybdenum by flame atomic absorption spectrometry (AAS) after chelation and extraction has been described<sup>6</sup>, but this technique does not reach at the desired levels, while the use of graphite furnace atomic absorption spectrometry (GFAAS) was susceptible to matrix interferences<sup>,7,8</sup>. Molybdenum has been determined Spectrophotometrically by its catalytic effect on the liberation of iodine from iodide by hydrogen peroxide using an automated procedure<sup>9,10.</sup> Many Spectrophotometric methods for the determination of Molybdenum (VI) based on its reaction with thiocyanate in the presence of a reducing agent<sup>11</sup> and with toluene-3, 4-dithiol<sup>12</sup> are not too selective, as they are subject to interference from many other elements. Some of the recently proposed methods using thiosemicarbazones<sup>13</sup> are found to be less sensitive. The methods employing different organic reagents have also been reported<sup>14</sup>. Although, several methods have been reported for the Spectrophotometric determination of Molybdenum (VI) 15-18, they suffer from drawbacks such as lack of reproducibility, stability, interferences and requirements of prior extraction requirement of heating.

Experimental part: materials and methods: The absorbance and pH measurements were made on a Shimadzu UV-visible spectrophotometer (Model UV-160A) fitted with 1.0 cm Quartz cells and Elico digital pH meter (Model LI 120) respectively. Suitable settings for derivative were as follows. The spectral band length was 5 nm, the wavelength accuracy was 0.5 nm with automatic wavelength correction and the recorder was a computer controlled thermal graphic printer with a cathode ray tube and one degree of freedom in the wavelength range 300–800 nm.

Facile synthesis of CMTSC: The reagent (CMTSC) was prepared by the Sah and Daniels  $^{16}\mbox{ procedure}$ 

**Characterization of CMTSC:** The structure and characterization of CMTSC was confirmed by IR, NMR and Mass spectral data

**Results and discussion:** The chromogenic reagent CMTSC was used for the spectrophotometric determination of the Molybdenum (VI). The Molybdenum (VI)-CMTSC complex shows the maximum absorbance at 400 nm, where the reagent blank does not absorb appreciably.

The calibration graph follows the straight-line equation Y= a c + b; where c is the concentration of the solution, Y is measured absorbance or peak height and a and b are constants. By substituting the Mo (VI)-CMTSC complex experimental data in the Beer's law equation, the calibration equations are calculated as  $\lambda$ max 400 nm =0.5226X+0.00239 for zero order data and  $\lambda$ max 420 nm =0.43074X+9.74116x10<sup>-5</sup> for first derivative data and  $\lambda$ max 440 nm = 0.67626X+0.00421 for second order derivative data which gives the straight lines.

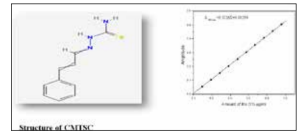


Fig: 1: Absorbance Vs Amount of Mo (VI) Mo (VI) =  $1.0x10^{-6}$  M CMTSC =  $2x10^{-3}$  M

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Triton-X100 (5%) = 0.5 ml pH =4.0 λmax =400 nm



	o (VI)-CMTSC- vative spectra	Fig.3: Mo (VI)-CMTSC-Sec- First derivative spectra			
Mo (VI)	= 1.0x10 <sup>-6</sup> M	Mo (VI) = $1.0 \times 10^{-6}$ M			
CMTSC	= 2x10 <sup>-3</sup> M	$CMTSC = 2x10^{-3} M$			
Triton-X100 = 0.5 ml		Triton-X100 = 0.5 ml			
рН	=4.0	pH =4.0			
λmax	= 420 nm	λmax = 440 nm			

Interference: In order to assess the analytical potential of the proposed methods, the effects of some diverse ions which often accompany Molybdenum (VI) was examined by carrying out the determination of 0.497 µg/ml of Molybdenum (VI) in the presence of foreign ions for CMTSC. The tolerance limit of a foreign ion was taken as the amount of foreign ion required to cause an error of  $\pm 2\%$  in the absorbance or amplitude. The data obtained in the derivative method was also incorporated. The data suggest that several associated anions and cations do not interfere when they are present in large excess, such as iodide, Nitrate, Thiosulphate, Thiocyanide, Bromide, Sodium (I), Bismuth (III), Tungsten (VI) and Zirconium (IV). The tolerance limit values for many anions and cations are more in derivative method. The interference of associated metal ions such as Iron (III) and Copper (II) is decreased by adding masking agents Phosphate and Thiourea respectively.

Applications: The method proposed in the present studies was applied for the determination of Molybdenum (VI) in Foodstuffs, Pharmaceutical samples and in Alloys.

Table:	Determina	ation of	f Molybdenur	n (VI) ir	n Foodstuffs,
Pharm	aceutical, A	Alloy sai	mples		

Foodstuff	Stand- ard Method	Amount of Molybdenum (VI) found* %					
		Zero order	R (%)	D1	R (%)	D2	R (%)
Banana	0.479	0.476	99.37	0.478	97.79	0.478	97.79
Apple	0.239	0.234	97.9	0.238	99.5	0.238	99.5
Cabbage	1.439	1.42	98.67	1.431	99.65	1.431	99.65
Tomato	0.959	0.954	99.47	0.955	99.5	0.955	99.5
Rice	0.718	0.705	98.18	0.711	99.0	0.711	99.0
Potato	0.240	0.238	99.16	0.239	99.58	0.239	99.58
Pharmaceutical sample							
Pantobionta	2.0	1.94	-3.0	1.96	-2.0	1.96	-2.0
Alloy samples							

BCS 406 Mn (0.53) Ni (1.69) Mo (1.03) Cr (2.12) Cr (2.12) Cu (0.32) V (0.02)	1.03	1.010	-1.92	1.020	-0.97	1.020	-0.97
JSS17 1-3							
Mo (0.035) Ti (0.036) As (0.045) Ni (0.011) Sn (0.034) Al (0.040) Ca (0.0013) Cr (0.067)	0.035	0.033	-5.71	0.034	-2.85	0.034	-2.85
Die Steel H 13 Grade Mo (1.254) Si (0.974) Mn (0.364) P (0.024) S (0.006) Cr (4.962) Ni (0.318) V (0.994)	1.254	1.250	-0.31	1.252	-0.15	1.252	-0.15

Conclusion: The proposed methods are simple, accurate and have advantages over the reported methods which suffer from interference by large number of ions or require either heating or extraction or are less sensitive. Some factors such as initial cost of instrument, technical know-how, consumable and costly maintenance of technique restrict the wider applicability of these techniques, particularly in laboratories with limited budget in developing countries and for fieldwork are to be taken as the consideration for the determination of Molybdenum (VI) at microgram guantities, a number of hydrazone derivatives and several analytical techniques such as AAS, ICP-AES, ICP, X-Ray fluorescence spectroscopy and UV-visible spectrophotometry are employed.

Among them spectrophotometric methods are preferred because they are cheaper and easy to handle. In general the technique of solvent extraction was widely used in the spectrophotometric determination of metal ions. However, organic solvents such as benzene and chloroform are often carcinogenic, toxic and cause environmental pollution. It is significant to develop a method which does not involve solvent extraction.

The average daily intake of Molybdenum is 0.3 mg. Daily intake above 0.4mg can be toxic. Molybdenum deficiency less than 0.05 mg/day can cause stunned growth, reduced appetite and impaired reproduction.

Precision and accuracy: The precision and accuracy of the proposed methods are studied by analyzing (10 replicates) of Molybdenum (VI) and the RSD value was found to be 0.49%.

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**REFERENCE** [1].Singh RB, Jain P, Singh RP, Hydrazones as analytical reagents, Talanta, 29 (1982) 77-84. | [2]. Hayashi, K.; Sasaki, Y. Nakanishi, M. and Ito, S.;Bunseki Kagaku, 19 (1970) 1673. [3]..., Hayashi K.; Sasaki, Y. and Ito, K.; Bunseki Kagaku, 21 (1972) 1338. | [4]. Hayashi, K.; Sasaki, Y.;Taga Shira S. and Hirate K; Anal Chem. acta., 198 (1987) 271. | [5]. Underwood, E.J., "Trace Elements in Human and Animal Nutrition" 4th edition, Academic press, New York, (1977) | [6]. Khan,S.W.,CloutierR.O and Hidiroglou, J.,J.,Assoc.off.Anal.Chem., *62* (1979) 1062. | [7]. Studnicki, M., Anal. Chem., *51* (1976) 1336. | [8]. Barbooti, M. and Jasim, F., Talanta, 28 (1981) 359. | [9]. Quin, B.F. and Woods, P.H., Analyst, 104 (1979) 555. | [10]. Bradfield, E.G and Stickland, J.F., Analyst, 100 (1975)1. | [11]. Cruthamel, C.E. and Johnson, C.E., Anal. Chem., *26* (1954) 1284. | [12]. Allen, S.H. and Hamilton, M.B., Anal. Chim. Acta, 7 (1952) 483 [13]. Di, J., Ku, Y., Tu, Y. and Bi; S., Anal. Sci., 18 (2002) 125. | [14]. Elwanki, M.B., Seetharamappa, J. and Masti, S.O., Anal. Sci., 17 (2001) 1121. | [15]. Lkan, M., Kharun, M. and Chmilenlco, F., Talanta, 59 (2003) 605. | [16]. Filik, H., Tutem, E. and Apak, R., Annal. Chem. Acta, 50 (2004) 77 | [17]. Zaijun Li; Jan Tang and Jiaomai Pan; Analyst, 126 (7) (2011) 1154. | [18]. Kumar, A. Dass, R. and Sharma, B.G.: L. India Chem. Sci. 81 (2004) 1177. | [19]. Havashi, K.; Sasaki, Y. Nakanishi, M. and Iton, S. Bunseki Kagaku, 19(1970) 1673. Kumar, A.; Dass, R. and Sharma, R.G.; J. Indian, Chem. Soc., 81 (2004) 177. | [19]. Hayashi, K.; Sasaki, Y.; Nakanishi, M. and Ito, S.; Bunseki Kagaku, 19 (1970) 1673 | [20]. Hayashi, K.; Sasaki, Y. and Ito, K.; Bunseki Kagaku, 21 (1972) 1338. |