

EFFECT OF ORGANOTIN SPECIES ON THE EMISSION SIGNAL IN ICP-AES

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INTRODUCTION

It is generally accepted that the analytical response in ICP-AES is not influenced by the chemical form of the analyte. Nevertheless, differences between signals of organometallic and inorganic compounds of the same analyte have been reported [1,2]. Thus, it has been proved that inorganic tin solutions provide higher emission signals than organotin ones. Moreover, this behaviour has been observed operating with different spray chambers and experimental conditions[3].

The aim of this work is to evaluate the analyte chemical form on the emission signal in ICP-AES. To this end, different tin solutions have been analysed using a pneumatic concentric nebulizer coupled to a cyclonic spray chamber. The effect of the liquid and gas flow rates on the aerosol drop size distribution, solution transport rate and emission signal in ICP-AES have been studied.

REFERENCES:

- (1) L.C. Alves, M.G. Minnich, D.R. Wiederin, R.S. Houk, J. Anal. At. Spectrom., 9, 399 (1994).
- (2) C. Rivas, L. Ebdon, S.J. Hill, J. Anal. Spectrom., 11, 1147 (1996).
- (3) J. Montiel, J. Mora, Influence of the analyte chemical form and the sample introduction system on the analysis of organotin compounds by ICP-AES, Euroanalysis XIII, Salamanca, 2004.

EXPERIMENTAL

➤ 10 ppm Sn in EtOH 0.75% from: SnCl₄; ⁿBuSnCl₃ (MBT); ⁿBu₂SnCl₂ (DBT); ^tBu₂SnCl₂ (DTBT)

➤ Sample introduction system:



Meinhard ® TR-30-A3 nebulizer



Cyclonic Spray Chamber

➤ ICP-AES operating conditions:

PERKIN ELMER OPTIMA 4300 DV	
RF Power (W)	1300
Plasma gas flow rate (L/min)	15
Auxiliar gas flow rate (L/min)	0,2
Nebulizer gas flow rate, Q _g , (L/min)	0,6
Liquid flow rate, Q _l , (mL/min)	0,2 - 1,0
i.d. injector (mm)	1,2
Sample and integration time (s)	Automatic
Torch position (mm)	15
Vision view	axial

RESULTS

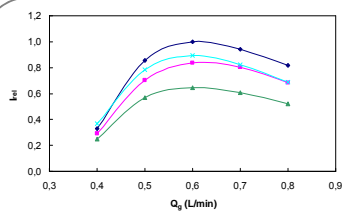


Figure 1. Effect of Q_g on the relative tin emission signal for the different solutions tested: (—●—) SnCl₄; (—■—) MBT; (—▲—) DTBT; (—▼—) DBT. Q_l = 1,0 mL/min. Sn II 283,998 nm

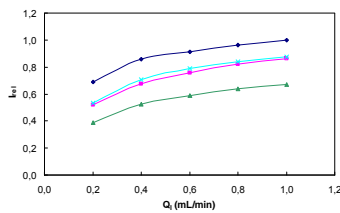
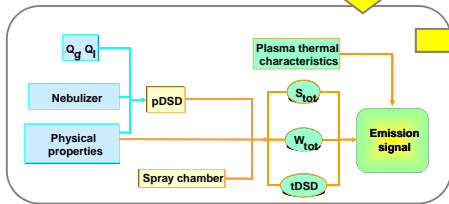


Figure 2. Effect of Q_l on the relative tin emission signal for the different solutions tested: (—●—) SnCl₄; (—■—) MBT; (—▲—) DTBT; (—▼—) DBT. Q_g = 0,6 L/min. Sn II 283,998 nm

$$I_{SnCl_4} > I_{DTBT} \approx I_{MBT} > I_{DBT}$$

EMISSION SIGNAL DEPENDS ON THE TIN COMPOUND



- There is no effect of analyte chemical form on:
 - Primary droplet size distribution (pDSD) (1)
 - Tertiary droplet size distribution (tDSD) (2)
 - Solvent transport rate (3)
 - Plasma characteristics (4)
- For a given Q_l, the analyte transport rate, W_{tot}, is independent on the analyte and the solution employed.

Q _l (mL/min)	Analyte	W _{tot} (μg/min)				
		Blank*	SnCl ₄	MBT	DBT	DTBT
0,2	Mg	0,50 ± 0,06	0,57 ± 0,02	0,59 ± 0,05	0,53 ± 0,06	0,56 ± 0,06
0,2	Mn	0,52 ± 0,07	0,54 ± 0,03	0,54 ± 0,05	0,48 ± 0,05	0,52 ± 0,05
1,0	Mg	1,62 ± 0,11	1,67 ± 0,13	1,66 ± 0,11	1,74 ± 0,02	1,9 ± 0,2
1,0	Mn	1,74 ± 0,12	1,63 ± 0,09	1,67 ± 0,05	1,77 ± 0,04	1,8 ± 0,3

* Ethanoic matrix without Sn

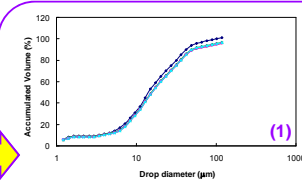


Figure 3. Primary droplet size distributions for the different solutions tested: (—●—) SnCl₄; (—■—) MBT; (—▲—) DTBT; (—▼—) DBT. EtOH 0,75%. Q_g = 0,6 L/min; Q_l = 1,0 mL/min.

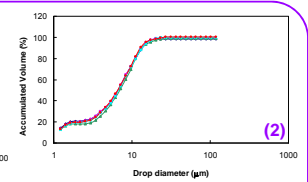


Figure 4. Tertiary droplet size distributions for the different solutions tested: (—●—) SnCl₄; (—■—) MBT; (—▲—) DTBT; (—▼—) DBT. EtOH 0,75%. Q_g = 0,6 L/min; Q_l = 1,0 mL/min.

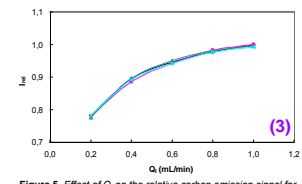


Figure 5. Effect of Q_g on the relative carbon emission signal for the different solutions tested: (—●—) SnCl₄; (—■—) MBT; (—▲—) DTBT; (—▼—) DBT. Q_l = 0,6 L/min. C I 193,030 nm

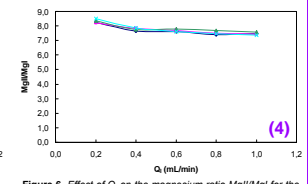


Figure 6. Effect of Q_g on the magnesium ratio Mg/II for the different solutions tested: (—●—) SnCl₄; (—■—) MBT; (—▲—) DTBT; (—▼—) DBT. Q_l = 0,6 L/min.

COMPOUND VOLATILITY

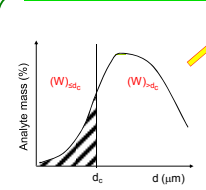


Figure 7. Analyte mass distribution of the primary aerosol (including a hypothetical value of the cut diameter of the spray chamber, d_c)

$$W_{tot} = (W)_{>d_c} + (W)_{<d_c}$$

✓ Low volatile analyte ⇒ W_{tot} ≈ (W)_{>d_c}
 ✓ High volatile analyte ⇒ W_{tot} ≈ (W)_{>d_c} + (W)_{<d_c}

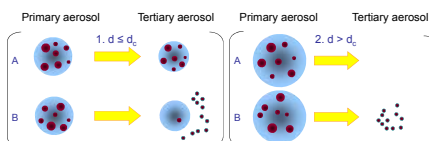


Figure 8. Effect of analyte volatility and primary aerosol drop size on the tertiary aerosol composition. 1: Analyte initially contained in drops with diameter equal or less to d_c; 2: Analyte initially contained in drops with diameters higher than d_c; A: Low volatile analyte; B: High volatile analyte.

Compound	Boiling Point (°C)
MgCl ₂	1410
MnCl ₂	2160
SnCl ₄	110
DTBT*	210
MBT*	240
DBT*	300

* Estimated values (T-H-E rule).

$$(W)_{>d_c} > (W)_{>d_c} \approx (W)_{>d_c} > (W)_{<d_c}$$

$$I_{SnCl_4} > I_{DTBT} \approx I_{MBT} > I_{DBT}$$

THE HIGHER COMPOUND VOLATILITY THE HIGHER EMISSION SIGNAL IN ICP-AES