

ACSAD / Soil Science Division

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# ATOMIC ABSORPTION SPECTROSCOPY

Instrumental Conditions

DAMASCUS - 1978

THE ARAB CENTER FOR THE STUDIES OF ARID ZONES AND DRY LANDS ( A C S A D ) T

#### THE ARAB CENTER

For The Studies of Arid Zones & Dry Lands

## (ACSAD)

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- ATOMIC ABSORPTION SPECTROSCOPY -

INSTRUMENTAL CONDITIONS

Damascus 1978

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- A.C.S.A.D. -

- TECHNICAL NOTES - N- 1

- Atomic Absorption and Emission in Flame.

## INTRODUCTION

This booklet intends to facilitate the choice of experimental parameters for beginners in Atomic.- Absorption - It includes main elements usually determined in soil and water samples.

Optimization of experimental conditions may serve either the purpose of getting the best precision or the best sensitivity. These qualities are somewhat contradictory. So are most the parameters on which depend the final analytical result. This is why an average setting is usually given in operation booklets of instrument makers. The following informations include average and optimum adjustments.

It is assumed, that experimental conditions leading to accurate results are realised. These conditions may be summarized as follow:

- a. Viscosity and acidity (percent of acid in water) of unknown and standard solutions are identical.Optimum in percentage of acid is around 1 to 2%. Above 5% viscosity an anion effect become noti-ceable.
- b. The solution of soil or plant should be preferably done in HCl, HClO<sub>4</sub> or HNO<sub>3</sub>. Others minerals acids as H<sub>2</sub>SO<sub>4</sub> or H<sub>3</sub>PO<sub>4</sub> produce a considerable loss in sensitivity.
- c. It is also assumed that the matrix of the sample is known so that chemical interferences and effects, due to ionization and matrix can be overcome.

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d. The solution is free from solid particles which show a non-specific absorption and cause signal instability.

## DEFINITIONS

a) we used notation p.p.m (part per million) for micro-gram/milliliter; for an easier typing, despite the fact that they usually represent slightly different numbers.

We have used:

- m : maximum concentration expressed in ppm, for which the calibration curve is a straight line.
- s : sensitivity: concentration (expressed as ppm) giving 1% Absorption, or 4,4 milliabsorbance; sensitivities are average of datas given by several makers and represent usually the best conditions obtainable in aqueous solvent.
- 1 : detection limit: minimum concentration (expressed as ppm) for which ten successive measurments give an average absorption at least equal to two times average ground noise.
- p : precision: if n measurments are made on a solution having a concentration around m/2 and if v is variance of this n measurments : then: p = 100 v/M (n must be higher than 10).

M = mean value of these n measurements. How to use given indications:

- Experimental conditions are given in chronological order of operation.
  - Preparation of standard solutions.
  - Lamp current (Approx.).
  - flame stoechiometry.
  - burner eight.

- Main Line/secondary line with associated slit width.
- Interferences and their suppression.
- Internal standard.
- Depressing and Enhancing salvents.
- no indication has been given concerning photomultiplier voltage setting; it has to be adjusted, after the choice of optimum band pass (slit-width) so that saturation of P.M is not reached.
- sensitivities of secondary lines are given relatively to sensitivity of main line taken as unity.
- Stoechiometry is expressed as ratio of gaz flows (fuel/oxidant).
  Some instruments indicate pressures, making a conversion necessary.
- no data is given for aspiration rate, as departing from the optimum of each nebulizers will usually increase backgroung noise while decreasing sensitivity.

Preparation of interference suppressors:

- Strontium chloride:
  - weight 152,5 g SrCl<sub>2</sub>,  $6H_2O_2$ , dissolve in on lirer of bi-distilled water final solution is 5% Sr ( $\omega/\omega$ ).

Lanthonum chloride:

- Ca and Mg stock solutions and samples should be done in a 0.5% La aq. solution. It is therefore useful to have a large quantity of 5% La and 0.5% stock sol. for making dilutions when necessary.
- weight 5,85 gr of La<sub>2</sub>O<sub>3</sub> <u>slowly</u> dissolved in minimum 50% HCl (violent reaction) adjust to 1 l with deionized water. Solution is 5% La (w/w). (using the chloride directly is just much more expensive at equivalent purity ).

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- ALUMINIUM -

Al n = 13 M = 26,981  $f(Al_2O_3/Al_2) = 1,8895$  1/f = 0.5292

## MODE :

Emission has a better sensitivity than obsorption.

## ABSORPTION:

- 1. Stock Solutions:
  - a. 1,000 g Al 99,99% dissolved int. 100 ml conc. HCl after having added 1 drop of Hg. The solution adjusted to one liter is 1000 ppm.
  - b. 0.5294 g/l of Al gives: a 1000 ppm sol.in Al<sub>2</sub>0<sub>3</sub>.
  - organo metallic compound: Aluminium.
    cyclohexane butyrate or Aluminium 2 ethylhexanoate.

## 2. Lamp:

Sensitivity decrease as current increase. (best sensition for  $8 \rightarrow 10$  mA).

## 3. Flame:

The only usable flame is  $C_{2}H_{2}/N_{2}O$  stoechiometric.

## 4. Borner Height:

5 m/m. (given as height of light path above the burner).

## 5. Main Line:

 $309_{27} + 309_{28} \text{ nm} \text{ (band pars 1 nm)}$ . 1 = 0.05 m = 30 - 60s = 0.6 p = 0.3%

## 6. Secondary Lines:

line		band	pass
396, 15	D <b>.</b> 8	0.5	nm
308,22	0 <b>.7</b>	0.5	nm
394,4	0.5	0.5	nm
237,34	0,25	0.5	nm
237,84	0 <b>.1</b> 6	0.5	nm
257,54	0. 12	0.5	nm
256,80	0.08	0.5	nm

## 7. Interferences:

- No interferences of Ca Zn Pb Mg P
- Slight int. from Fe Cl. (Adjust Cl conc. of standard and unknown if HCl is used for sample preparation).

## 8. Correction of Interferences:

- Fe add the same quantity of Fe to samples and standards to correct enhancement due to Fe.
- <u>Al</u> is slighty ionised in hot C<sub>2</sub>H<sub>2</sub>/N<sub>2</sub>O flame. An ionization buffer (K, Cs, Sr, 1% as chloride) is added, and chloride concentration needs to be adjusted.

## 9. Enhancing solvent:

n-butanol, isopropanol, Ethanol.

## 10. Depressing solvent:

Hydrochloric Acid, chloride rich solutions.

## EMISSION

- Flame: C<sub>2</sub>H<sub>2</sub>/N<sub>2</sub>O stoech./rich (red feather 20/30 m/m).
- main line: 396, 15 nm  $(1 = 0.01 \ s = 0.5)$ .
- Secondary Lines: 394,4 nm and 309,3 nm.

## INTERFERENCES:

if solution is rich in Ca, emitting at 396,80 nm, the 396,15 nm line is to be avoided.

## CONCENTRATIONS:

- Soils: from 2 to 30% in Al<sub>2</sub>O<sub>3</sub>
- Plants: from 2 to 5000 ppm
- Waters: up to 5 ppm.

## REFERENCES:

- GUEST. Mc Pherson. The use of flame procedures in metalurgical analysis. Port II. Determination of Al in sulphide and silicate material, ores and slags. A.C.A. 78, 299, 1975.
- C.A.B.: Chemical det. of Al in soil and plant material Annotated Bibliography (1967 to 1974) N- 1674 Com. Agric. Bureau, Slough, ENGLAND.

## - CALCIUM -

В

Ca n = 20 M = 40,0 f(CaO/Ca) = 1,399 1/f = 0,714

1 eq/l = 20,04 ppm f(CaCO<sub>3</sub>/Ca) = 2,495 1/f = 0,401 1 ppm = 0,05 meq/l

#### MODE

emission is more stable, more sensitive and less subject to chemical interf. than absorption.

## ABSORPTION

## Stock Solutions

a 1000 ppm sol of Ca (i.e 50 meg/l) is obtained from:

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a. 2,7693 gr/l CaCl<sub>2</sub>
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b. 2,4973 gr/l CaCO<sub>3</sub> dissolved in minimum 10% HCl, complete to have final concentration 1,25% in HCl.

organometallic compound: calcium - 2 - ethylhexanoate.

- Lamp: sensitivity decreases as current increases if max. lamp current is 10 wA, best sensitivity is for 6 wA and best precision for 9 wA.
- Flame: C<sub>2</sub>H<sub>2</sub>/air (1/3.5 4.5) stoeck or lean but with maximum gas flow; this flame is non ionizing but subject to many interferences. C<sub>2</sub>H<sub>2</sub>/N<sub>2</sub>O (1/1,5 - 1,8) stoeck. less sensitive but free from main interferences. Needs an ionization buffer (0.1% KCl, KNO<sub>3</sub>, Cs...) to reduce Ca<sup>++</sup> concentration.

Burner height:

between 5 and 10 m/m for  $C_{p}H_{p}/air$  flame.

<u>Main Line</u>: 422,7 nm (band pass 0,4 - 1,0 nm) for  $C_2H_2/air$  s = 0.05 m = 10 for  $C_2H_2/N_20$  s = 0.02 m = 5 (Conc. given in ppm).

## Secondary Lines:

239,86 relative sensit. 0.005 (band pass 0.15 nm). allow to work in the range of 200 - 1000 ppm.(poor signal to noise ratio)

#### Interferences:

- a. Important interf. from PO<sub>4</sub>, SO<sub>4</sub>, Al, Si, NO<sub>3</sub>. corrected by a 1% La addition in standard and unknown if not too much Mg is present.
- b. if concentration of Mg is important, standard and unknown must be Mg saturated(ratio Mg/Ca about 100) care must be taken for samples containing phosphate, gypsum, dolomite.
- c. no interference of K, NH<sub>L</sub>, OH, C1, CO<sub>3</sub>.
- d. all mineral acids cause a slight decrease in Ca concentration.
- e. in solution rich in Si & Al it is necessary to use  $C_2H_2/N_2O$ flame and odd Sr to both standard and unknown.

#### EMISSION

🗕 flame:

- C<sub>2</sub>H<sub>2</sub>/N<sub>2</sub>O lean (5 m/m red feather).
- C<sub>2</sub>H<sub>2</sub>/air with decreased sensitivity.
- main line: 422,67

1 = 0.0003 m = 5 to 10

s = 0.003

A narrow bandpass is recommended to avoid disturbance due to the nearby strong emission of CN at 421 nm.

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CONCENTRATIONS

- Plants : 0,1 to 10% (105<sup>0</sup> dry weight basis)
- Water :-drinkable: less than 150 ppm
  - -from Limestone areas 50 to 150 ppm
  - -from gypseous rich areas: 100 to 500 ppm.
- -Soils, Rocks-from 0 to almost 40% Ca (equiv. to 100% CaCOz).

## **REFERENCES:**

- WARD, BIECHLER: "Rapid, direct determination of Ca in natural waters by A.A.S.

A.A. Newsletters 14, 29, 1975.

- MAHJOURY: Clay Mineralogy, physical and chemical properties of some soils in arid regions of Iran.
   S.S.S.A. proceedings 39, 1157, 1975.
- C.A.B.: Chemic. det. of Ca and Mg in soil and plant material.
  Annotated Bibliography N- 1675 Commonwealth Agricultural Bureau,
  1975.

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## - COBALT -

Co n = 27 M = 58,9332 f(CaO/CO = 1,2714) 1/f = 0.7864

MODE

Absorption is more sensitive than emission.

ABSORPTION

Stock Solutions 1000 ppm of Ca

- 1. 1,000 gr of metal dissolved in minimum conc. HCl.
- 2. 4,038 gr of CoCl<sub>2</sub>, 6H<sub>2</sub>0 or 4,9379 gr of Co(NO<sub>3</sub>)<sub>2</sub> in one liter. Since these salts are extremely hygroscopics, checktitration by complexometry is necessary. (see appendix).
- organometallic compound: Cobalt cyclohexanebutyrate.

Lamp: sensitivity decrease: slightly as lamp current inc. a good compromise is obtained at 50% of max. lamp current.

Flame:

C <sub>2</sub> H <sub>2</sub> /Air	stoech.	the	most	sta	ble.			
C <sub>2</sub> H <sub>2</sub> /Air	reducing	the	most	sen	sit.			
C <sub>3</sub> H <sub>8</sub> /Air	reducing							
C_H_/N_0	gives bet	ter	resul	ts	than	other	combinat	ions.

Main Line:

240,73 nm (band pass maximum  $\Omega_{\bullet}3$  nm) s =  $\Omega_{\bullet}1$  m = 5

Secondary Lines:

242,49 nm r.s = 0.8 range 2 to 50 ppm 241,46 nm 0.5 252,14 nm 0.3 - 12 -

243,58	nm r.s	= 0,3		
304,40	mu	0.08	band pass	0 <u>,</u> 5 mm
352 <b>,</b> 59	nm	0,06	range 100	to 2000 ppm
346,58	nm	0,03	band pass	0 <b>.15</b> nm
341 <b>,</b> 26	nm	0,03	bend <b>pa</b> ss	0 <b>.15</b> nm
347,40	nm	0,02	bend pass	0 <b>.15</b> nm
301,76	nm	0,01		
391,0	nm	0.005	band pass	0 <b>.5 nm</b>

#### Interference

Non specific absorption around the main line (240 nm) corrected either by Deuterium - continuous emitting Lamp background correction or by measuring non specific absorbance at 238,8 nm and "substracting" it from the final result (or adding if dealing with absorbance readings)

No noticeable interference in diluted solutions.

Interference of Fe, Ni, Cr if present in high amounts (1500 ppm Ni produce a 50% decrease of the signal).

## Enhancing Solvants:

 70% isopropanol in water is a solvant doubling sensitivity (for 240,73 nm line).

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#### Particularities:

 Precision at low concentration is very poor and depends strongly on flame conditions which are to be adjusted with utmost care.

#### EMISSION:

Main Line: 345,35 nm

Flame : C<sub>3</sub>H<sub>8</sub>/Air: best flame

 $C_2H_2/N_2O$ : (red feather 2 m/m) (caution : to much lean a flame is likely to explode inside the burner).

## CONCENTRATION

Soils

- 🛥 🔰 1 to 40 ppm (average around 5 ppm).
- ultrabasic (serpentines...) 0 to 2%

## Plants

- 0.01 to 1 ppm in usual soils (wide range of concentration in Co rich soils, in endemic species).

## REFERENCES:

- WARREN CARTER: Det. of trace amounts of Cu, V, Cr, Ni, Co, Ba
  in silicate rocks by flame A.A.S. Can.J. of Spectroscopy 20,1,1975.
- SAPEK: A.A.S. Det. of Pb, Ni,Co in soil extracts Chem. Analysis
  (WARSAW) 19, 687, 1974.
- SINHA-BANERJEE. Interf. in estimation of trace amounts of Co, Cu and Zn in soils by A.A.S. Technology (SINDRY - INDIA) 11, 263, 1974.
- HOCQUELLET. Det. of Co in animal feeding stuffs by flame and flameless A.A.S.

Annales. Falsif. Expert Chimiste. 67, 495, 1974.

 HARA, SONDRA, IWAI. Growth response of cabbage plants to transition elements under water culture conditions. Part II. Co, Cu, Zn, Mo.

S.S. Plant. Nut. 22, 317, 1976.

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- Cu -

n = 59 M = 63,646 f(CuO/Cu) = 1,251 1/f = 0.798

## MODE :

Absorption is more sensitive.

## Absorption:

- Stock Solutions: 1000 ppm Cu:
  - 1. 1,000 gr of metallic Cu dissolved in 3 ml HND<sub>3</sub> conc. (cold)
  - 2. 3,929 gr CuSO<sub>4</sub>, 5H<sub>2</sub>O/1 H<sub>2</sub>O.
  - 3. 3,798 gr Cu(NO<sub>3</sub>)<sub>2</sub>, 3H<sub>2</sub>O/1 H<sub>2</sub>O. If this option is chooser, Cu concentration has to be checked by complexometry (see appendix).

Lamp current:

If max. lamp current is  $5 - 7 \text{ mA}_{,}$  optimum is 4 maximum sensitivity is 3 and max. precision  $4 - 5 \text{ mA}_{,}$ 

Flame:

C<sub>2</sub>H<sub>2</sub>/Air stoech. of Lean. C<sub>3</sub>H<sub>8</sub>/Air gives a better sensitivity.

Burner height:

11 m/m for C<sub>2</sub>H<sub>2</sub>/Air stoech. flame.

Main Line: 324;75 nm

1 = 0.006 s = 0.06 m = 4 to 6 (1 ppm produce around 100 milliabs.).

Secondary Lines:

line	relative seas.	band pass (nm)
327,4 nm	r.s 1/2	0.5
217 <b>,</b> 89	1/5	0.3
218,2	1/5	0.3
246,51	1/6	0.3
222,5	1/10 1/20	0 <b>.15</b>
249,22	1/100	0.5
244,16	1/300	1

## Interferences:

- large excess of transition elements give a depressed reading, especially for Zn.
- Cu is one of the elements giving the most easy-to-read
  results. (if no computored integration is available).
- virtually no other interf. by other elements in  $C_2H_2$  air flame.

## EMISSION:

Line: 327,40 nm : max. sensitivity 324,70 nm : less sens. both line have to be used in conjonction with the minimum band pass (and maximum photo multiplication) a lean  $C_2H_2/N_2O$ flame is more sensitive that  $C_2H_2/Air$  stoech.

## Concentration:

Soils : 2 - 200 ppm plants: 1 - 25 ppm Water : drinking water : max. 1 ppm irrigation water: max. 0.5 ppm

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- Fe -

Fe n = 56 M = 55,847  $f(Fe_2O_3/Fe_2) = 1,429$   $1/f = O_{\bullet}699$ f(FeO/Fe) = 1,286  $1/f = O_{\bullet}7773$ 

## ABSORPTION:

- Stock solution (1000 ppm Fe)
- 1,000 gr pure Fe in 5 ml HCl conc. (add few drops of HNO3 conc.); adjust to 1 l.
- 7,021 gr (NH<sub>4</sub>)<sub>2</sub> Fe(SO<sub>4</sub>)<sub>2</sub>, 6H<sub>2</sub>O Dissolv. Adj. to 1 1.
- 4,840 FeCl<sub>3</sub>, 6H<sub>2</sub>C, control Fe concentration by complexometry.

All stock solution must be kept in a dark place in polythene container.

Lamp current:

max. precision is near max current max. sensitivity is 70% of max. current.

Flame:

C<sub>p</sub>H<sub>p</sub>/Air Lean (make the yellow fringe just disappear).

Main Line: 248,33 nm

 $1 = 0_{\bullet}01$   $s = 0_{\bullet}1$   $m = 5_{\bullet}10$ 

Other Lines:

Fe has many	absorbing li	nes.	Among them:			
248,81	r.s	1/2				
252 <b>,</b> 21	r.s	1/2				
271,98	r.s	1/3	(relatively	large	band	pass
			poss.)			
302,06	<b>r</b> .9	1/4				

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252,74	r.s 1/5
371,99	1/8 (band pass 0.3 nm, very stable
	meas.)
269,69	1/9
385,99	1/12 (band pass 0.3 nm - doublet).
344,06	1/100
392,03	1/250 (band pass 0 <sub>0</sub> 3 nm).

## Interferences:

- Especially from Si and PO<sub>4</sub>. This chemical interference is depending on instrument settings.
- With C<sub>2</sub>H<sub>2</sub>/Air axydant (very Lean), Fe interf. are decreased, together with sensitivity.
- Interference of Si may be overcome by using a 0.2% CaCl<sub>2</sub> sol as int. standard (see annex), or by using the hotter  $C_2H_2/N_2O$  flame.
- No Interference from
  - 3000 ppm K, Ca
  - 1000 ppm Na, Mg, SO<sub>4</sub>, Cl, transit. metals.
- Background correction for 248,81 Line.

## EMISSION:

- Line : 371,99 minimum slit-width and max. photomult.
- <u>flame</u>: C<sub>2</sub>H<sub>2</sub>/N<sub>2</sub>O Lean (red feather 5 m/m).
- <u>Concentration</u>:

Water : surface water alcaline surface: less than 1 ppm. under ground water: 1 - 3 ppm. drinking: less than 0.1 to 0.2 depending on local regulations. - 18 -

REFERENCES FOR TRANSITION ELEMENTS - (see also Co page 13.)

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  ANALUSIS, 3, 345, 1975.
- WALLINGFORD & al: Effects of beef feedlot manure and lagoon water on Fe, Zn, Mn, Cu content in corn and DPTA soil extracts.
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- WARD BROOKS REEVES. Det. of Cu, Cd, Pb, Zn in soils, stream sediments, waters, and nat. vegetation around Tui Mine.
  N. acaland J of Science 19, 81, 1976.
- C.A.B. annotated bibliography on chemical det. in soil and plant
  N- 1679 for copper, N- 1681 for Fe, Mn. Commonwealth Agricultural
  Bureau, Slough, ENGLAND.

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## - POTASSIUM -

n = 19 N = 39,102 f(K<sub>2</sub>O/K<sub>2</sub>) = 1,204 1/f = 0.830 1 millieq = 39,102 milligram.

## MODE :

The best advantage of emission over absorption is to **ellow** to work at 404,4 nm and give a very low **sensitivity**, without supressing interferences.

#### ABSORPTION:

Stock solutions: 1000 ppm of K: (25,62 meg/1.).

- 1,907 gr KCl; dry 105<sup>0</sup> dissolve in 1 l.
- 2,586 gr KND, dry 105<sup>0</sup> dissolve in 1 1.
- organo-metallic Cyclohexanebutyrate of K.

#### Lamp:

max. precision 80% max. current. max. sensitivity 50% max. current.

#### Flame:

The best flames are in the order:  $C_4H_{10}/Air$ : cylindricol burner, no interferences.  $C_3H_8$  /Air: few interf.  $C_2H_2$  /Air: (Lean) important interf. of sodium.

#### burner height:

Critical for reducing interference of sodium. Less than 1 m/m with  $C_{2}H_{2}/air$  flame.

Main Line: 766,5 nm (resonnance Line).

1 = 0.002 s = 0.05 m = 2 ppm (data for  $C_4 H_{10}/air$  flame).

Secondary Lines:

769,9	r.s 1/2	(band pass 1 nm – resonna <b>nc</b> e
		Line)
404,72/404,41	r.s 1/200	) (band pass 0.4 nm - doublet).

## Interference:

As for all **claime** metals, ionization causes a depression of the response. Thus, easily ionizable elements, Na, Rb, Cs, Li enrich the flame in electrons reduce the number of K in the sample and gives a final overestimated result. A ionization buffer 0.1% Cs of Sr<sup>\*</sup> supress this effect (it has to be added to both standard and samples). A decrease in flame temperature is also of benefit.

Enhancing solvent:

- Butanol 4% (in water).

## Particularity:

Potassium emits and absorbs in red-band of visible wavelengths. It may be necessary to change photomultiplier to keep on with max. sensitivity.

## EMISSION:

Flame :  $C_2H_2/Air$ : blue flame (oxydant).  $C_3H_8/Air$ : blue flame (oxydant).

 Sr connot be used with sulfate containing solution, because of strontium sulfate precipitation. Burner height:

from 10 to 25 m/m depending on burner geometry.

Concentration:

Soils : 0 to 5% K<sub>2</sub>0. Plants : 0.2 to 10%. Waters : surface:5 - 15 ppm. ground : usually less than 20 ppm. may reach higher conc. occasionally.

# REFERENCES:

(see Ca and Na also)

- NEUNER & al: Automated flame photometric det. of K<sub>2</sub>O in fertilizers.
  - J. Asso. off. Anol. Chemists 58, 923, 1975.

- Mg n = 12 A = 24,305 f(Mg/MgD) = 1,658 1/f = 0.603 1 meq = 12.15 mg

## MODE:

## ABSCRPTION:

Stock solution: 1000 ppm (82,4 meq/1.).

1,000 gr of Mg ribbon diss. in 7 ml conc. HCl and adj. to
 1 1.

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- 3,916 gr MgCl<sub>2</sub>, 6H<sub>2</sub>O/11 H<sub>2</sub>D.
- organo-metallic compound: many organo-magnesians.

## Lamp current:

- Max. precision obtained for max. lamp current.
- Max. sensitivity 50% max. Lamp current.

#### Flame:

- C<sub>p</sub>H<sub>p</sub>/air reducing (rich yellow).
- C<sub>p</sub>H<sub>p</sub>/N<sub>p</sub>D (1/1:25) stoeck.

## Burner height:

8 to 10 m/m.

Main Line: 285,21 (band pass 0.1 nm).

1 = 0.002 s = 0.05 m = 0.4 (0.1 ppm gives, 100 milliabsor bance if correct adj. are made). - 23

Secondary Line:

202,58 r	•8	1/25 (band pass 0.8 nm)
279,58 r	• 8	1/100 (ionic Line)
260,27		(ionic line)

ionic line could be used if much Mg is present in a medium free of other easily ionizable elements.

## Interference:

- a) Aluminium, phosphate, chloride, Sulphur, Si, depress the response in the C<sub>2</sub>H<sub>2</sub>/air flame. This effect is reduced by making solution in 1% (w/w) Lanthanum, or Strontium.
- b) enhancing effect of acetic ion<sup>\*</sup> and alcaline metals. Effect reduced by using an ionization buffer (KCl 0.1%  $\omega/\omega$ )
- c) Varying interferences of Cu, Ti, Fe, Al. All interferences are strongly dependent on size of droplets reaching the flame and can be almost supressed in laminor flow.(except for Al).

## EMISSICN:

Flame:

 $C_2H_2/N_2O$  Lean. wave Length 285,21 nm. No advantage over absorption.

## Concentrations:

- Water:

surface (on acid soils or rocks) 0 - 5 ppm
 (on limestone - dolomite) 10 - 50 ppm
 wells up to 4000 ppm.

 Particular attention is to be paid to exchangeable Mg due to high concentration of acetate in many methods.

## MODE :

## ABSCRPTION:

## Stock solutions: 1000 ppm Mn.

- → 3,607 gr MnCl<sub>2</sub>/1 or 3,076 gr/1 MnSO<sub>4</sub>, 4 H<sub>2</sub>0 in water.
- 1,000 gr metallic Mn in minimum conc. HCl.
- Organometallic: Mn cyclohexanebutyrate.

## Lomps

Sensitivity doesn't change too much with lamp current, therefore current should be set to the value giving the best precision (80% max. current).

## Flame:

C2H2/air: reducing best sensitivity. C2H2/air: oxydizing best precision. C3H8/air: best stability (good sensitivity). C2H2/N2O: Lean, 5 m/m red feather. Suitable for 403,08 nm. (burner height: 10 m/m).

## Burner height:

5 to 8 m/m for C<sub>2</sub>H<sub>2</sub>/air oxidizing.

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## Main Line:

279,48 nm minimum available band pass (279,83 and 280,11 peaks must be separated from 279,48).

## Secondary Lines:

403,08	r.s	1/20 (wide band pass possible).
321,70	<b>r</b> .8	1/2000

## Interferences:

- Si even in low concentrations. (correction by addition of
  50 to 2000 ppm CoCl<sub>2</sub> to standard and sol).
- Phosphates, perchlorates, transition metals (Fe in excess of 25 ppm), in reducing flames.
- No interference of 1000 ppm from Na, K, Ca, Mg. If concentration in these metals.

#### Concentration:

Sol : 0 - 1% en MnO. Plant : 5 - 5000 ppm. Water : less than 0.5 ppm. (surface). less than 0.1 ppm. (drinking water).

#### EMISSION:

403,08 nm using a lean  $C_{2}H_{2}/N_{2}O$  flame

REFERENCES: (see also transition elements)

- SIMAN CRADOCK HUDSON Development of Mn toxicity in posture legumes under extreme climatic conditions P.S 41, 129, 1974.
- MORTVEDT GIORDANO: crop resp. to Mn sources applied with ortho and polyphosphate fertilizers. SSSAP 39, 782, 1975.

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- Na -

Na n = 11 A = 22,9898  $f(Na_2O/Na) = 1,3479$  1/f = 0.74181 millieq = 23 mgr 1 ppm = 0.0434 meq/1.

## MODE :

ABSORFTION: for 1000 ppm Na : (i.e 43,5 meq/1).

- 2,5420 gr/l NaCl (from 120<sup>0</sup>C dried NaCl)
  for 20 meg/l Na.
- 1,1714 gr/l NaCl.
- Organometallic Na cyclohexanebutyrate.

#### Lamp:

Sensitivity doesn't decrease when increasing lamp current. can be used for best precision.

#### Flame:

 $C_3H_8$ /air max. sensitivity. Effect of ionization of other alcaline metal is reduced by the low temp. of the flame.

C<sub>2</sub>H<sub>2</sub>/air: Lean (1/5,5) less sensitive, less stable.

Main Line: 589,0 with narrow band pass)

 $1 = 0_0001$   $s = 0_0005$  m = 1 ppm

## Secondary Lines:

	589,59	r.8 1/2	(resonnance line - needs a narrow
			band pass).
	589,0 + 589,59	г.в 1/1.2	(entire doublet band pass 0.6 nm).
	330,2	r.s 1/300	(narrowest possible band pass).
	330,3	r.s 1/300	(narrowest possible band pass).
or :	330,2 + 330,3	r.s 1/200	

#### Interference:

Few interfering elements, except from high concentration of Cl or SO<sub>4</sub> ion. Effect from easily ionizable elements is supressed by 1000 ppm of Ca, Li or K as buffer.

## Enhancing solvant:

n. butanol 6% (w/v in water).

## EMISSION:

- flame : C<sub>3</sub>H<sub>A</sub>/air Le**e**n (minimum burner height).
- Line : 589,0 nm (minimum band pass).

## Interferences:

Unlike Potassium, there are more interferences in emission mode than in absorption Fe, Ca are enhancing, in addition, large quantities of Ca cause an increase of background noise due to emission bands of Ca hydroxydes (from 572 nm to 602 nm peak to peak).

#### Concentration:

 The range of concentration, in natural water, in soils and in plants is very wide as just an idea: for soils : from 0.1 to 500 meq/l in saturated extract. for plants: from 0.01 to 5%. - 28 -

- Si •

n = 14 A = 28,086  $f(SiO_{2}/Si) = 2,1393$  1/f = 0.4674

## ABSORPTION:

- <u>Stock solution</u>: 1000 ppm Si
  - a. 117,86 g/1 Si02, 12N03, 26H20
  - b. 7,552 g/l Na<sub>2</sub>SiO<sub>3</sub>, 5H<sub>2</sub>O (diluted in hot water)<sup>\*</sup>.
  - c. 2,1393 gr pf pure silica used with 10 gr de solid NaOH
    dilute residue of fusion in hot water.
  - d. organometallic : octaphenyl cyclotetrasi loxane.

#### Lamp:

best sensitivity: 50 - 70% max. current. best precision : 80 - 100% max. current.

## Flame:

Only  $C_2H_2/N_2O$  Lean with maximum gas flow (to avoid flash - back).

**Burner Height:** 

Silicium absorbs in a narrow region above the burner (7 to 8  $m/m_{\odot}$ 

Main Line: 251,61 m/m

s = 1,5 m = 150

 Exact concentration is to be verified by standard gravimetric analysis. 29

## Secondary Line:

250,69	r.s 1/2,5	252,41	r.s 1/5
251,43	r.s 1/3	221,67	<b>r</b> .s 1/5
252,85	r.s 1/4	221,89	r.s 1/10

## Interference:

- HF, H<sub>3</sub>80<sub>3</sub> give a depression of the signal.
- presence of an easily ionizable element gives an enhancement.
  Therefore with excess of Na, K, Mg, Ca, a 0.1% buffer of any alcaline chloride should be used.

## EMISSION:

## Line:

251,61 nm - (narrow band pass) flame  $N_2O/C_2H_2$  stoech. (15 m/m red feather). Measurment at 7 - 9 m/m.

## Concentrations:

Plants: 0.01 to 5% **Water** : potability limit: 25 ppm.

## REFERENCES

MORRIS - Det. of Si in cured tobacco leaf by A.A.S. Tobacco Science 18, 120, 1974. - 310 -

## – Zn –

Zn n = 38 A = 65,37 f(ZnD/Zn) = 1,2447 1/f = 0.803

## MODE:

## ABSORPTION:

## Stock solutions: 1000 ppm

- a. 1,000 gr Zn dissolved in 3 ml of conc. Hcl
- b. 1,245 gr ZnO, few drop of water, dissolved in diluted HC1

## Lamp:

- sensitivity decreases sharply as lamp current increases.
- max. sensitivity 50% max. lamp current.
- max. precision 80% max. Lamp current.

## Flame:

- C<sub>2</sub>H<sub>2</sub>/air: reducing (fuel rich).
- C<sub>3</sub>H<sub>2</sub>/air: (flame absorbing).

## Burner Height:

6 m/m for  $C_{p}H_{p}/air$  flame.

#### Main Line:

213,06 nm (minimum band pass to exclude the 213,60 Line of Cu) s = 0.02 ppm m = 1 ppm

(data given for isopropanol 50% and  $C_3H_{\rm A}/{\rm air}$  flame).

#### APPENDIX

1. ABREVIATIONS:

AAN Aas A.C.A.	:	Atomic Absorption Newsletters. (Ed. Sabina Savin. Perkin- Elmer Corporation, Main Ave, Norwalk CT 06856 U.S.A.) Atomic Absorption Spectroscopy). Analytical Chemica Acta. (Elsevier Publ. Co P.O.Box 211 - Amsterdam Netherlands).
AN ANA ANCH Adacj C.A.B.		Analusis - Société de Productions documentaires & Ave. du 18 Juin 92500 Rueil-Macmaison France. Analyst (Ed. J.9 Atrill 9-10 Savile Road LONDON WIXIAF). Analytical Chemistry. Association of Offic. Anal. Chemists (Ed. Reynolds, P.O.Box 540 Benjamin Franklin Station, Washington D.C 20044). Commonwealth Agricultural Bureau. Slough - England.
CJS NZJS	:	Canadian Journal of Spectroscopy.(Moltiscience Publications Ltd. 1253 Me Gill College. Suite 404 - Montreal 110 - CANADA New Zealand Journal of Science.(Ed. J.C GREORY, Dep. of Sc. and Indus. Research, Private Bay, Wellington New Zealand.
P.S.	:	Plant and Soils. (Ed. M. Nihjoff 9–11 Lange Voorhout Box 269. The Hague, Netherlands.).
SSPN	:	Soil Science and Plant Nutrition. Japan Publications Trading Co Ltd. Box 5030 Tokyo - Interna- tional. Tokyo JAPAN.
SSSAP	:	Soil Science Soc. of America proceedings. SSSA, 677 St. GEORGE ROAD MADISON Wis. 53711.

TAL : Talanta: Headington Hill Hall Oxford OX3 OBW ENGLAND.

## 2. Definitions:Methodology

a. Accuracy:

measure the agreement between a measured value: m and the value accepted as "true": mo. for instance: a = 100 (m - mo)/mo. the smaller a, the more accurate the measured value m. is.

b. Precision:

measure the agreement among tests results as expressed in terms of standard deviation or relative standard deviation if n differents values are obtained by the same operator using the same method on the same sample: if further more: M is the average of these n values. S is the standard deviation n values. precision is: p = 100 S/M. the smaller p, the more precise the <u>method</u> concerned is. - 32 -

## c. <u>Repeatability</u>:

measure how much a given method is reliable as far as the constency of its results is concerned: Several series of tests results are performed on the same sample by the same operator using the same method at different periods of time. (two for instance). If S1 and S2 are the standard deviation, the smaller is the difference (S1 - S2), the more repeatable is the method.

d. Reproductibility:

measured by the standard deviation of a serie of test results given by differents laboratories, using the same method on the same sample.

3. Definitions: Apparatus

## a) Intrinsic properties of apparatuses:

- Resolution:

The resolution, given in nm units, is a fixed property of a given monochromator (single or double grating). A usual figure is 0.2 nm (from 0.05 to 0.4 nm). A resolution of 0.2 nm means that two lines distant of 0.2 nm (peak to peack) are folly separated.

## - slit width:

measure the amount of light reaching the photomultiplier. One should always choose the widest slit compatible with the selection of only one resonance line.

- reciprocal linear dispersion: (r.1.d)

measure the spectral bandwidth for a geometric slit width of 1 mm. A small linear dispersion (nm/mm) means that a small spectral bandwidth is associated to a wide slit, the result being a high signal to noise ratio. (Double-grating may have a r.l.d of 1 nm/mm in v.v) value of 1.3 to 4 nm/mm are reported by manufactures.

## b) display and read-out accessories:

a wide range of display and read-out recording systems is octually ovailable, among the mode of display: - <u>scale voltmeters</u>: (scale calibrated in % Absorption or Ab-

- sorbance units).
- Digital voltmeters: connected to a micro-compoter allowing:
  - discontinuous display (various sequences of time)
  - integration over a given period of time
  - peak height measurment
  - peak surface value
  - average.

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Among the read-out recording systems.

- in addition to the above mentioned devices, record of results can be achieved either by a digital printer or a strip-chart potentiometric recorder. Each system has its pros and cons, but only a strip-chart recorder provides a complete show of what-is-going-on during the all process. It is a good proctice to keep in mind how much reach the absorption percentage during each sample aspiration. Digital voltmeters are very accurate and versatile but one has to know that when a result is given in integrated mode (i.e an average, a surface) the fluctuation of the signal doesn't appear. The same remark is valid for artificial zeroing of the baseline.
- 4. General Bibliography:

ANGINO E.E and G.K BILLINGS "Atomic Absorption Spectrometry in Geology 1972. (ELSEVIER Pub.) CORNIL-LECENT Spectrographic determination of some non-metallic elements in water and in biological or geological samples. AN. 3, 11, 1975. cresser; A course on Analytical Techniques. Ministry of Agriculture IRAN (1975) ISAAC-KERBER Instrumental methods for soils and plant tissue analysis. SSSAP. 1971. DINNIN. (General review of methods for agriculture) ANCH. 47, 5, 1975 GABRIELS Analysis of Horticultural products, soil mixes and irrigation waters by FE and AAS. AN, 3, 139, 1975. JEANROY Analyse totale par A.A des roches, sols, minerais et ciments après fusion au metaborate de Sr. AN. 2, 10, 1973. LIDDEL. Noise at detection limit levels in AAS. ANCH 48, 1931, 1976. MITCHEK The Spectrochemical Analysis of soils, plants and related material. TA, 20, 5, 1973. PINTA. Spectrometrie d'Absorption Atomique (200 L). MASSON-ORSTOM-PARIS 1971. SCHRENK Analytical Atomic Spectroscopy. (Plenum Press N. YORK 1976). UELZ.B. Atomic Absorption Spectrophotometry, verlag chemie-wienheim.N.YORK 1976. WHITESIDE Methods of dissolution of siliceous samples for AAS (Pye unicam 1975. ENGLAND).

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## 5. CHECK TITRATION OF Co STOEK SOLUTION:

Stock solution supposeddly 1000 ppm is diluted 10 times. To a 20 ml aliquot add: 3 ml of PH 4,3 buffer (ACoNa 105 gr/l in Acetic Acid 10%), 5 drops of Alizarin complexone indicator and 3 drops of xylene cyanol FF. Heat to 80° and titrate with 2 x 10-2M EDTA to a green end-point. (change of color is from grey to red and to green.

#### optional method:

To a 20 ml aliquot add 25 ml dist.water, 5 ml of PH 4,5 buffer (ACoNa 136 gr/l in Acetic Acid 6%), 5 drops of xylenolorange tetrasodic salt (0,2% in water) heat to 70-80°C, and titrate to violetto-yellow end-point.

#### 6. CHECK TITRATION OF Mn STDEK SOLUTIONS:

 when dissolving metallic Mn in acid, a brown precipitate of MnO2 may be formed. Add 10% sodium hydrosulfite Na2S2O4 untill it redissolves.

#### Titration of Mn (II):

- Dilute 10 ml of the supposed 1000 ppm stoek solution to 100 ml, add 1 gr of ascorbic acid, dissolve. Add 5 drops of solochrome black T as indicator 10 ml of buffer PH 10 (70 amonium chloride added to 570 ml of conc. amonia sol, diluted to 1 l) titrate immediately with 2.10<sup>-2</sup>M EDTA to the red to blue end-point, (from T.S West, complexome try with EDTA and related agents, BDH publ.).

- Mn (II) is converted to MnO4<sup>-</sup> by amonium persulfate or sodium bismutate oxydation. Pink MnO4<sup>-</sup> is titrated back to Mn(II) by ferrous salt.

- To 10 ml solution add sufficient nitric acid to have a final concentration 2,4 M about add 15 ml silver nitrate 0,1 M and 50 ml of ammonium persulfate 10% - Heat to  $80^{\circ}$  -  $100^{\circ}$  for 15 min titrate by Mohr salt N/10 just to disapearance of the pink color.

#### References:

#### G. CHARLOT

Chimic analytique quantitative - T.II MASSON - 1974

T.S West

complexometry with EDTA and related reagents. BDH Chem. Pub. 1969

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