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PROFESSOR SORIN MAGER BIOGRAPHY

A brilliant scientist, the leader of the Organic Chemistry School of Cluj in the last decades, was the dean of the Faculty of Chemistry and Chemical Engineering for 8 years (1992-2000) in a difficult period of structural changes. At his 75 anniversary he is the same active, creative, exigent and productive professor as he is known by the academic community for nearly 50 years.

Borned on May 7th 1931 in Arad, a town charged with history in the western part of Transylvania, in a family with a strong attachment for the culture and national traditions. He distinguished himself already during his years as student in the lyceum and faculty, as an outstanding personality



dedicated for study, with scientific curiosity and interest in organic chemistry.

He obtained the PhD degree in Organic Chemistry in 1964, at "Babeş" University, under the supervision of Professor Maria Ionescu with the thesis: "*o*-Nitrobenzylydene acetals of myoinositol".

He held different positions: 1953-1958 teaching assistant ("V. Babeş" University, Cluj-Napoca); 1958-1961 chemist in "Terapia" drug plant; 1961-1963 teaching assistant (Pedagogical Institute of Cluj-Napoca); 1963-1968 lecturer (Pedagogical Institute of Cluj-Napoca); 1968-1970 lecturer, 1970-1990 associated professor and from 1990-2001 professor and from 2001 consulting professor at "Babeş-Bolyai" University.

His creative scientific work was strongly influenced by the one year post-doctoral research stage spent at Notre Dame University at Notre Dame, Indiana, USA in 1971-1972 in the group of professor Ernest Eliel, the most important researcher in the field of stereochemistry and of saturated six membered ring heterocycles. After his return in Roumania he edited books and papers which are the first ones (in Roumania) in Stereochemistry and in Organic Structural Analysis and he is considered effectively as being the parent of these disciplines in our university.

Concerning his teaching activity, many series of graduates of our faculty strongly appreciated his clear, logic, systematic and enthusiastic

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lectures. The colleagues and the academic community also appreciated his devotion towards Alma Mater Napocensis, his spirit of justice, his morality and collegiality. Professor Sorin Mager gave many lectures in important universities and International and National Conferences, he was visiting professor at University of North Carolina, Chapel Hill, North Carolina and Michigan State University, East Lansing, Michigan, USA (1994).

He published 6 books (one in a French Publishing House) and he was collaborator to the "Encyclopedia of Chemistry" volumes 2, 3, 6 - 123 papers (54 in foreign journals). in stereochemistry, organic structural analysis and fine organic synthesis.

Professor Mager was Member of American Chemical Society (ACS) and he is member of Romanian Chemical Society. He was advisor for 12 PhD Thesis and member in 64 PhD. Juries. His research works are cited in many important publications and he was awarded with C. D. Nenitzescu Prize of the Romanian Academy (1992).

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SYNTHESIS AND STRUCTURAL ANALYSIS OF SOME NEW PRECURSORS OF MACROCYCLIC CYCLOPHANES: 1,4-BIS (2-R-5,5-DIMETHYL-1,3-DIOXAN-2-YL) BENZENE

NICULINA BOGDAN, FLAVIA POP, ANDREI SERBAN GÂZ, ELENA BOGDAN, ANAMARIA TEREC, LUMINIȚA DAVID AND ION GROSU^{*}

Dedicated to Professor Sorin Mager on the occasion of his 75th birthday

ABSTRACT. The synthesis and the stereochemistry of some new 1,3-dioxane derivatives of 1,4-diacetylbenzene are reported. The structural analysis carried out using NMR investigations revealed the anancomeric structure of the compounds. The favorable arrangement of the different functions for the participation to the macrocyclisation is insured by the axial orthogonal dispositions of the aromatic group with respect to both 1,3-dioxane rings.

Introduction

The investigations on the stereochemistry of 1,3-dioxane derivatives with aryl groups at position 2 of the heterocycle showed important structural aspects. The A value (free conformational enthalpy) of aryl groups at position 2 of the 1,3-dioxane ring are high (*e.g.* A_{Ph} = 13.04 kJ/mol [1,2]). The 2-aryl-1,3-dioxanes are anancomeric compounds and the conformational equilibrium between I and II is shifted toward I which exhibits the aryl group in the *equatorial* orientation (Scheme 1 [3-5]).

The conformational equilibrium of 2-alkyl,2-aryl-1,3-dioxanes (III, IV; *e.g.* 2-phenyl-2-methyl-1,3-dioxanes, Scheme 1) is shifted toward the conformer exhibiting the aryl group in the *axial* position [1,6-8].

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Scheme 1

The higher A value of the methyl group at position 2 ($A_{Me} = 16.63 \text{ kJ/mol}$ [1]) than the A value of the phenyl group ($A_{Ph} = 13.04 \text{ kJ/mol}$ [1]) at the same position suggests the *axial* preference of the phenyl group. Thermodynamic measurements of 2-methyl-2-phenyl-1,3-dioxanes showed a considerably higher preference of the methyl group for the *equatorial* position ($\Delta G_{III-IV}^0 = 10.11 \text{ kJ/mol}$ [1]) than that calculated from the A- values of methyl and phenyl groups ($\Delta A = A_{Me} - A_{Ph} = 3.63 \text{ kJ/mol}$).



Scheme 2

The equatorial aryl groups exhibit a weak preference for the *bisectional* rotamer (I_a) [9-11] and the axial phenyl group shows the *orthogonal* orientation (IV_b , Scheme 2 [7,8,12-15]). This is consistent with the ¹H NMR shifts [7,8].

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Investigations on the stereochemistry of compounds bearing two 1,3dioxane rings on the same aromatic system, obtained from benzenedicarboxaldehydes, have revealed the *equatorial* orientation of the aromatic ring for both heterocycles [3-5], while the investigations of similar derivatives of 1,4-diacetylbenzene [16] and of 2,6-diacetylbenzene [17] showed the axial orientation of the aromatic group with respect to both heterocycles. Some of these 1,3-dioxane derivatives were already successfully used in the synthesis of macrocyclic cyclophanes [18, 19]

It was considered of interest to investigate the stereochemistry of new 1,3-dioxane derivatives of 1,4-diacetylbenzene bearing different substituents at the positions 2 and 5 of the 1,3-dioxane rings and to determine the conformational behaviour of the heterocycles, the orientation of the aromatic substituent in both heterocycles, and to identify the possibility to use these compounds for the synthesis of cyclophanes.

Results and Discussions

In order to obtain new bis(2-R-1,3-dioxan-2-yl) derivatives bearing the 1,4-phenylene unit the 1,3-dioxane derivative **1** [16] was brominated in good yields (Table 1) using the typical procedure for the bromination reaction of cyclic ketals (Scheme 3)[20,21].

The bromination reaction generated as side product the tribrominated derivative **3** (Scheme 4, Table 1), too.



Scheme 3

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Scheme 4

Dibrominated derivative **2** was reacted with several nucleophyles (e.g. HO⁻, $C_6H_5COO^-$) in order to have access to other derivatives with appropriate substitution for the macrocyclisation reaction. All the attempts failed and the conclusion was that the steric hindrance of the carbon atoms bearing the bromine substituents is very high and nucleophyle substitutions cannot take place in these positions.

Due to this result we changed our strategy and we decided to obtain first the desired derivatives of 1,4-diacetylbenzene and then to use these derivatives for the synthesis of the 1,3-dioxane rings.

In the first step we brominated in good yields 1,4-diacetylbenzene with bromine in acetic acid [22] and after that we carried out the substitution of bromine atoms of the dibrominated derivative **4** with $C_6H_5COO^-$ (Scheme 5, Table 1), and we obtained the diester **5**.



Scheme 5.

The dibenzoate derivative **5** was subjected to the condensation reaction with neopentylglycol or with dimethyl-bis(hydroxymethyl)malonate (Scheme 6, Table 1) in order to obtain the 1,3-dioxane diester **6** and hexaester **7**. The diester **6** was deprotected with LiOH at 0° C, to give diol **8**.

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Scheme 6

Diol **8** was used to obtain the protected α , α '-dihidroxi-1,4-diacetylbenzene **10**, *via* the di-1,3-dioxane derivative **9** (Scheme 7, Table 1).



Scheme 7

Table 1. Results of the synthesis of compounds 2, 3 and 5-10

Compound	2	3	5	6	7	8	9	10
Yields %	75	15	92	52	74	91	78	86

Compounds 2, 3, 7, 9, 10 are new ones, while compounds 5, 6, 8 were already reported [19].

The structural analysis was carried out using NMR investigations. The spectra of **2**, **3**, **6-9** exhibit different signals for the axial and equatorial protons of the 1,3-dioxane rings and for the similar groups located on it (Table 2).

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Table 2. NMR data for compounds 2, 3 and 6-9

Compound	4'(4")-H;	6'(6")-H	5'(5")-CH ₃		5'(5")-COOCH ₃	
Compound	ax.	eq.	ax.	eq.	ax.	eq.
2	3.45	3.52	1.36	0.63	-	-
3	3.40	3.55	1.36	0.63	-	-
6	3.47	3.47	1.30	0.60	-	-
7	3.98	4.62	-	-	3.71	3.64
8	3.47	3.52	1.29	0.63	-	-
9	3.48	3.54	1.31	0.61	-	-

The NMR data suggest the anancomeric structure of these derivatives and the axial orthogonal orientation of the aromatic ring.

Conclusions

New 1,3-dioxane derivatives were obtained in good yields by condensation reactions or using the α , α ' bromination reaction of cyclic ketals. The NMR based structural analysis of these derivatives revealed the rigid structure for all compounds and the axial orientation of the aromatic ring with respect to both saturated heterocycles. The reported derivatives are important precursors for the synthesis of cyclophane derivatives.

Experimental

¹H-NMR (300 MHz) and ¹³C-NMR (75 MHz), COSY, HMQC, and HMBC spectra, were recorded at rt in CDCl₃ on a Bruker 300 MHz spectrometer, using the solvent line as reference. Mass spectra were recorded on an ATI Unicam Automass.

Melting points were measured with a Kleinfeld melting point apparatus and are uncorrected. Thin-layer chromatography (TLC) was conducted on silica gel 60 F_{254} TLC plates purchased from Merck. Preparative column (flash) chromatography was performed using PharmPrep 60 CC (40-63 μ m) silica gel purchased from Merck. Chemicals were purchased from Aldrich or Acros and were used without further purification.

General Procedure for the Synthesis of Compounds 2 and 3

The 1,3-dioxane derivative (2.1 mmol) and dry dichloromethane (40 ml) were introduced into a three-necked flask equipped with a reflux condenser, a thermometer and a dropping funnel. Bromide (9.25 mmol) in 10 ml dry dichloromethane was added drop wise, under magnetically stirring, to this mixture cooled into an ice bath at 0-5°C, the ensuing reaction being monitored initially by the fading of the solution color. After the addition of the bromide, the ice bath was removed and the stirring was continued for 1h,

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the contents in the flask being allowed slowly to reach room temperature (20-25°C). The mixture was evaporated *in vacuum* and the residue was crystallized from ethanol.

1,4-Bis(2'-bromomethyl-5',5'-dimethyl-1',3'-dioxan-2'-yl)benzene (2)

White solid, m.p.=164-5°C, yield 75% (flash chromatography, elution: DCM/pentane/diethyl acetate 1/3/0.1 v/v, Rf = 0.60). ¹H-NMR (300 MHz,CDCl₃, \bar{o} ppm): 0.63 (6H, s), 1.36 (6H, s), 3.45 (4H, d, *J*=10.6 Hz), 3.46 (4H, s), 3.52 (4H, d, *J*=10.6 Hz), 7.51 (4H, s). ¹³C-NMR (75 MHz, CDCl₃, \bar{o} ppm): 21.9 (CH₃) 23.0 (CH₃), 30.2 (C_{quat}), 40.8 (CH₂), 72.2 (CH₂), 98.9 (C_{quat}), 128.44 (CH), 137.8 (C_{quat}). **MS** (EI, 70 eV); m/z (rel. int., %): 493 (2) [M+H]⁺, 477 (10), 399 (100).

1-(2'-dibromomethylene-5',5'-dimethyl-1',3'-dioxan-2'-yl)-4-(2''-bromomethyl-5'',5''-dimethyl-1'',3''-dioxane-2''-yl)benzene (3)

White solid, m.p.=134-5°C, yield 15% (flash chromatography, elution: DCM/pentane : 3/1 v/v, Rf = 0.52). ¹**H-NMR** (300 MHz,CDCl₃, δ ppm): 0.63 (6H, s), 1.36 (6H, s), 3.40 (4H, d, *J*=10.9 Hz), 3.46 (2H, s), 3.55 (4H, d, *J*=10.9 Hz), 6.88 (1 H, s) 7.63 (2H, d, *J*=8.3 Hz) 8.16 (2H, d, *J*=8.3 Hz) ¹³**C-NMR** (75 MHz, CDCl₃, δ ppm): 21.8 (CH₃) 22.9 (CH₃), 30.2 (C_{quat}), 49.6 (CH), 40.1 (CH₂), 72.3 (CH₂), 98.6 (C_{quat}), 128.6 (CH), 130.4 (CH), 131.4 (C_{quat}) 143.9(C_{quat}). **MS** (EI, 70 eV); m/z (rel. int., %): 475,477 (2.5), 391,393 (49), 314 (12), 305 (14), 209 (11), 104

General Procedure for the Synthesis of Compound 7

2,2-dicarboxymethyl-1,3-propanediol (11 mmol) and ketone **5** (5 mmol) with catalytic amounts of *p*-toluenesulphonic acid (0.1 g) were solved in 200 mL toluene. The mixture was refluxed and the water produced in the reaction removed using a Dean-Stark trap. When 80 % of the water had been separated, the mixture was cooled to room temperature and the catalyst was neutralized (under stirring 0.5 h) with CH₃-COONa powder in excess (0.2 g). The reaction mixture was washed twice with 100 mL of water. After drying over MgSO₄, the benzene was removed and **7** was purified by flash chromatography.

1,4-Bis[2'-benzoyloxymethyl-5',5'-bis(carboxymethyl)-1,3-dioxan-2'-yl]benzene (7) White solid, m.p.= 245-6°C, yield 74% (flash chromatography, elution: DCM, Rf = 0.43. ¹**H-NMR** (300 MHz,CDCl₃, δ ppm): : 3.64 (6H, s), 3.71 (6H, s), 3.98 (4H, d, *J*=11.3 Hz) 4.31 (4H, s) 4.62 (4H, d, *J*=11.6 Hz) 7.42 (4H, dd, overllaped *J*=7.2 Hz, *J*=7.5 Hz) 7.54 (2H, t, *J*=7.5 Hz) 7.60 (4H, s,) 8.0 (4H, d, *J* = 7.2 Hz). ¹³**C-NMR** (75 MHz, CDCl₃, δ ppm): 53.16 (CH₃) 53.19 (CH₃), 53.23 (C_{quat}), 63.8 (CH₂), 69.2 (CH₂), 100.4 (C_{quat}) 128.4 (CH), 128.6 (CH), 129.9 (C_{quat}), 130.0 (CH), 133.2 (CH) 136.5 (C_{quat}), 165.9 (CO), 166.9 (CO), 168.1 (CO). **MS** (EI, 70 eV); m/z (rel. int., %): 616(100) [M-134], 442 (40).

General Procedure for the Synthesis of Compound 9

To a suspension of NaH (8.4 mmol) in dry THF (50 mL), diol **8** (2.1 mmol) in 10 ml THF was added under argon. The mixture was heated to reflux and benzyl bromide (2.1 mmol) solved in 5 mL THF was added slowly to the

NaH solution. Reflux of the solvent was continued for 5h, after which the system was brought to room temperature and the solid phase removed by filtration. The solvent was evaporated and the crude product was solved in CH_2CI_2 (20 mL) and then washed twice with water (20 mL). After drying over magnesium sulfate, the solvent was removed and the crude product was purified by crystallization from ethanol.

1,4-Bis(2'-benzyloxymethyl-5',5'-dimethyl-1,3-dioxan-2'-yl)benzene (9)

White solid, yield 76% ¹**H-NMR** (300 MHz,CDCl₃, δ ppm): 0.61 (6H, s), 1.31 (6H, s), 3.48 (4H, d, *J*=10.6 Hz) 3.54 (4H, d, *J*=10.6 Hz), 3.56 (4H, s), 4.54 (4H, s) 7.11-7.23 (10H, m, overlapped signals), 7.50 (4H, s). ¹³**C-NMR** (75 MHz, CDCl₃, δ ppm): 22.05 (CH₃), 30.4 (<u>C</u>H₃), 31.1 (C_{quat}), 71.6 (CH₂), 73.8 (CH₂), 77.25 (CH₂) 100.9 (C_{quat}), 127.38-127.42 (**CH**) 128.2 (CH), 132.3 (C_{quat} 1-C, 4-C), 137.5 (C_{quat}).

General Procedure for the Synthesis of Compound 10

To a stirred solution of **9** (6 mmol) in MeCN (3mL) and water (3mL) was added solid cerium ammonium nitrate (CAN, 18 mg, 4 mol %). The solution was heated at reflux for 2 h. After cooling at room temperature, water (10 mL) was added and the mixture was extracted with dichloromethane (3x20 mL). The organic layers were dried over MgSO₄, filtered and the solvent was evaporated *in vacuum*. The crude product was further purified by crystallization from acetone.

1,4-Bis(2'-benzyloxymethyl) benzene (10)

White solid, yield 86%. ¹**H-NMR** (300 MHz,CDCl₃, δ ppm): 4.68 (4H, s), 4.74 (4H, s) 7.31 -7.37 (10H, overlapped signals) 7.98 (4H, s).

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SYNTHESIS AND STRUCTURAL ANALYSIS OF SOME NEW PRECURSORS OF MACROCYCLIC CYCLOPHANES: 1,4-BIS (2-R-5-R¹-1,3-DIOXAN-2-YL) BENZENE DERIVATIVES

NICULINA BOGDAN, ANDREI SERBAN GÂZ, FLAVIA POP, ELENA BOGDAN, ANAMARIA TEREC, LUMINIȚA DAVID AND ION GROSU^{*}

Dedicated to Professor Sorin Mager on the occasion of his 75th birthday

ABSTRACT. The synthesis and the stereochemistry of some new 2,5-substituted-1,3dioxane derivatives of 1,4-diacetylbenzene are reported. The structural analysis carried out using NMR investigations revealed the anancomeric structure of the compounds and the *cis* or/and *trans* disposition of the substituents. These experiments showed the favorable arrangement of the different functions for the participation to the macrocyclisation reactions insured by the axial orthogonal disposition of the aromatic group with respect to both 1,3-dioxane rings.

Introduction

The studies on the conformational equilibriums of 2,2-disubstituted-1,3-dioxanes bearing alkyl and aryl groups in the ketal part of the heterocycle (I, II; e.g. 2-phenyl-2-methyl-1,3-dioxanes, Scheme 1) showed the shifting of these equilibriums toward the conformer exhibiting the aryl group in the *axial* position [1-3].

The *axial* preference of the phenyl group is suggested by the higher A value of the methyl group at position 2 (A_{Me} = 16.63 kJ/mol [1]) than the A value of the phenyl group (A_{Ph} = 13.04 kJ/mol [1]) at the same position. Thermodynamic measurements in 2-methyl-2-phenyl-1,3-dioxanes showed a considerably higher preference of the methyl group for the *equatorial* position (ΔG_{II-I}^0 = 10.11 kJ/mol [1]) than that calculated from the A-values of methyl and phenyl groups ($\Delta A = A_{Me} - A_{Ph} = 3.63$ kJ/mol).

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Scheme 1

Investigations on the stereochemistry of compounds bearing two 1,3-dioxane rings on the same aromatic system, obtained from benzenedicarboxaldehydes, have revealed the *equatorial* orientation of the aromatic ring for both heterocycles [4-6], while the investigations on the stereochemistry of derivatives of 1,4-diacetylbenzene [7] and of 2,6-diacetylpyridine [8] showed the axial orthogonal orientation of the aromatic ring with respect to both 1,3-dioxacyclohexane systems. The derivatives of these compounds bearing different substituents in positions 5' and 5" of the 1,3-dioxane rings show three diastereoisomers (*cis, cis, cis, trans, and trans, trans*) in agreement with the disposition of the groups with higher precedence at positions 5' (5") on the same side (*cis*) or on different sides (*trans*) with the aromatic group at position 2' or 2" (*e.g.* the derivatives of 1,4-diacetylbenzene and of 2,6-diacetylpyridine were already investigated and they were successfully used in the synthesis of macrocyclic cyclophanes [9, 10].



Scheme 2

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It was considered of interest to investigate the stereochemistry of new 1,3-dioxane derivatives of 1,4-diacetylbenzene (or of some α , α' derivatives) bearing different substituents at the positions 2 and 5 of the 1,3-dioxane rings and to determine the conformational behaviour of the heterocycles, the orientation of the aromatic substituent for both heterocycles, and to identify and to characterize the possible *cis* and *trans* isomers.

Results and Discussions

New 1,3-dioxane derivatives of 1,4-diacetylbenzene having nonsymmetric substitution at the positions 5' and 5" of the heterocycles were obtained in good yields by the condensation reaction of 1,4-dicetylbenzene with several 1,3-propanediol derivatives (Scheme 3, Table 1).



Scheme 3

The yields for the reactions with esters were higher and for all cases the raw products were mixtures of isomers: *cis-cis, cis-trans* and *trans-trans*. The ratios and the yields are presented in table 1. At position 2 we considered as reference the ester group, which is of highest precedence.

Table 1.

Results of the synthesis of compounds 5-8

Compound	5	6	7	8
Yields	45	49	53	36
Cis-cis/cis-trans/trans-trans	0/2/1	0/2/1	10 / 12 / 1	1/1.4/0

The *trans-trans* isomers of **5** and **6** and *cis-cis* isomer of **7** and **8** were separated by flash chromatography and were investigated as single compounds.

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The *trans-trans* isomer of the tetrol **9** was synthesized (yields = 94 %) using the LAH reduction (Scheme 4) of the separated *trans-trans* isomer of **5**.



Scheme 4

The NMR investigation showed the anancomeric structure of the compounds. In all investigated compounds the aromatic substituent occupies the axial orientation for both 1,3-dioxane rings, while the substituents at positions 5' and 5" are either equatorial or axial and generate the *cis* and *trans* configurations. The anancomeric structure of the compounds is proved by the recording in the NMR spectra of different signals for the axial and equatorial protons of the dioxacyclohexane rings (Table 2). The axial or equatorial orientation of the substituents at positions 5' and 5" was deduced from the values of the coupling constants. If the substituent at position 5 of the 1,3-dioxane ring is equatorial, the axial proton at the same position exhibits a large coupling constant with the axial protons of positions 4 and 6. The signal belonging to the axial protons at positions 4 and 6 is in this case a doublet of doublets with two large coupling constants (many times overlapped into a triplet; the second coupling with large constant is due to the geminal equatorial proton). If the substituent at position 5 of the 1,3-dioxane ring is axial, the proton of the position exhibits couplings with small constants with the protons of positions 4 and 6, including the axial ones. The pattern for the signal belonging to the axial protons of positions 4 and 6 shows a doublet of doublets with a large coupling constant (due to the equatorial protons of the same positions) and another one with small value due to the coupling with the vicinal equatorial proton at position 5.

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Table 2.

¹H NMR data (δ , ppm; J, Hz; CDCl₃) for compounds **4-8**

Compd	Isomer	Position of	4'(4")-H, 6'(6")-H (δ)		J		
Compu.	isoniei	R at 5'(5")	ax	eq	4(6)eq- 4(6)ax	4(6)ax-5ax	4(6)ax-5eq
5	trans- trans	3.61	3.84	4.20	11.7	11.7	-
6	trans- trans	1.18(t), 4.06(q)	3.84	4.19	11.7	11.7	-
7	cis-cis	3.95-3.96 (ov peak	verlapped	4.09	11.7	-	2.3
8	cis-cis	4.46	3.80 (overlapp	-3.92 ed peak)			
9	trans- trans	3.22	4.53	4.96	11.3	11.3	-

Conclusions

The high yields synthesis of some new *cis-cis* and *trans-trans* bis-1,3dioxane derivatives having as spacer between the two heterocycles an 1,4phenylene unit is reported. The structural analysis carried out by NMR investigations reveals the axial orientation of the aromatic ring with respect to both dioxacyclohexane units. These derivatives are precursors for the synthesis of simple or layered cyclophanes.

Experimental

¹H-NMR (300 MHz) and ¹³C-NMR (75 MHz), COSY and HMQC spectra, were recorded at *rt* in CDCl₃ or MeOD on a Bruker 300 MHz spectrometer, using the solvent line as reference. Mass spectra were recorded on an ATI Unicam Automass.

Melting points were measured with a Kleinfeld melting point apparatus and are uncorrected. Thin-layer chromatography (TLC) was conducted on silica gel 60 F_{254} TLC plates purchased from Merck. Preparative column (flash) chromatography was preformed using PharmPrep 60 CC (40-63 µm) silica gel purchased from Merck. Chemicals were purchased from Aldrich or Across and were used without further purification.

General Procedure for the Synthesis of Compounds 5-8

(5 mmol) diols **1-4** and (2 mmol) ketone (1,4-bis(2'-benzoyloxyacetyl)benzene) with catalytic amounts of *p*-toluenesulphonic acid (0.1 g) were solved in 100 mL CDCl₃. The mixture was refluxed and the water produced in the reaction removed using a modified Dean-Stark trap. When 80 % of the water had been separated, the mixture was cooled to room temperature and the catalyst was neutralized (under stirring 0.5 h) with CH_3 -COONa powder in excess (0.2 g). The reaction mixture was washed twice with 100 mL of water. After drying over MgSO₄, the solvent was removed and the crude products were purified by flash chromatography or by crystallization.

t-1,*t*-4-Bis(*r*-2'-benzoyloxymethyl- *t*-5' – carboxymethyl-1,3-dioxan-2'-yl)benzene (5 *trans-trans*)

White solid, m.p.=208-9°C, yield 24 % (flash chromatography, eluent: DCM, Rf = 0.24. ¹**H-NMR** (300 MHz,CDCl₃, δ ppm): 3.14 (2H, m), 3.61 (6H, s), 3.84 (4H, dd overlapped, *J*=*J*'=11.7 Hz), 4.20 (4H, dd, *J*=11.7 Hz, *J*'= 4.6 Hz), 4.36 (4H, s) 7.39 (4H, dd overlapped, *J*=8.7 Hz, *J*'=8.3 Hz), 7.54 (2H, t, *J*=8.7 Hz) 7.56 (4H, s), 7.95 (4H, d, *J*=8.3 Hz). ¹³**C-NMR** (75 MHz, CDCl₃, δ ppm): 40.0 (CH), 52.0 (CH₃), 62.1 (CH₂), 69.8 (CH₂), 100.0 (C_{quat}) 128.4 (CH), 128.7 (CH), 130.0 (C_{quat}), 129.8 (CH), 133.2 (CH) 136.8 (C_{quat}), 165.9 (CO), 170.3 (CO).

t-1, *t*-4-Bis(*r* -2'-benzoyloxyethyl-*t*-5'- carboxyethyl-1,3-dioxan-2'-yl)benzene (6 *trans-trans*)

White solid, m.p.=141-2°C, yield 26 % (flash chromatography, eluent: DCM, Rf = 0.27. ¹H-NMR (300 MHz,CDCl₃, δ ppm): 1.18 (6H, t, *J*=7.2 Hz) 3.11 (2H, m), 3.84 (4H, dd overlapped, *J*=*J*'=11.7 Hz), 4.06 (4H, q, *J*=7.2 Hz) 4.19 (4H, dd, *J*=11.7 Hz, *J*'= 4.5 Hz), 4.36 (4H, s), 7.39 (4H, dd overlapped , *J*=7.9 Hz, *J*'=7.2 Hz) 7.52 (2H, t, *J*=7.9Hz) 7.57 (4H, s), 7.95 (4H, d, *J*=7.2 Hz).

t-1,*t* -4- Bis(*r*- 2'-benzoyloxymethyl- *c*-5'-hydroxymethyl-1,3-dioxan-2'yl)benzene (7 *cis-cis*)

White solid, m.p.= 154-5°C, yield 17% (flash chromatography, eluent: DCM/ethanol = 2/0.1, Rf = 0.15. ¹H-NMR (300 MHz,CDCl₃, δ ppm): 1.47 (2H, dt *J*=7.2, *J*'=2.3), 1.78 (2H, t, *J*=5.3 Hz), 3.95-3.96 (8H, overlapped picks), 4.08 (4H, dd, *J*= 11.7 Hz, *J*= 4.5 Hz), 4.33 (4H, s), 7.41 (4H, t, *J*= 7.1 Hz) 7.52 (2H, dd overlapped), 7.99 (4H, d, *J*= 7.1 Hz). ¹³C-NMR (75 MHz, CDCl₃, δ ppm): 36.2 (CH), 62.05 (CH₂), 62.12 (CH₂), 70.12 (CH₂), 100.3 (C_{quat}), 128.5 (CH), 128.6(CH), 129.8 (CH), 129.9 (C_{quat}), 133.2 (CH), 137.0 (CH), 166.0 (CO).

t-1,*t* -4- Bis(*r*- 2'-benzoyloxymethyl- *c*-5'-tosyloxymethyl-1,3-dioxan-2'yl)benzene (8 *cis-cis*)

White solid, m.p.=213-4°C, yield 14% (flash chromatography, elution: diethyl ether/pentane = 1/2, Rf = 0.21. ¹**H-NMR** (300 MHz,CDCl₃, δ ppm):

1.69 (2H, m), 2.43 (6H, s),C 3.80-3.92 (8H, overlapped picks), 4.21 (4H, s), 4.46 (4H, d, *J*= 7.5 Hz), 7.31 (4H, d, *J*=8.3 Hz) 7.41 (4H, dd, *J*=8.3 Hz, *J*=7.2 Hz) 7.53 (4H, s) 7.55 (2H, t, *J*=7.2 Hz) 7.77 (4H, d, *J*=8.3 Hz) 7.97 (4H, dd, *J*=8.3 Hz) 13 **C**-**NMR** (75 MHz, CDCl₃, \bar{o} ppm): 21.8 (CH₃), 33.9 (CH), 61.4 (CH₂), 68.9 (CH₂), 69.9 (CH₂), 100.4 (C_{quat}), 128.0 (CH), 128.56(CH), 128.62(CH), 129.8 (CH), 130.1 (CH) 132.8 (C_{quat}), 133.2 (CH), 136.7 (C_{quat}), 145.1 (C_{quat}), 165.9 (CO).

General Procedure for the Synthesis of Compound 9

A mixture of **5** (1mmol) and (5 mmol) of LiAlH₄ in 50 mL of THF was stirred under reflux for 5 h. Aqueous NaOH (5%; 10 mL) was very slowly

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dropped into the stirred mixture at 0-5°C to destroy the excess LiAlH₄. After being stirred for 1 h, the mixture was filtered and washed several times with hot THF. The filtrate was evaporated, and the residue was purified by crystallisation from DCM.

t-1,*t*-4- Bis(*r*-2'-benzoyloxymethyl- *t*-5'-hydroxymethyl-1,3-dioxan-2'-yl)benzene (9 *trans-trans*)

White solid, m.p.= 218-9°C, yield 94%. ¹**H-NMR (**300 MHz, MeOD, δ ppm): 2.2-2.35 (2H, m), 3.22 (4H, d, *J*= 6.4 Hz), 3.46 (4H, s) 3.53(4H, dd, *J*=*J*'=11.3 Hz), 3.96 (4H, dd, *J*=11.3 Hz, *J*'=4.6 Hz), 7.49 (4H, s). ¹³**C-NMR (**75 MHz, MeOD, δ ppm): 40.0 (CH), 62.7 (CH₂), 66.0 (CH₂), 72.4 (CH₂), 103.6 (C_{quat}), 130.8 140.4 (C_{quat}). **MS** (70 eV) (rel. int. %) m/z: 366 (M⁺, 0.6), 179 (7), 147 (9), 117 (12), 89 (22), 71 (96), 59(100).

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REACTIONS OF THE 3-CYANO-10-METHYL-PYRIDO[3,2-g]QUINOLIN-4-ONE. II*

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Dedicated to professor Sorin Mager at his 75th anniversary

ABSTRACT. The synthesis of 4-alkoxy-3-cyano-10-methyl-pyrido[3,2-g]quinoline derivatives is described. The structural assignments were performed by high resolution 2D NMR spectroscopy.

Introduction

The reaction of 3-cyano-10-methyl-pyrido[3,2-g]quinoline-4-one **1** with alkylhalides under PTC conditions, in strong alkaline media was previously described [1]. In the presence of KOH or NaOH, the hydration reaction of the cabonyl bond generates a stable gemminaldiol as the main reaction product and the target alkylation products are minor reaction products under these experimental conditions (scheme 1). The mixture of N-alkyl- and O-alkyl-3-cyano-10-methyl-pyrido[3,2-g]quinoline derivatives contains the two alkylated derivatives in 3:1 ratio.



Scheme 1

^{*} Part I¹

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Another reaction strategy is thus required in order to obtain the O-alkyl-3-cyano-10methyl-pyrido[3,2-g]quinoline derivatives in good yields.

Results and Discussions

The two steps reaction path proposed for the regioselective synthesis of 4-alkoxy-3-cyano-10-methyl-pyrido[3,2-g]quinolines **3** is presented in scheme 2.



Scheme 2

As previously described, the chlorination of 3-cyano-10-methyl-pyrido[3,2-g]quinolin-4-one **1** with phosphorous oxychloride generated 4-chloro-3-cyano-10-methyl-pyrido[3,2-g]quinoline **2** in good yields [2]. The chlorine substituent in position 4 of the quinoline ring can be easily replaced by nucleophilic aromatic substitution. The reactivity of **2** is enhanced by the electron withdrawing effect of the nitrile group situated in position 3 of the heterocycle. For these reasons **2** was considered as an efficient intermediate for the synthesis of 4-alkoxy-3-cyano-10-methyl-pyrido[3,2-g]quinolines **3**.

The reaction of **2** with several alkoxides in the corresponding alcohols (such as *n*-butanol, *i*-butanol, amyl alcohol) were performed. The reaction conditions were summarized in table 1.

Table 1.

The	reaction	conditions	for	the	alkylation	of 2	with	alkovides
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Alkoxide	Reaction conditions		
	Temperature [°C]	「ime [h]	Yield [%]
Sodium n-butoxide	110	2	42%
Sodium <i>i-</i> butoxide	110	2	48%
Sodium n-pentoxide	130	2	30%

The structures of the reaction products **3** were assigned by high resolution NMR experiments. 2D-NMR spectra were used for the complete structural assignments, as follows:

- The ¹H-¹H homocorrelations that generate the splitting pattern of the protons in the structure **3** were observed by the proton 2D COSY with gradients experiment (parameter set COSY 45gs-BBI). Figure 1 presents a detail of this 400 MHz 2D COSY spectrum, showing the cross peaks determined by the homogenous spin-spin couplings in the aromatic region and the corresponding structural assignments.



Fig 1. 400 MHz 2D NMR COSY spectrum of 3 (detail of the aromatic region) and the corresponding structural assignments

The tertiary carbon atoms of the heterocycle **3** were assigned according to the inverse ¹H-¹³C correlation with z-gradients experiment (parameter set: HMQCgs-BBI). Figure 2 presents the 400 MHz 2D ¹H-¹³C hetercorrelation spectrum, and the corresponding structural assignments.

The quaternary carbon atoms were assigned according to the ¹H-¹³C inverse long range heterocorrelation with z-gradients experiment (parameter set: HMBCgs-BBI). In figure 3, the cross peaks in the HMBC spectrum show the ¹H-¹³C heterocorrelations through two and three covalent bonds that enabled the structural assignments.

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Fig 2. 400 MHz 2D NMR HMQC spectrum of 3 (detail of the aromatic region) and the corresponding structural assignments



Fig.3. 400 MHz 2D NMR HMBC spectrum of 3 (detail of the aromatic region) and the corresponding structural assignments.

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Conclusions

The reaction path proposed for the regioselective O-alkylation of 3cyano-10-methyl-pyrido[3,2-g]quinoline-4-one afforded 4-alkoxy-3-cyano-10methyl-pyrido[3,2-g] quinolines **3** in moderate yields.

Experimental

General procedure for 4-alkoxy-3-cyano-10-methyl-pyrido [3,2g]quinolines preparations.

7 mL anhydrous alcohol was placed in a round bottom flask, treated with 0.04 g clean sodium (2 mmol) and warmed gently until all the sodium has reacted. 4-Chloro-3-cyano-10-methyl-pyrido[3,2-g] quinoline 0,25 g (1 mmol) were added and the solution was refluxed for 2 hours. The solution was cooled, then filtered. The filtrate was poured in water. The precipitate thus formed was filtrate and the solution was extracted with ether and then the solvent was evaporated. The product, a yellow powder, was recrystallised from ethanol.

4-(1-butoxy)-3-cyano-10-methyl-pyrido[3,2-g]quinoline

IR $[cm^{-1}]$ 2850, 2900 (C-H stretching vibration), 2220 (C=N stretching vibration), 1603, 1500 (aromatic C=C stretching vibration), 1220 (C-O stretching vibration).

¹H-RMN (400 MHz): δ_{H} (DMSO-d₆ solution): 8.93 (dd, 1H, J = 4Hz, J = 1.2 Hz, H₈) 8.60 (s, 1H, H₅), 8.44 (dd, 1H, J = 8.4 Hz, J = 1.2 Hz, H₆), 8.34 (s, 1H, H₂), 7.38 (dd, 1H, J = 8.4 Hz, J = 4 Hz, H₇), 4.40 (t, 2H), 3.44 (m, 4H), 3.02 (s, 3H), 1.06 (t, 3H).

 13 C-RMN δ_{C} 176.45, 154.57, 151. 03, 148.37, 146.35, 138.13, 131.92, 128.07, 124.03, 122. 76, 122.17, 119.77, 87.92 (CN), 56.54, 19.06, 12.67.

4-(2-methyl-1-propoxy)-3-cyano-10-methyl-pyrido[3,2-g]quinoline

IR [cm⁻¹] 2215 (C<u>=</u>N stretching vibration), 1603, 1565 (aromatic C=C stretching vibration), 1276, 1047 (C-O stretching vibration).

¹H-RMN (400 MHz): δ_{H} (DMSO-d₆ solution): 8.93 (dd, 1H, J = 4Hz, J = 1.2 Hz, H₈) 8.60 (s, 1H, H₅), 8.44 (dd, 1H, J = 8.4 Hz, J = 1,2 Hz, H₆), 8.34 (s, 1H, H₂), 7.38 (dd, 1H, J = 8.4 Hz, J = 4 Hz, H₇), 4.39 (d, 2H), 3.43 (m, 1H), 3.02 (s, 3H), 1.07 (d, 6H).

 13 C-RMN $\delta_{\rm C}$ 176.47, 154.61, 151.02, 148.47, 146.35, 138.12, 131.99, 128.10, 124.03, 122.80, 122.15, 119.76, 87.90 (CN), 56.54, 19.05, 12.69.

4-(1-pentoxy)-3-cyano-10-methyl-pyrido[3,2-g]quinoline

¹H-RMN (400 MHz): δ_{H} (DMSO-d₆ solution): 8.91 (dd, 1H, H₈) 8.58 (s, 1H, H₅), 8.42 (dd, 1H, H₆), 8.34 (s, 1H, H₂), 7.38 (dd, 1H, H₇), 4.2 (t, 2H), 3.5 (m, 2H), 3.02 (s, 3H), 1.07 (m, 5H).

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MODELLING URANIUM(VI) EXTRACTION USING ARTIFICIAL NEURAL NETWORKS

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Dedicated to professor Sorin Mager at his 75th anniversary

ABSTRACT. The use of ANN simulators for the mathematical modelling of the uranium(VI) extraction from an aqueous solution, using polar and/or nonpolar solvent containing the chelating agent di-(2-ethylhexyl)-ditiophosphoric acid, at different pH values is described. The ANN has been trained with experimental data using either 1-butanol or kerosene solvent. Good prediction results show the incentives of this modelling approach and the perspectives of its using for simulating processes with the aim of optimising both laboratory and industrial units.

Introduction

Artificial Neural Networks (ANNs) are founded on an idealized representation of the biological cell that processes information. ANNs can be represented as weighted directed graphs consisting in simple elements, neurons (grouped in layers) operating in parallel, and the connections between neurons. The weighted connection paths link every two neurons from adjacent layers such as the weighting structure is able to provide the overall network performance. As the ability to learn is a fundamental trait of intelligence, the ANNs may be considered simple but powerful artificial intelligence elements. In the ANNs context the learning process consists in updating the network architecture and connection weights so that the network can efficiently achieve a specific task. Usually, the tasks an ANN may effectively perform are: pattern classification, clustering or categorization, function approximation, prediction and associative memory applications [1].

But the main benefits of the ANN approach consist in their remarkable ability for generalization. The use of ANN models for control purposes has gained considerable attention in the field of modelling chemical process, being the subject of several scientific reports, and they are increasingly applied for system identification and simulation [2]. As a consequence, the ANNs may be successfully used for modelling systems in which detailed

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governing rules are unknown or difficult to formalize under the form of first principle models [3, 4]. This last capability is exactly the type of application the present work is aimed to achieve.

The extraction of uranium(VI) from aqueous solutions with immiscible polar or nonpolar solvents using a dithiochelating agent was extensively studied. The choice of the proper solvent [5, 6] as well as the choise of the ligand [7, 8] and the optimal pH range were previously investigated.

This paper presents a first attempt to use Artificial Neural Networks for predicting the extraction of uranium(VI) from an aqueous solution, using mixtures of polar/nonpolar solvent of different ratio, in the presence of the chelating ligand di-(2-ethylhexyl)-ditiophosphoric acid. The chelate formation reaction is presented in scheme 1.



Scheme 1

Results and Discussions

The Artificial Neural Network employed in this study is of feedforward type and the backpropagation algorithm has been used for training the network's biases and weights. The network configuration is of multilayer structure and the quasi Newton Levenberg-Marquardt backpropagation algorithm has been used for learning.

The ANN architecture has been designed on the basis of heuristic methods. Its structure consists in an input layer of three neurons, a hidden layer of four neurons and a single-neuron output layer. The activation function of the first and second layer is of *tansig* (tan-sigmoid) form but the activation function function of the output neuron is *linear*.

First, the ANN has been trained using the following input-output (input-target) pairs of data:

Inputs: experimental data for the extraction of uranium(VI) from an aqueous solution having the uranium(VI) concentration of 0.1212 g/l, with either 1-butanol or kerosene solvent containing the chelatization ligand di-(2-ethylhexyl)-ditiophosphoric acid, at different pH values; these experimental data are presented in Table 1.

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 Outputs: experimental data consisting in the percentage values of uranium(VI) extraction obtained for the above mentioned inputs, also presented in Table 1.

Table 1.

Experimental data for the extraction of uranium(VI) from an aqueous solution $(C_{U(VI)}=0.1212 \text{ g/I})$, with either 1-butanol or kerosene solvent containing the chelatization ligand di-(2-ethylhexyl)-ditiophosphoric acid, at different pH values.

Sample no.	1-Butanol sc	olvent	Kerosene so	Kerosene solvent		
	Intputs	Outputs	Intputs	Outputs		
	pH	E [%]	pH	E [%]		
1	1.2	94.5				
2	2.0	96.4				
3	4.1	99.2				
4	4.7	99.8				
5	5.6	99.9				
6			1.05	13.4		
7			1.21	21.4		
8			2.0	55.4		
9			2.8	85.9		
10			4.1	87.8		
11			4.7	87.2		
12			5.6	85.8		

The ANN has been successfully trained. The simulation results produced by the trained network for modelling the percentage values of uranium(VI) extraction confirm this statement. The results (outputs) obtained by simulation for the already trained ANN, having as inputs the data presented in Table 1, are compared to the experimental outputs presented in the same table. The comparative results between experimental and ANN simulated data are shown in figure 1.

The relative errors between the ANN simulated data and the experimental data are considered to be small, as their offset is limited to ± 1.4 %.

Second, as one of the most appreciated properties of the ANNs is their ability to make predictions, this was also the main purpose of the presented investigation. The predictions consist of the ANN's aptitude to provide values of the outputs for the inputs not yet seen during the training step.

During this second testing step the already trained ANN has been used for predicting the values for the percentage extraction of uranium(VI) from the aqueous solution ($C_{U(VI)}=0.1212$ g/I) containing 1-butanol and kerosene solvents with concentrations of various ratios, having totally different values compared to the ones used during the training step.



Fig. 1.

Comparative results between ANN simulated data and experimental data, for the set of input-output data used for training the ANN (*learning*)

In order to test the prediction capability of the trained ANN a new set of inputoutput experimental data has been used, as they are presented in Table 2.

Table 2.

Experimental data for the extraction of uranium(VI) from an aqueous solution (C_{U(VI)}=0.1212 g/l), using various 1-butanol-kerosene solvent mixtures containing the chelating ligand di-(2-ethylhexyl)-ditiophosphoric acid, at the pH values 2.5-3

Sample no.	C _{butanol} [% vol. in kerosene]	E [%]
1	5	97.00
2	10	99.84
3	20	99.60
4	30	99.76

The values for the percentage extraction of the uranium(VI) obtained by ANN simulation are compared with the experimental values shown in Table 2 and the comparative results are presented in figure 2.

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Figure 2.

Comparative results between ANN simulated data and experimental data, for the set of input-output data used for ANN testing (*prediction*).

As it may be noticed from figure 2, the ANN has good predictions capabilities proved by the errors situated in a narrow interval of \pm 2.5%. Additionally it may be mentioned that testing has been performed on an input-output set of data not encountered during the ANN training procedure, and with the inputs having values very different to those used for training.

Conclusions

Although a relatively limited set of data has been used for training the results obtained on the basis of an ANN model for the simulation of the extraction of uranium(VI) from an aqueous solution ($C_{U(VI)}$ =0.1212 g/l), using various 1-butanol-kerosene solvent mixtures containing the chelating ligand di-(2-ethylhexyl)-ditiophosphoric acid are very promising, The simulator may be used for getting new sets of data for uranium(VI) extraction using 1-butanol-kerosene solvent mixtures, sparing experimental time and costs. Further improvement of the model may be performed by the use of a more extended training set of data.

The use of ANN simulators for the mathematical modelling of the uranium(VI) extraction from an aqueous solution with 1-butanol - kerosene solvent mixtures containing the chelating ligand di-(2-ethylhexyl)-ditiophosphoric

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acid may become a very useful tool for both laboratory and industrial scale optimisation of this investigated process.

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SYNTHESIS OF CIS - 7,8 – EPOXY - OCTADECANE, SPECIES -SPECIFIC COMPONENT OF THE SEX PHEROMONE OF NUN MOTH LYMANTRIA MONACHA (LEPIDOPTERA, LIMANTRIIDAE)

IOAN OPREAN^{a,b}, ANA AURELIA BOTAR^b, LUCIA GÂNSCĂ^b AND IULIANA VASIAN^b

ABSTRACT. The synthesis of *cis*-7,8-epoxy-octadecane, species-specific component of the sex pheromone of nun moth *Lymantria monacha* was based on a $C_{10}+C_2=C_{12}$ and $C_{12}+C_6=C_{18}$ coupling scheme, by acetylenic route, starting from 1-bromo-decane. The second coupling reaction took place between di-n-dodecyne mercury and 1-bromo–hexane in a transmetallation reaction.

Keywords: nun moth, Lymantria monacha, cis-7,8-epoxy-octadecane

Introduction

Nun moth (NM), *Lymantria monacha*, in Europe and gypsy moth (GM) *Lymantria dispar*, in Europe and North America are most important defoliators of coniferous and deciduous forests, respectively.



Figure 1

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For sexual communication GM and NM females produce the pheromone disparlure; NM and GM males respond to the (+) - disparlure enantiomer [(7R, 8S)-cis-7,8-epoxy-2- methyloctadecane] as a single attractive but unspecific pheromone component [1-29].

Three epoxies [(-) - disparlure, (+) - and - (-) - monachalure] and two hydrocarbon volatile components [2- methyl- Z7- octadecene and Z7octadecene] [30] synergistically prevented cross-attraction of coseasonal GM males and imparted specificity to NM sexual communication. Fig.1

In the literature are not mentioned the synthesis of cis-7,8-epoxyoctadecane, species-specific component of the sex pheromone of nun moth *Lymantria monacha*.

The paper presents a route for the synthesis of *cis*-7,8-epoxy-octadecane.

Results and Discussion

For the synthesis of *cis*-7,8-epoxy-octadecane (1) we explored the pathway outlined in Scheme 1, based on a $C_{10}+C_2=C_{12}$ and $C_{12}+C_6=C_{18}$ coupling scheme, by acetylenic route, starting from 1-bromo-decane.

In our variant, the first coupling reaction was effected in order to obtain 1-dodecyne (4) by adding 1-bromo-decane to monosodate acetylene, obtained in situ from acetylene in DMSO with sodium hydride. 1-Dodecyne (4) with mercuric chloride in an aqueous solution containing potassium iodide and sodium hydroxide (Nessler's reagent [31]) was quantitatively precipitated as di-n-dodecynyl mercury (5).

The key step in our acetylenic route consisted in transmetallation of didodecynyl mercury (5), which was directly lithiated by heating with lithium metal in diglyme and then alkylated with 1-bromo-hexane obtaining 7-octadecyne (7).

Z-7-Octadecene (8) was prepared from 7-octadecyne (7) by hydrogenation using Ni-P2/EDA catalyst, obtained by reducing nickel acetate with sodium borohydride and subsequent poisoning with ethylenediamine [32]. Using NiP-2 catalyst, the hydrogenation takes place quantitatively and with a very high degree of stereoselectivity.

To prepare *cis*-7,8-epoxy-octadecane (1) the olefine 8 was epoxidated with monoperphtalic acid.



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Scheme 1

Experimental

GC-MS analyses were performed on a Hewlett Packard GL-MSD 5890-5972 instrument, using a HP-5MS 30mX0.25mmX0.25µm capillary columm. A Perkin Elmer Spectrometer Model 700 was used for IR spectra.

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1-Dodecyne (4)

Aparatus: 1L reactor provided with a thermometer dipping into the liquid, a dropping funnel, a stirrer and a 20-30 cm long tube for preventing any splasting out of the reaction mixture.

All operations took place in inert atmosphere and under vigorous stirring.

4,41g (92 mmoles) of natrium hydride were added to 56 mL of dry DMSO. The temperature was risen up to $65^{\circ}-68^{\circ}$. The conversion was finished when no more hydrogen was evolved.

Acetylene (freed from acetone) was introduced for 15 min. at a rate of about 2L/min with external cooling in order to moderate the reaction.

During this process the temperature is maintained between 20° and 30° C.

28 mL of DMSO were added and afferwards, between 20^{0} - 30^{0} , 8,17 g (37 mmoles) of 1-bromo-decane with external cooling.

The reaction was led to completion by stirring for 3 hours at room temperature. The reaction mixture was poured into 100 mL of ice-water and extracted with four portions of 100 mL petroleum ether.

The combined extract were washed with brine and dried over MgSO₄. A satisfactory purity could be achieved by vacuum distillation collecting the fraction at $92^{0}-94^{0}C/28$ mmHg; yield: 6,14g (79%)

IR spectrum (film, cm⁻¹): 2160 s (-C≡C), 3280 vs (≡CH).

Mass spectrum (m/z; %): 151(<1), 137(<1), 123(1), 109(10), 95(40), 81(100), 67(77), 55(53), 41(85), 29(37).

Di-n-dodecynyl mercury (5)

18,4 g (110 mmoles) of 1-dodecyne (4) dissolved in 500 mL ethanol was added to a vigorous stirred solution of Nessler's reagent prepared according to the standard procedure [31].

The product 5 was obtained in yield of 93%, m.p.= $86^{\circ}-87^{\circ}$.

7-Octadecyne (7)

16,33 g (30 mmoles) of di-n-dodecynyl mercury (5) in 84 mL diglyme was treated with 0,44 g of lithium under inert atmosphere at $100-110^{\circ}$ for 2 hours.

11g (66 mmoles) of 1-bromo-hexane was added dropwise then the temperature raised to 120° - 130° under stirring for 6 hours. The mixture was left overnight, decanted, diluted with water and extracted with ethyl ether. The ethereal solution was washed to neutral and dried over MgSO₄.

After the removal of the solvent 12,1g (75%) of 7 were obtained, GC purity: 92%.

Mass spectrum (m/z; %): 250(<1), 235(<1), 221(<1), 207(<1), 180(1), 165(4), 152(2), 123(12), 109(29.9), 95(44.5), 81(53), 55(76.04), 67(54), 40(100), 29(24).

Z-7- Octadecene (8)

7-Octadecyne (7) was hydrogenated at Z-7-octadecene (8) in the presence of the NiP2/EDA catalyst, prepared in situ according to the standard procedure [32]. Z-7-octadecene (8) was obtained in yield of 93%, GC purity: 95%.

Mass spectrum (m/z, %): 252(10), 224(1), 196(<1), 182(<1), 168(1), 158(1.9), 125(11), 111(31.2), 97(59.8), 83(65.5), 69(73.4), 55(100), 41(99.5), 29(44.9).
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Z-7,8-Epoxy-octadecane (1)

7,56g (30 mmoles) of Z-7-octadecene (8) was trated with 68 mL of ethereal solution of monoperphtalic acid (1 mL containing 93 mg of peracid) and kept at room temperature for 24 hours. The mixture was then filtered, the Synthesis of *cis* - 7,8 - epoxY- octadecane filtrate washed successively with 10% NaHCO₃, diluted NaOH and water. After drying over MgSO₄, the solvent was removed and the residium was **1**. Yield: 95%, GC purity: 95%.

Mass spectrum (m/z, %): M=268(<1), 252(<1), 225(<1), 211(<1), 197(<1), 183(12), 152(2.5), 141(<1), 127(15), 97(55.9), 83(40.9), 69(59), 55(100), 41(84), 29(45.5).

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CONETORI OF HIGH GENERA

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Dedicated to Professor Sorin Mager, on his 75 years, for his bright contribution to the Stereochemistry.

ABSTRACT. Nanocones are taken as starting structures for designing various 3D nanostructures. Toroidal objects of genera from one to five are built up by using conical zones and appropriate nanotubes. In the same sense, units derived by opening the Platonic solids operated by various map operations are used. The topology of the resulting objects is discussed in connection with the strain estimated at the POAV1 level of theory.

1. Introduction

A map *M* is a combinatorial representation of a closed surface.¹ Several transformations (*i.e.*, operations) on maps are known and used for various purposes.^{2,3}

Let us denote in a map: *v* - number of vertices, *e* - number of edges, *f* - number of faces and *d* - vertex degree.

Recall some basic relations in a map:⁴

$$\sum d v_d = 2e \tag{2}$$

$$\sum s f_s = 2e \tag{3}$$

where v_d and f_s are the number of vertices of degree *d* and number of *s*-gonal faces, respectively. The two relations are joined in the famous Euler⁵ formula:

$$v - e + f = \chi(M) = 2(1 - g)$$
 (4)

with χ being the Euler *characteristic* and *g* the genus⁶ of a graph (*i.e.*, the number of handles attached to the sphere to make it homeomorphic to the surface on which the given graph is embedded; *g* = 0 for a planar graph and 1 for a toroidal graph). Positive/negative χ values indicate positive/negative curvature of a lattice.

If a graphene sheet is divided into six sectors, each with an angle of 60° (Figure 1), and if *m* of these sectors (with *m* varying from 1 to 5) are

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selected sequentially with the dangling bonds being fused together (see below), a series of five single-walled nanocones is obtained, with a linear angle α at the cone apex equal to 112.9°, 83.6°, 60.0°, 38.9°, and 19.2°. These values correspond to the formula:⁷

$$\alpha = 2 \arcsin(m/6) \tag{1}$$

One can add two extreme cases: (i) the graphene sheet, with all m = 6 sectors being involved, corresponding to a "cone" with an angle of 180°; and (ii) when m = 6, one obtains a "cone angle" equal to zero, corresponding to a nanotube capped at one end with any combination of hexagons and six pentagons, e.g., a "half-buckminsterfullerene". Thus *nanotubes capped at one end can be considered to be a particular case of nanocones,* and indeed Ebbesen has observed by transmission electron microscopy how a blunt nanocone with m = 5 on adding a sixth 60° sector becomes converted into a nanotube.^{8,9}



Figure 1. The six sectors on a honeycomb lattice

In the hereafter text we will consider only cones ending in polygons of s = 3 to 5 with no other polygonal defect of the graphite sheet. The name of such objects includes: [tip polygon]CN_length of the cone body, in number of hexagon rows (Figure 2).







Figure 2. A nanocone ending in a trigon

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2. Single Conetori

Conical zones may be involved in the construction of the so-called "distinct-walled tori" DWT, proposed by Nagy *et al.*¹⁰ to appear by sealing a double-walled carbon nanotube DWNT in two distinct position, by an electron beam. Proposals of toroidal structures bearing polygonal defects are known since the pioneering times of nanoscience.¹¹⁻¹³

Such structures, of genus 1, called hereafter conetori, are built up, in general, by two conical zones joined by two tubes, one internal and the other external, having corresponding distinct diameters. Their name includes: [cone tip polygon]CT_cone length_type and length of internal tube(tiling)_type and length of external tube(tiling), the length being given in number of hexagon rows. The tube is either an H (zig-zag) or V (armchair) one.¹⁴

Function of the composing parts the conetori are classified as follows.

2.2.1. Conetori with Internal H-tube, External V-tube

Even the small conetori (Figure 5) of this type are strained structures, it could be imagined that by increasing the number of atoms, they could become more relaxed molecules, as suggested in Figure 6. This huge object can be covered by disjoint coronenes (by the generalized map operation $(2,2)^{15}$); it appears the opportunity to address the question of aromaticity of such total resonant molecular systems and consequently the possible increase of molecule stability.



[5]CT 2 H0(7) V4(5,7,5)

[5]CT_2_H1(7)_V5(5,7,5) v =370; S=5.05



Figure 3. Conetori with internal H-tube, external V-tube

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Figure 4. Conetorus with disjoint coronenic units

2.2.2. Conetori with Internal V-tube, External V-tube

Cones of this type show (5,8,5) tiling at the junction with the inner tube while (5,7,5) at that with the outer tube (Figure 5a). The octagons can be eliminated by a Stone-Wales operation (*i.e.*, edge rotation, specified by RO in Figure 5b). Systematic search on the stability of tori belonging to this type has been reported in ref.¹⁰

(a) [5]CT_2_V2(5,8,5)_V5(5,7,5) V = 420; S=3.64



(b) [5]CT_2_V3(6,7)RO_V5(5,7,5) V = 420; S=3.61



Figure 5. Conetori with internal V-tube, external V- tube

3. Multi Tori

The torus of g = 1 is the simplest one in the series of objects of high genera. Such structures can be generated by opening the Platonic solids, operated by some map operations. In the following some of the most interesting classes of multi tori are illustrated and characterized from the topological point of view.

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3.1. Tori of g =2 and g =3

By intersecting two simple tori in one or two points, multi tori of g=2 and g=3, respectively, can be constructed (Figure 6).

Systematic construction of such multi tori is based on the Platonic solids: they are operated by some map operations, *e.g.*, quadrupling *Q*, capra *Ca*, etc., and next every original face opened. In this way, repeat units in possible infinite lattices or finite multi tori, by joining every pair of open faces by an appropriate nanotube segment, can be constructed. The objects in Figure 7 originate in the tetrahedron T while those in Figures 8 to 11 in the cube C.

DT; *v*=936; *e*=1404; *f*₇=12; *f*₆=454; *g*=2; S=5.81







Figure 6. Multi tori of genera g=2 and g=3

DT(T); v = 124; e = 186 $f_7=12$; $f_6=48$; g = 2; S = 14.44







Figure 7. Double torus of g=2, and its core derived from the tetrahedron T

Observe the objects in Figure 8 are isomers having the open faces of the cube ended as the "armchair" - V - tube (a) and "zig-zag" - H - tube (b). Clearly, the attached tubes have to have the same open ends. Also note the very low strain in the (b) object, which is further observed in the derived triple torus (Figure 9b, compared with Figure 9a). The both objects 43 in Figure 8 were inferred in the molecular realization of the celebrate Dyck graph,¹⁶⁻¹⁸ on 32 vertices of valence 3, 48 edges, 12 octagons, girth 6, diameter 5, and chromatic number 2. It is non-planar and has the genus g = 1 (*i.e.*, there exists an embedding of the graph on the torus).¹⁷ Cycle counting revealed 12 octagons and 16 hexagons.



Figure 8. Core units for m*ulti tori of high genera*, derived from the cube C



Figure 9. *Triple tori*, *g*=3 with the core derived from the cube C

The strain S, in terms of the POAV1 theory, $^{19-22}$ of such objects of high genera¹⁸ is relaxed as the number of atoms increases. It is evident, when compare the structures in Figures 10 (320 atoms) and 11 (2240 atoms), with a clear drop in their strain.

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Figure 10. *Triple torus*, *g*=3 and its core derived from the cube C

Ca_S(TTH_CA_R); v = 2240; e = 3360; $f_7 = 24$; $f_6 = 1092$; g = 3; S = 2.54



Figure 11. A more relaxed t*riple torus*, *g*=3, by Capra operation

3.2. Structures of High Genera

Structures of high genera can be generated, as above mentioned, by opening the Platonic solids, operated by some map operations. Units derived as above may form either infinite lattices of negative curvature²³⁻²⁶ or closed cages, showing porous structure. Spongy carbons have been recently synthesized.²⁷ Objects like those in Figure 12, of genus 5, can be built up, on the unit derived by opening the chamfering/quadrupling transform of cube (see Figure 8), by closing together six cones emerging from the open faces. They are isomers comprising either a V or an H tubular inner unit.

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Objects of high genera have also been modeled by Lenosky et al.,^{23,24} Terrones *et al.*^{25,26} and more recently by Lijnen and Ceulemans.¹⁸ Structures have been optimized at the Amber MM HyperChem level

(Polak-Ribiere conjugate gradient, at RMS = 0.005) and the strain computed by the JSChem software program.²⁸

Conclusions

Conical zones are involved in several nanostructures, starting from the intersecting units of nanotubes to finite or infinite structures. Multi tori appear in zeolites or in spongy (carbon) structures, as parts of the infinite P- or FRD-type negatively curved surfaces. The toroidal objects including conical zones could appear as real molecular structures, as the nanotubes filled by spherical fullerenes turn in double walled nanotubes, the sealing of which enabling (elongated) toroidal structures with distinct walls.

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RETRO MAP OPERATIONS

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ABSTRACT. Operations on maps are well known theoretical tools for transforming a given polyhedral tessellation. Several properties of fullerenes, such as their π -electronic structure and stability, need information on their associate graph, which eventually resulted by a map operation applied on a smaller molecular graph. In this respect, retrooperations, particularly those of the most used leapfrog, chamfering and capra operations, appear particularly useful in searching the properties of fullerenes. A series of analyzed fullerenes proved to be leapfrog transforms of smaller cages. This helped in understanding their closed π -electronic structure and stability.

Introduction

Fullerenes are carbon allotropes, with finite cage associate graphs, that already entered in the realm of real chemistry: they have been functionalized or inserted in supramolecular assemblies.¹⁻⁷

A fullerene is, according to a classical definition, an all-carbon molecule consisting entirely of pentagons (exactly 12) and hexagons (N/2-10). Non-classical fullerene extensions may include rings of other sizes.⁸⁻¹⁰

A map M is a combinatorial representation of a closed surface.¹¹ Several transformations (*i.e.*, operations) on maps are known and used for various purposes.

Operations on maps are topological-geometrical transformations enabling modification of a polyhedral tessellation. Basic simple map operations, such as dualization, truncation, stellation, etc., are supposed known. About this subject, the reader is referred to. ¹¹⁻¹³

Let us recall some basic relations in a map:¹⁴

$$\sum d v_d = 2e \tag{1}$$

$$\sum s f_s = 2e \tag{2}$$

In the above, *v* is the number of vertices, *e* - number of edges, *f* - number of faces, *d* - vertex degree, v_d - number of vertices of degree *d* and f_s - number of *s*-gonal faces, respectively. The two relations are joined in the famous Euler formula:¹⁵

$$v - e + f = \chi(M) = 2(1 - g)$$
 (3)

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with χ being the Euler *characteristic* and *g* the genus¹⁶ of a graph (*i.e.*, the number of handles attached to the sphere to make it homeomorphic to the surface on which the given graph is embedded; *g* = 0 for a planar graph and 1 for a toroidal graph).

This paper presents three of the most used map operations and their retro-pairs and their possible use in investigating π -electronic structure and stability of some fullerenes.

Leapfrog

Leapfrog Le is a composite operation, ^{12,17-20} which can be written as:

$$Le(M) = Du(P_3(M)) = Tr(Du(M))$$
(4)

A sequence of stellation-dualization P_3 -Du rotates the parent *s*-gonal faces by π/s . Leapfrog operation is illustrated, for a pentagonal face, in Figure 1.



Figure 1. *Le* operation on a pentagonal face

The map transformed parameters are as follows:

Le(M):
$$v = s_0 f_0 = d_0 v_0; e = 3e_0; f = v_0 + f_0$$
 (5)

Retro-leapfrog *RLe* operation is based on the following sequence:

$$RLe = -P_3(Du(Le(M)))$$
(6)

performed by cutting all vertices in the dual (of leapfrogged map) with degree lower than the maximal one (Figure 2). As a 3D realization, *RLe* is illustrated in Figure 3.

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Figure 2. Retro-Leapfrog RLe operation on a pentagonal face

Le (M); v = 48

Du(Le(M)); v = 26



v = 20

M = Cubeoctahedron; v = 12





Figure 3. Retro-Leapfrog operation

Quadrupling

Quadrupling Q is another composite operation, achieved by the sequence: 12

$$Q(M) = -E(Tr_{P_3}(P_3(M)))$$
(7)

where -*E* means the (old) edge deletion (dashed lines, in Figure 4) of the truncation Tr_{P3} of each central vertex of P_3 capping.

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The complete transformed parameters are:

Q(M):
$$v = (d_0 + 1)v_0; e = 4e_0; f = f_0 + e_0$$
 (8)

Q operation leaves unchanged the initial orientation of the polygonal faces.



Figure 4. Q operation on a pentagonal face

Retro-quadrupling *RQ* operation is based on the sequence:

$$RQ(M) = E(-Tr_{P_3}(P_3(M)))$$
(9)

performed by adding new edges parallel to the boundary edges of the parent faces (Figure 5) and deletion of these faces. As a 3D realization, RQ is illustrated in Figure 6.



Figure 5. Retro-Quadrupling *RQ* operation on a pentagonal face





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Figure 6. Retro-Quadrupling operation

Capra

Capra Ca - the goat, is the Romanian corresponding of the *leapfrog* English children game. It is a composite operation, ^{13,21,22} necessarily coming from the Goldberg's²³ multiplying factor m:

$$m = (a^{2} + ab + b^{2}); a \ge b; a + b > 0$$
(10)

that predicts *m* (in a 3-valent map) as follows: *Le*, (1, 1), *m*= 3; *Q*, (2, 0), *m*= 4; *Ca*, (2, 1), *m*= 7. By this reason, *Le* is also called *tripling* and *Ca septupling*. *Q* was originally called *chamfering*.²³

Capra is achieved by truncating the vertex located in the center of parent faces of a pentangulation P_5 transform (Figure 7). Note that, P_5 involves an E_2 (*i.e.*, edge trisection). This operation results in a map that preserves the original vertices while the parent *s*-gonal faces are twisted by $\pi/(3/2)s$.

The transformation can be written as:

$$Ca(M) = Tr_{P_5}(P_5(M))$$
 (11)

with Tr_{P5} meaning the truncation of each vertex where P_5 capping faces of a parent face are incident. *Ca* insulates any face of *M* by its own hexagons, which are not shared with any old face (in contrast to *Le* or *Q*).

The complete transformed parameters by Ca are:

Ca(M):
$$v_1 = v_0 + 2e_0 + s_0 f_0 = (2d_0 + 1)v_0; \ e_1 = 7e_0; \ f_1 = s_0 f_0 + f_0$$
 (12)

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Figure 7. *Ca* operation on a pentagonal face

Retro-capra *RCa* operation is achieved by the sequence:

$$RCa(M) = -E_2(-Tr_{P5}(M))$$
 (9)

In words, delete the smallest faces of the actual map and continue with - E_2 (Figure 8). As a 3D realization, *RCa* is illustrated in Figure 9.



Figure 8. Retro-Capra R*Ca* operation on a pentagonal face

 C_{140}

- $Tr_{P_5}(D); N = 80$





Figure 9. Retro-Capra operation

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Discussion

The properly closed PC shell *fz*-tubulenes²⁴ (*i.e.*, cages having fullerene halves as caps for a "zig-zag" tube) obey a true leapfrog rule, but written for zig-zag cylinders, Z-LER: N = 13k + 3km; k = 4,6,8,...; m = 1,2,...

A closed structure of π -electronic shell was found²⁵ to be a general property of cages derived by *Le* operation from smaller objects. We just confirmed this hypothesis by drawing the parent small cages (with non-trivalent atoms corresponding to the polar rings different from hexagon) by the retro *RLe* operation (Figure 10).

The closed shell π -electronic structure of these cages is reflected in the HOMO-LUMO gap, in both simple Hűckel level of theory and PM3 calculations (Table). The heat of formation HF values of the cages with the polar ring between k = 6 and 10 are pretty low (note that the PM3 energy for C₆₀ is about 13.5 kcal/mol).

Observe the Hückel gaps close in LUMO being NBO (for $0 \pmod{(k,4)}$ cages, Table, entries 1 and 3). Also note the cap $C_{13k/2(k(56)^{k/2}-2[3k,0])}$; k = 6, is C_{60} deducible, by taking a hexagon as the polar ring. The subscripts name is given in terms of the spiral code.²⁴

The strain energy SE, in terms of POAV1 theory,²⁶⁻²⁹ decreases as the cage size increases (Table). For comparison, SE for C_{60} is about 8.26 kcal/mol.



Figure 10. Fullerenes of general formula $\ C_{_{N(k\,(56)^{k/2}-Z[3k,0])}}$

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Table.

				PM3		SE	Hűckel [Data		
	Cage	V	Sym.	HF/at.	Gap	POAV1	HOMO	LUMO	Gap	Shell
	k			(kcal/mol)	(eV)	(kcal/mol)			(β	
									units)	
1	4	52	D_{2h}	21.585	5.533	12.526	0.256	0	0.256	PC
2	6	78	C_{3h}	12.294	6.083	6.794	0.516	-0.118	0.634	PC
3	8	104	C_{4h}	11.453	5.730	5.218	0.256	0	0.256	PC
4	10	130	C_{5h}	12.518	5.998	4.999	0.458	-0.087	0.545	PC

Data for <code>fz-tubulenes C</code> $_{\mathit{N(k(56)^{k/2}-Z[3k,0])}}$

It is quite strange that, for the cage with k=8, both the heat of formation HF (Table, row 3) and strain energy SE (in POAV1 terms, Table 2, row 3, column 2) data show lower values than for the cage with k=6, suggesting a higher stability for the former. However, the HOMO-LUMO gap for the cage with k=8 is lower than for the cage with k=6, both at Hűckel and PM3 level of theory (Table), indicating a kinetic instability, which is supplementary supported by the non-bonding character of the LUMO orbital, in Hűckel theory.

Conclusions

Operations on maps and their retro-transformations, particularly those of the most used leapfrog, chamfering and capra operations, proved to be useful in investigating the properties of fullerenes, particularly their π -electronic structure and stability. Thus, a series of Z-LER cages proved to be leapfrog transforms of smaller cages. Among these, the cage with k = 6, was described as the most stable among the discussed structures, which indeed, corresponds to the isolated fullerene C₇₈.

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PSEUDOCERAMIDES AND THEIR DERIVATIVES. 3. BENZYLIDENE ACETALS OF N-ACETYL-N-METHYL GLUCAMINE – PRELIMINARY STUDY.

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ABSTRACT. Acetalization of N-acetyl-N-methyl-1-amino-1-deoxy-D-glucitol with 2.5 equivalents of 1,1-dimethoxybenzaldehide in N,N-dimethylformamide at reflux temperature with camphorsulphonic acid as catalyst yielded 5,6-O-benzylidene-N-acetyl-N-methyl-glucamine as the major product. 2,3:4,6-di-O-benzylidene-N-acetyl-N-methyl-glucamine, 4,6-O-benzylidene-N-acetyl-N-methyl-glucamine and a mixture of 2,3:5,6- and 3,4:5,6-di-O-benzylidene-N-acetyl-N-methyl-glucamine byproducts have been also isolated. Separation and purification conditions of these new products are described. The structures have been assigned by ¹³C-NMR and MS-EI.

Keywords: aminoalditol acetals, pseudoceramide, flash chromatography, 5,6-O-benzylidene-N-acetyl-N-methyl-glucamine, 2,3:4,6-di-O-benzylidene-N-acetyl-N-methyl-glucamine, 4,6-O-benzylidene-N-acetyl-N-methyl-glucamine, 3,4:5,6-di-O-benzylidene-N-acetyl-N-methyl-glucamine, 3,4:5,6-di-O-benzylidene-N-acetyl-N-methyl-glucamine.

Introduction

The development of new biocompatible and biodegradable surfactants based on natural feedstocks is subject of an increasing scientific and industrial interest [1]. Aminoalditol derivatives represent a class of compounds that can be used both as surfactants as well as pseudoceramides [2, 3]. The use of Nmethyl-glucamine (an accessible and cheap aminoalditol) allows two strategies in order to obtain a compound with amphiphatic character that could exhibit ceramide properties:

a) attachment of a nonpolar tail to the amine function followed by the attachment of a second nonpolar tail to the alcohol function

b) creation of an amide bond at the amine function (by acylation) followed by the attachment of two nonpolar tails at the alcohol functions.

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These two options can be realized with biocatalysts, using the catalytic capacity and the specificity of lipases [4, 5]. A new approach in the synthesis of these compounds starts from N-acetyl-N-methyl glucamine and consists in the enzymatic attachment of fatty acids by esterification or transesterification reactions catalyzed by lipases [6]. The reaction monitorization can easily performed by HPLC or GC, whether mono- and diesters of N-acetyl-N-methyl-glucamine are available as standard compounds. The selective esterification of each alcohol function in the aminoalditol requires acetal derivatives as intermediates. This paper presents the results of acetalization of N-acetyl-N-methyl-1-amino-1-deoxy-D-glucitol with dimethoxy benzaldehyde in acid catalysis.

Results and Discussion

The condensation of aldehydes and ketones with alcohols is one of the first reactions studied in organic chemistry. E. Fischer obtained in 1895 for the first time, fructose acetals and then glucose acetals, in acid catalysis [7]. Acetals can be obtained in acidic, basic or neutral conditions or using other methods [8]. A series of well documented reviews present results of the acetalization reaction for alditols [9], aldoses, aldosides [10,11] and ketoses [12] but not of aminoalditols.

The acetalization of N-acetyl-N-methyl-glucamine was performed by transacetalization reaction in acidic media using dimethoxy benzaldehyde (see the experimental part). The reaction was monitorized by TLC.and the inspection of the plate showed 5 spots at the following Rf values: spot 1 (0.73), spot 2 (0.67), spot 3 (0.55), spot 4 (0.24) and spot 5 (0.15) (AcOEt :MeOH = 90:10).

Considering the TLC plate polarity and the eluent polarity we used, the spots on the TLC plate can be assigned as follows: spots 1, 2 and 3 for dibenzylidene derivatives and spots 4 and 5 for mono-benzylidene derivatives. After separation by flash chromatography, the compounds have been analyzed by MS-EI. Spots 1, 2 and 3 are di-benzylidene derivatives and have the molecular peak at m/z= $413(M^{+})$ for a 22 eV ionization energy, thus confirming the proposed assignments. Spot 1 was characterized as acetylated derivative, having the molecular peak at m/z= $455(M^{+})$, 24 eV.

The analysis of ¹H-NMR spectra is difficult because the 8 protons in the linear side chain are localized in the 3.5-4.3 ppm interval and the coexistence of cis-trans conformers of the amide bond (Figure 1) determines the doubling of the total number of signals.

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Figure 1.

Cis-trans conformers of N-acetyl-N-metyl-glucamine and derivatives

The ¹³C-NMR spectroscopy offers valuable data concerning the nature of the acetal rings as well as their positions [13, 14]. Buchanan and coworkers [15, 16] observed that the chemical shift of the acetal carbon atom is strongly influenced by the size of the ring and is situated in the following intervals:

108.1-115.7 ppm for 1,3-dioxolane rings

97.1-101.1 ppm for 1,3-dioxane rings

100.8-102.3 ppm for 1,3-dioxepane rings

The ¹³C-NMR spectra analysis of spot 1 show 8 signals in the169.4 - 171.2 ppm interval, which indicate the existence of 2 isomers, each one with 2 conformers. The 4 signals situated in the 103.7 – 104.2 ppm interval correspond to 4 acetalic C atoms from 1,3-dioxolanic rings and belong to 2 dibenzylidene isomers, both with 2 conformers. These chemical shifts are deshielded (they are present at lower ppm values: 103.7 - 104.2 ppm) comparing with literature data (108.1 - 115.7 ppm for 3-dioxolanic rings [15, 16]) because of the influence of Ph. Hereby the 2 isomers from spot 1 can be only the compounds **2** and **3** from Figure 2: 2,3:5,6-di-O-benzylidene-N-acetyl-N-methyl-glucosamine and 3,4:5,6-di-O-benzylidene-N-acetyl-N-methyl-glucosamine. Assigning position 5,6 to one of the benzylidene groups will be explained below.

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Figure 2.

Mono- and diacetal reaction products

¹³C-NMR spectra analysis of spot 2 relieves 4 signals:

- at 103.7 and 104.5 ppm the signals correspond to an acetalic C from 1,3-dioxolane ring (the signals are shifted to a higher value of the magnetic field because of the influence of the phenyl group comparing with the literature data [15, 16])

- at 100.7 and 101.0 ppm the signals correspond to an acetalic C from 1,3-dioxane ring.

Admitting that the initial and kinetically more stable 5, 6-benzylidene derivative is stabilized during time to the thermodynamically more stable 4,6-benzylidene isomer, results that spot 2 can be assigned solely to compound **4a** (Figure 2) namely 2,3:4,6-di-O-benzylidene-N-acetyl-N-methyl-glucamine. The 3,5:4,6 and 2,5:4,6 isomers are excluded because the chemical shifts of acetalic C at 105.0 and 104.1 ppm are clearly assigned to a dioxolanic ring.

Although isopropyliden and cyclohexyliden aminoalditol acetals have been already described [20, 21] there are no data about benzylidene acetals, consequently the position of benzylidene group had to be established. The analysis of ¹H-NMR spectra is difficult and doesn't give any valuable information. To spread the hydrogen atom attached on carbon atom with free hydroxyl groups from compound of spot 4 the free hydroxyl groups were derivatized by acetilation and than analysed by ¹H-NMR.

The type of ring was deduced from chemical shifts of the acetalic C atom from the ¹³C-NMR spectra. There are 2 signals at 103.5 ppm and 103.9 ppm which indicate a 1,3-dioxolane ring and thereby the spot 4 correspond to compound **1** (Figure 2): 5,6-O-benzyliden-N-acetyl-N-methyl-glucamine. The position of the 1,3-dioxolane ring on the linear side chain was determined by NMR experiments: monodimesional APT and bidimensional COSY and HETCOR. The assignment of C atoms from the linear side chain was established by HETCOR and APT experiments only for C1 and C6.

The presence of cis-trans conformers was proved previously by our group [20, 21]. Obtaining mainly 5,6-O-acetals of N-acetyl-N-methyl-glucamine 62

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is in agreement with the literature concerning kinetic acetalization of diethyldithioacetals of glucose, galactose and arabinose [17-19] and for isopropyliden and cyclohexyliden acetals of N-acetyl-N-methyl-glucosamine [20, 21].

The ¹³C-NMR spectra analysis of spot 5 relieve the existence of 1,3dioxane ring (100.4 and 100.5 ppm) and it was assigned to compound **4** from Figure 2 (4, 6-benzilidene isomer of N-acetyl-N-methyl-glucamine). The benzylation of compound **1** (spot 4) (NaH, BnBr) followed by hydrolysis of benzyliden acetal (I_2 /MeOH) leads finally to 2,3,4-tri-O-benzyl-N-acetyl-N-methylglucozamine. This compound was also obtained from 5,6-isopropilydene-Nacetyl-N-methyl-glucozamine [21] and from 5,6-cyclohexylidene-N-acetyl-Nmethyl-glucozamine[22], their structure were previously established [21]. These compounds were obtained by the same reaction path, the obtained compounds presenting identical IR spectra. Therefore the assigning structure of compound **4** (spot 5) were confirmed also by chemical way.

The procedure and the complete description of the ¹³C-NMR spectra are presented in the experimental part. The description of structure was done for both conformers together and the notation used is: (cis, trans), meaning that both signals are present and also that the number of signals is double.

Equipment and Methods

NMR spectra were recorded on a Varian Gemini 300 MHz apparatus against the line of the solvent (CDCl₃). Chemical shifts are expressed in ppm. Mass spectra were recorded on a Varian Finnigan Mat 212 mass spectrometer. TLC analysis was performed on Kieselgel Merck plates with fluorescence indicator. Separation by flash chromatography was performed on 220-400 mesh silica (Aldrich). Visualization of the chromatograms was accomplished with UV lamp and by developing with a 20% H_2SO_4 in ethanol solution, then heating the plate at 120°C. Anhydrous solvents have been used.

Experimental

2 g (8.4 mmol) N-acetyl-N-methyl-glucamine is solved in 5 ml DMF (kept on 4Å molecular sieves) and a catalytic amount of camphorsulphonic acid is added under stirring. The reaction mixture is cooled to 0°C and 3.6 ml (24 mmol) dimethoxy benzaldehyde are drop by drop added under argon atmosphere and stirring. The reaction mixture is stirring at reflux temperature (48 h) until TLC control (AcOEt:MeOH = 90:10) indicates the disappearance of the starting material. The reaction is stopped by adding 0.5 ml Et₃N and the solvents are vacuum distilled with toluene. The mixture is then separated and purified by flash chromatography (Table 1).

The 5,6-O-benzyliden-N-acethyl-N-methyl glucosamine was obtained in 52% yield. All products are as consistent clays at room temperature.

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Table 1.

Flash chromatography and TLC separation and analysis conditions of the acetalization products

Spots separation	FC elution system	TLC control system
1+2+3 spots by 4+5 spots	Tol:AcOEt=22:78	AcOEt :MeOH=95:10
1+2 spots by spot 3	Tol:AcOEt=22:78	AcOEt :MeOH=95:10
spot 1 by spot 2	Tol:AcOEt=22:78	AcOEt :MeOH=95:10
spot 3 by spot 2	Tol:AcOEt=25:75	AcOEt :MeOH=95:5
spot 4 by spot 5	Tol:AcOEt=25:75	AcOEt :MeOH=95:5
spot 5 by 4+5 spots	Tol:AcOEt=25:75	AcOEt :MeOH=95:5

Spot 1: 2,3:5,6- *di-O-benzylidene-(4-O-acetyl)-N-acetyl-N-methyl-glucosamine* + 3,4:5,6-*di-O-benzylidene-(2-O-acetyl)-N-acetyl-N-methyl-glucosamine* (compounds 2 and 3, Figure 2)

MS(EI), 22eV, (characterized as monoacetylated derivative $C_{25}H_{29}NO_7$) m/z= 455(M⁺.)

¹³CRMN (75MHz, CDCl₃)

171.25, 171.19, 170.82, 170.40, 169.96, 169.64, 169.52 (4C, 2X \underline{CH}_3 -CO, (cis, trans), (the isomers 2 and 3)) 137.34, 136.80 (4C, 2x \underline{C} from Ph, (cis, trans), the isomers 2 and 3)) 129.71, 129.66, 129.38, 129.28, 129.07, 128.34, 128.29, 128.20, 126.30, 126.21, 126.16, 126.09, (20C, 10x \underline{C} H from Ph, (cis, trans), the isomers 2 and 3)) 104.20, 104.01, 103.90, 103.74, (4C, 2x \underline{C} H-Ph, (cis, trans), the isomers 2 and 3)) 177.42, 76.99, 76.57, 74.98, 74.37, 74.24, 70.33, 70.28, 70.20, 70.06, 69.88, 68.09, 68.03, (8C, \underline{C}_2 , \underline{C}_3 , \underline{C}_4 , \underline{C}_5 , (cis, trans), the isomers 2 and 3)) 67.37, 67.25, 67.10, 66.59, (2C, \underline{C}_6 , (cis, trans), the isomers 2 and 3)) 50.77, 50.45, (2C, $\underline{C1}$, (cis, trans), the isomers 2 and 3)) 37.19, 37.07, 33.52, 33.41, (2C, $\underline{C}H_3$ -N, (cis, trans), the isomers 2 and 3)) 21.54, 21.50, 21.10, 20.75, 20.71, 20.67, 20.58, 20.53, 20.45, 20.38, (4C, $\underline{C}H_3$ -CO, (cis, trans) the isomers 2 and 3)) **Spot 2:** 2,3 :4 6-di-O-benzylidene-N-acetyl-N-methyl-glucosamine (compound **4**a, Figure 2)

MS(EI), 22eV, $(C_{23}H_{27}NO_6)$, m/z= 413(M⁺·) ¹³**CRMN (75MHz, CDCI**₃) 171.97, 171.46, (1C, N-C=O, (cis, trans)) 137.97, 137.90, (2C, C from Ph, (cis, trans)) 129.61, 129.57, 129.41, 129.35, 129.09, 128.99, 128.38, 128.28, 128.27, 128.17, 126.71, 126.61, 126.55, 126.43, 126.34, 125.92, (10C, CH from Ph, (cis, trans)) 104.52, 104.33, 103.70, (1C, CH-Ph, (cis, trans)), from dioxolane ring 101.02, 100.90, 100.76, (1C, CH-Ph, (cis, trans)), from dioxane ring 78.69-68.56 (multiple signals), (4C, C2, C3, C4, C5, (cis, trans)) 67.91, 67.55, (1C, C6, (cis, trans)) 49.02, 48.91, (1C, C1, (cis, trans)) 38.42, 38.16, (1C, CH₃-N, (cis, trans)) 21.75, 21.63, (1C, CH₃-CO, (cis, trans))

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Spot 4: *5*, *6*-*O*-*benzylidene-N*-*acetyl-N*-*methyl-glucosamine* (compound **1**, Figure 2) **MS(EI)**, (24eV) ($C_{16}H_{23}NO_6$), m/z=325, (M⁺.) ¹³**CRMN (75MHz, CDCI**₃) 173.06, (1C, N-<u>C</u>=O, (cis, trans)) 137.87, 137.29, (1C, <u>C</u> from Ph, (cis, trans)) 129.28, 129.13, 128.26, 128.19, 126.54, 126.44, 126.33, 126.31, 126.03, (5C, <u>C</u>H from Ph, (cis, trans)) 103.98, 103.57, (1C, <u>C</u>H-Ph, (cis, trans)) 74.02, 73.60, 72.47, 72.43, 70.09, 69.74, 69.67, 68.17, (4C, <u>C</u>2, <u>C</u>3, <u>C</u>4, <u>C</u>5, (cis, trans)) 68.71, (1C, <u>C</u>6, (cis, trans)) 53.77, 51.63, (1C, <u>C</u>1, (cis, trans)) 38.11, 37.95, (1C, <u>C</u>H₃-N, (cis, trans)) 21.63, 21.58, (1C, <u>C</u>H₃-CO, (cis, trans))

Spot 5: *4*, *6*-*O*-*benzylidene-N*-*acetyl-N*-*methyl-glucosamine* (compound **4**, Figure 2) **MS(EI)**, (24 eV) ($C_{16}H_{23}NO_6$), m/z=325, (M⁺.) ¹³**CRMN (75MHz, CDCI**₃) 172.18, (1C, N-<u>C</u>=O, (cis, trans)) 137.73, 137.33, (1C, <u>C</u> from Ph, (cis, trans)) 128.38, 128.27, 128.20, 126.75, 126.67, 126.56, 126.42, 126.23, 125.97, (5C, <u>C</u>H from Ph, (cis, trans)) 100.59, 100.47, (1C, <u>C</u>H-Ph, (cis, trans)) 79.53, 79.09, 78.40, 78.34, 78.04, 72.38, 72.24, 70.13, 69.57, 69.30, (4C, <u>C2</u>, <u>C3</u>, <u>C4</u>, <u>C5</u>, (cis, trans)) 63.44, 62.71, (1C, <u>C6</u>, (cis, trans)) 49.50, 49.11, (1C, <u>C1</u>, (cis, trans)) 38.47, 38.41, (1C, <u>CH₃-N</u>, (cis, trans)) 21.72, 21.59, (1C, <u>CH₃-CO</u>, (cis, trans))

Conclusions

The reaction between N-acetyl-N-methyl-glucamine and dimethoxy benzaldehyde in acid catalysis (camphorsulphonic acid) gives mainly the 5,6-monobenzylidene acetal (compound 1, Figure 2) with 52% yield. The new compounds: 5,6-O-benzylidene-N-acetyl-N-methyl-glucosamin (as main 2,3:4,6-di-O-benzylidene-N-acetyl-N-methyl-glucosamine, 4,6-Oproduct). benzylidene-N-acetyl-N-methyl-glucosamine and a mixture of 2,3:5.6- with 3,4:5,6-di-O-benzylidene-N-acetyl-N-methyl-glucosamine (not previously described in the literature), were characterized by MS-EI, ¹H-NMR, and ¹³C-NMR. The identity of products was confirmed by MS and the ring type was established by ¹³C-NMR, COSY and HETCOR experiments. The careful study of the spectra indicates the existence of cis-trans conformers. The existence of conformers makes difficult their characterization by ¹H-NMR, therefore they were characterized mainly by ¹³C-RMN. The optimal conditions for reaction monitorization and for flash chromatography separation and purification have been also established.

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THE INHIBITORY EFFECT OF THE ATENOLOL UPON THE ENZYME CATALYZED HYDROGEN PEROXIDE DECOMPOSITION

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ABSTRACT. The reaction of decomposition of hydrogen peroxide in the presence of catalase has been investigated by means of a Clark oxygen sensor, in the presence and absence of various concentrations of Atenolol – a β -blocker drug - having an inhibiting role. The Michaelis – Menten kinetic parameters have been determined from Lineweaver-Burk plots. The inhibition pattern we have deduced, suggested by the Lineweaver - Burk plots, corresponds to a fully mixed inhibition mechanism. Inhibition constants K_I and K_I were determined. A kinetic method for the determination of Atenolol has been suggested.

Keywords: Atenolol, Catalase, Hydrogen peroxide decomposition, Inhibition mechanism, Kinetic methods.

Introduction

Catalase, an oxidoreductase EC 1.11.1.6, is a tetrameric haeminenzyme consisting of a 4 identical tetrahedrally arranged subunits of 60000 g/mol each [1]. Catalase is widely distributed in nature It is found in all aerobic microorganism, in plant and animal cell [2]. The enzyme, when located in organelles, acts as a regulator of the H_2O_2 level, while in erythrocytes, catalase provides) a protection for hemoglobin against the oxidizing agents like H_2O_2 together with gluthathione peroxidase [3].

The enzyme - catalyzed reaction:

$$2H_2O_2 \xrightarrow{\text{catalase}} 2H_2O + O_2 \tag{1}$$

has been the subject of extensive investigations, both in regard to the kinetics and to the mechanism of the decomposition [4, 5, 6, 7, 8].

Similar reaction were also observed for other organic peroxides [9, 10] Well-known inhibitors of these reactions are ions like azide, cyanide and fluoride [11, 12, 13], or some organic compounds compounds like 3-amino-1,2,4-triazole [14] and atenolol[15].

Atenolol is a cardioselective β -adrenoceptor blocking agent (β -blocker).

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Atenolol (C₁₄H₂₂N₂O₃) according to European Pharmacopeia (±)-2-[4-(2-hydroxi-3-izopropylaminopropoxil)]-phenyl]-acetamide[16]



Atenolol has a chiral center, and is clinically used as a racemate [15]. B-adrenoceptor antagonists are a group of compounds that competitively inhibit the effects of catecholamines at β - adrenergic receptors [17]. These agents are used widely in clinical medicine for the treatment of varios deseases including hypertension, angina pectoris [18] cardiac arrythmias, hypothyroidism and glaucoma.

The main aim of this paper is the description of the inhibition mechanism. It would permit to develop a kinetic method of the determination of the drug Atenolol in very low concentration, based on the inhibitory effect on the rate of hydrogen peroxide decomposition.

Experimental

Equipment. The measurements were undertaken with a Clark oxygen sensor, attached to a Multiline P4 multimeter with automatic data acquisition on a PC. The sensors cover a measuring range from 0 up to 19.99 mg/L for the dissolved oxygen, with a resolution of 0.01 mg/L and an accuracy of $\pm 0.5\%$ from 5.30 °C. All experiments were performed in a vessel provided with a water jacket. The temperature was maintained constant at 20 \pm 0.1 °C, by means of a Falc 90 recirculatory water bath. The reaction mixtures were stirred with a magnetic stirrer, always with the same frequency.

Reagents and Solutions. We used bacterial catalase from *Micrococcus Lysodeikticus* 176340 U/ml (where the enzyme unit 1U is the amount of enzyme needed to transform 1 µmol of the substrate within 1 min, under standard conditions), from FLUKA with a purity index of 0.85, which came from the ratio of absorbances at 405 nm and 280 nm, $A_{405}/_{A280}$.

The molar concentration of catalase was determined spectrophotometrically at 407 nm, where the extinction coefficient is known to be $4x10^5$ mol⁻¹ dm³ cm⁻¹ [19].

All the other reagents were of analytical reagent grade. The solutions were prepared with de-ionized, four-times distilled water in order to avoid the interference of heavy metals. Stock solutions of catalase (9x10⁻¹⁰

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mol/L), H_2O_2 (4.9 x 10⁻² mol/L) and atenolol (10⁻⁵ mol/L) in phosphate buffer of pH=7.0 were freshly prepared before each set of experiments. Hydrogen peroxide has been standardized against potassium permanganate in acidic media.

Procedure

The reaction mixture of 10 mL volume was prepared directly in the reaction vessel connected to a thermostate, where the solutions of the reagents were kept. Measured volumes of buffer and atenolol stock solution were placed in the vessel, and the change in the concentration of oxygen was monitored while the mixture was continuously stirred. The oxygen concentration became constant after about 50 s, when a known volume of H_2O_2 stock solution was added to the mixture. After about 50 s the concentration of oxygen reached a constant value again, so that the catalyzed reaction was started by the injection of 2 ml of the stock solution of the enzyme. A typical oxygen concentration versus time curve obtained by monitoring the reaction with a Clark sensor is presented in Fig. 1.



A typical curve for O_2 evolution, recorded with the Clark sensor in the reaction mixture pH=7 at 20^oC

The rate has been measured after the introduction of the enzyme solution into the mixture as the slope of the oxygen concentration increase with time. Only the starting period, within 25 - 30 % of reaction, with a linear dependence has been considered. The degree of H₂O₂ transformation was calculated from the experimental measured oxygen concentration as

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$$X = \frac{2[O_2]_t - [O_2]_0}{[H_2O_2]_0}$$
(1)

where $[O_2]_t$ is the actual concentration of O_2 , and $[O_2]_0$ is the constant value of $[O_2]$ obtained after the addition and consumption of all amount of H_2O_2 .

Results and Discussion

A large amount of information is available in literature about the kinetics and the mechanism [5, 20- 22] of the reaction in the absence of inhibitor

The initial reaction rates, r_0 , were determined from the slopes of the early part of the O_2 evolution curves, after catalase addition. Although the catalytic cycle seems to be quite complicate [23] because of the various oxidation states of Fe in heme, our data of initial rates fitted well to Michaelis-Menten equation [24]

$$r_{0} = \frac{k[E_{0}][S]}{K_{M} + [S]} = \frac{r_{max}[S]}{K_{M} + [S]}$$
(2)

where $[E]_0$ and [S] stand for the initial concentration of the enzyme and for the substrate concentration respectively, K_M and r_{max} are the Michaelis-Menten parameters, and k is the rate constant of breakdown of the enzyme substrate complex to the product. This equation can be brought into the double-reciprocal form:

$$\frac{1}{r_0} = \frac{1}{r_{max}} + \frac{K_M}{r_{max}} \cdot \frac{1}{[S]}$$
(3)

and used to obtain the Lineweaver-Burk plots [25]. The value of the Michaelis-Menten constant $K_{\rm M}$ = 3.94x10⁻⁴ mol/L and the maximum velocity $r_{\rm max}$ =4.64x10⁻² mol L⁻¹ s⁻¹, obtained by us, are comparable to those mentioned in literature [26].

The Influence of Atenolol on the Enzyme-Catalyzed Decomposition of Hydrogen Peroxide

Several kinetic runs performed in the presence of different concentrations of atenolol proved the inhibitory effect of this compound. When the concentration of H_2O_2 was varied at several fixed concentrations of atenolol, Lineweaver-Burk plots for the inhibited reaction

$$\frac{1}{r_{o(I)}} = \frac{1}{r_{max(I)}} + \frac{K_{M(I)}}{r_{max(I)}} \cdot \frac{1}{[S]}$$
(4)

resulted in a family of straight lines (Fig. 2), which have a common intersection point on the left side of the ordinate and below the abscissa.



Fig.2.

Lineweaver Burk plots obtained for different concentration of atenolol

As the slopes of Lineweaver- Burk plots depend on the inhibitor concentration, a systematiuc study could lead to a kinetic method of determination of this inhibiting agent concentration.

The Michaelis-Menten parameters in the presence of the inhibitorl were determined from the parameters of the lines in figure 2. Their values are:

$$r_{max(-I)} = r_{max} \quad \frac{1 + \beta \frac{[I]}{K'_{I}}}{1 + \frac{[I]}{K'_{I}}} \qquad \text{and} \qquad K_{M(I)} = K_{M} \left(\frac{1 + \frac{[I]}{K_{I}}}{1 + \frac{[I]}{K'_{I}}}\right) \tag{5}$$

where: $K_{M(I)}$ is the apparent Michaelis constant in the presence of inhibiting agent, K'_{I} and K_{I} are the dissociation constants of the enzyme – substrate – inhibitor complex, ESI, and the enzyme inhibitor complex, EI, respectively.

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The constant β is equal to zero for full inhibition, while for partial inhibition $0 < \beta \le 1$. Lineweaver-Burk plots of Fig. 2 correspond to the inhibition pattern of full or partial mixed inhibitors [26].

In order to distinguish between full and partial inhibition, the slopes and intercepts from the primary Lineweaver- Burk plots were re-plotted against the corresponding inhibitor concentration. They are presented in figure 3 Both





The plots of intercepts and slopes versus the inhibitor concentration

these plots gave straight lines with good correlation coefficients. The straight lines equations are intercept: $y = (8,0 \pm 2.07) \cdot 10^7 x + (23,55 \pm 5,08)$, and slope: $y = (42 \pm 7.76) \cdot 10^2 x + (85 \pm 1,9) \cdot 10^{-4}$. This behavior is considered as typical for full inhibition [26]. Considering the possibility of a

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Table 1.

Mean values or the maximum velocity $r_{\text{max}(l)}$ and Michaelis Menten constant in the presence of the inhibitor $K_{M(l)}$

[atenolol] ₀ x 10 ⁷	$r_{max(I)} \times 10^2$	$K_{M(I)} \times 10^4$
(mol/l)	(mol/ls)	(mol/l)
1.00	2.98	2.68
2.00	2.53	2.35
3.00	1.94	3.19
4.00	1.87	1.66

partial inhibition, a non-linear fit of the intercept $1/r_{max(I)}=f([I] \text{ and the slope } K_{M(I)}/r_{max(I)}=f[I]$ gave negative values for β .

Therefore it is very likely that atenolol acts as a reversible full mixed inhibitor according to the reaction scheme:



Scheme 1.

Mixed inhibition mechanism for atenolol, where E stands for catalase, S for hydrogen peroxide and I for atenolol.

The plot (Fig.3) of slopes and intercepts respectively against the inhibitor concentration were used to determine the dissociation constants mentioned in the scheme above, $K_I = (2.023 \pm 0.2) \times 10^{-6}$ M of EI and $K_I = (2.94 \pm 0.2) \times 10^{-7}$ M Of ESI. The constant K_I was obtained from the slope of the $K_{M(I)}/r_{max(I)}$ versus [atenolo] graph:

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$$K_{I} = \frac{K_{M}}{r_{\max \cdot slope}}$$
(6)

while the K_1 constant was determined from the slope of the $1/r_{max(1)}$ versus [atenolol] graph:

$$K'_{I} = \frac{1}{r_{\text{max} \cdot \text{slope}}}$$
(7)

Conclusions

The catalase –catalyzed decomposition of hydrogen peroxide in phosphate buffer was studied in presence of atenolol. The atenolol acts like a fully mixed inhibitor, as observed from Lineweaver-Burk plots.

Michaelis Menten parameters were determined for the decomposition of H_2O_2 in the presens of catalase yielding the values of K_M =3.94x10⁻⁴ mol/L and r_{max} =4.64x10⁻²mol/L⁻¹s⁻¹

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NEW HALO COMPOUNDS OF SILICON AND TIN, POTENTIAL PRECURSORS OF >E=C=P- HETEROALLENIC SYSTEMS

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ABSTRACT. New halo compounds of Si and Sn potential precursors of doubly-bonded derivatives of type >E=C=P- (E = 14 main group elements) have been synthesized. Compounds **1-3** (bearing a 2,4,6-tri-*tert*-butylphenyl group bound to phosphorus and one 2,4,6-tri-methylphenyl radical bonded to Si) are stabilized by the large steric hindrance which prevents the approach of monomers to form dimers or oligomers.

Introduction

The chemistry of the –P=C< phosphaalkenes has been intensively studied since the synthesis of the first such compound [1] in 1965. Its stabilization was then achieved by resonance effects; however, it was found that bulky organic groups, such as 2,4,6-tri-tert-butylphenyl (Mes*), 2,4,6-*tri-iso*-propylphenyl (Tip), 2,4,6-*tri*-metylphenyl (Mes), *tert*-butyl (t-Bu), etc offer a supplementary stabilization through their sterically protective effect.

The general available synthetic route to make a bond of an E 14 group element to the C=P moiety is shown in Scheme 1:



Scheme 1

Due to the pronounced π -electron accepting effect, phosphaalkenes containing low coordinated phosphorus atoms have been investigated in various catalytic processes (polymerization of ethylene, hydroformylation reactions, direct conversion of allylic alcohols) [2-5] and proved also to be useful synthons in organoelement chemistry [6].

Several metal complexes of phosphaalkenes in different coordination modes have been already synthesized and have been recently reviewed [6, 7].

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Silylated and stannylated phosphaalkenes are described in the literature [8-12]. The presence of group 14 elements in the phosphaalkene molecule could induce interesting electronic effects that would allow the fine tuning of the metal centre properties of the complexes. Their use as building block of new polymeric materials could also lead to interesting outcomes.

We report herein the synthesis and characterization of three novel phosphaalkenes substituted with group 14 organometallic groups starting from the sterically hindered Mes*PCCl₂.

Results and Discussion

The treatment of the dihalogenated phosphaalkene Mes*PCCl₂ with one equivalent of n-BuLi at -85°C leads to the formation of Mes*PC(Cl)Li, which can be used as a nucleophilic agent in the synthesis of silylated and stannylated compounds.

Thus, compounds **1** and **2** were obtained following the route shown in Scheme 2.



Scheme 2

Although these compounds were not completely purified, they were characterized through multinuclear NMR spectroscopy, and relevant data are given in Table **1**. The phosphorus and carbon chemical shifts of **1** are in the expected region for a P=C system [13,15].

In case of compound **2** the presence of a diastereotopic carbon atom (Me₃Si<u>C</u>H) leads to two different signals in ¹H-NMR for the methyl groups.

Compound **2** (the E isomer) show a doublet at 43.6 ppm (${}^{3}J_{P-Sn}$ =363.74 Hz) in the ${}^{119}Sn-NMR$ spectra and a singlet in the ${}^{31}P$ spectra [13,14].

A byproduct in the synthesis of **2** was the Z geometrical isomer, which gives signals at 306.07 ppm in ³¹P NMR spectra and 12.06 ppm (d, ${}^{3}J_{P-Sn}$ =374.93 78

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Hz) in ^{119}Sn NMR respectively. The relative proportion of the E/Z stereoisomers as shown by $^{31}\text{P-NMR}$ is 4/1.

Compound **2** has also been characterized through ²⁹Si-NMR spectroscopy and gives a doublet at -5.65 ppm with a silicon-phosphorus coupling constant of 58 Hz.

Table 1.

Relevant NMR data for compounds1 and 2

Compound	¹ H NMR	³¹ P NMR	¹³ C NMR
1	1.34 ppm (d, ⁵ J _{HP} = 0.33 Hz,	323.8 ppm (² J _{PSi}	173 ppm (dd, C=P)
	18H, <i>o</i> -t)	=58 Hz)	
	1.44 ppm (s, 9H, p-tBu) 2.25		
	ppm (s, 12 H, <i>o</i> -Me)		
	2.41 ppm (s, 6H, <i>p</i> -Me)		
	7.41 ppm (s, 2H, H _{arom})		
2	0.24 ppm (s, 9H, Me ₃ Si) 0.27	305.490 ppm	4.40 ppm
	ppm (s, 9H, Me ₃ Si) 1.33 ppm	$(^{2}J_{PSn} = 365.66)$	(wide signal, <u>C</u> H₃Si)
	(s, 9H, <i>p</i> -tBu) 1.49 ppm (s,	Hz)	174.25 ppm
	18H, o-tBu) 7.40 ppm (s, 2H,		(d, ² J _{PC} =102,85 Hz)
	H _{arom})		14.42 ppm (s, <u>C</u> HSn)
			135.09 ppm
			(d, ² J _{CP} = 71.61 Hz, ipso
			C Mes*)

The reaction of $MeSiCl_3$ with two equivalents of $Mes^*PC(Cl)Li$ (Scheme 3) lead to the formation of the sterically hindered silylated phosphaalkene **3**:

Mes*PCCL₂
$$\xrightarrow{\text{n-BuLi, THF}}$$
 Mes*PC
Li
 $1/2 \text{ MeSiCl}_3$ $\xrightarrow{\text{THF}}$ THF
 -70°C
Mes*P=C
Cl
Mes*P=C
Cl
3
Scheme 3

NMR data for **3** are given in Table 2.

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Table 2.

NMR chemical shift and coupling constants for 3.

Compound	¹ H NMR	³¹ P NMR	¹³ C NMR
	0.88 ppm (s, 3H, Me)	313.57 ppm	162.90 ppm
3	1.33 ppm (s, 9H, p-tBu) 1.44	$(^{2}J_{PSi} = 44.63)$	$(dd, {}^{2}J_{CP} = 75.46 \text{ Hz}, {}^{3}J_{CP}$
	ppm (s, 18H, o-tBu)	Hz)	= 5,5 Hz)
	7.41 ppm (s, 2H, Harom)		134.37 ppm
			(d, ² J _{CP} = 64.14 ppm, ipso
			C Mes*)

The ²⁹Si NMR spectrum shows a triplet at -1.61 ppm, with a ${}^{2}J_{SiP}$ coupling constant of 47.7 ppm, which is close to that observed in the ${}^{31}P$ NMR spectrum.

In the analogues halosilylphosphaalkenes Mes*PC(CI)SiPhTipX (X = CI, F) the silicon atom appears more shielded (-6.8 ppm and -10 ppm respectively), but the ${}^{2}J_{PSi}$ coupling constant is comparable (54 Hz in both cases).

The reactivity of these new compounds is under current study. Their potential application in the synthesis of new, stable heteroallenes of >E=C=P-type is also currently investigated. An approach to the obtaining of such unsaturated derivatives would be the reaction of **1-3** with t-BuLi, followed by intramolecular elimination of LiCl (Scheme 4):



Scheme 4

Conclusions

Three new compounds containing -P=C(CI)-E(CI)< (E = Si, Sn) skeleton stabilized by bulky substituents at phosphorus and E have been synthesized. The use of these dihaloderivatives for the synthesis of heteroallenes containing low coordinated phosphorus atoms is under current investigation.

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Experimental

All manipulations were carried out using Schlenk techniques under inert atmosphere. Solvents were dried and freshly distilled from sodium/benzophenone.

NMR Spectra were recorded in $CDCI_3$ on a Bruker AC 200 instrument at 200.1, 50.1, 81.01 MHz (for ¹H, ¹³C and ³¹P respectively) and an AC 300 instrument at 100.6 MHz for ¹³C, ²⁹Si and ¹¹⁹Sn.

MeSiCl₃ and MesH were available commercially (Aldrich). 1.6 M solution of n-BuLi and t-BuLi (Aldrich) in hexane were used throughout. Mes*PCCl₂ has been prepared according to the literature procedure [16].

Synthesis of {Chloro-[chloro-bis-(2,4,6-trimethyl-phenyl)-silanyl]-methylene}-(2,4,6-tri-tert-butyl-phenyl)-phosphane (1):

To a solution of 1 g Mes*PCCl₂ (2.7 mmols) in 20 mL THF 1.9 mL of n-BuLi 1.6M were added at -90°C. The reaction mixture was stirred for an hour at -85°C and than added dropwise to a solution of 0.98 g Mes₂SiCl₂ (2.7 mmols) in 30 mL THF, cooled at -70°C. The solution was allowed to warm up at room temperature and the solvent was removed under vacuum and replaced with pentane. LiCl was removed by filtration. MS(m/z): 624 (M+), 609 (M⁺-Me), 491 (M⁺-Mes-Me), 379 (M⁺-Mes*), 119 (Mes)

Synthesis of {[Bis-(bis-trimethylsilanyl-methyl)-chloro-stranyl]-chloro-methylene}-(2,4,6-tri-tert-butyl-phenyl)-phosphane (2):

To a solution of 2.12 g Mes*PCCl₂ (5.9 mmols) in 30 mL THF 3.3 mL of n-BuLi 1.6M were added at -90°C. The reaction mixture was stirred for an hour at -85°C and than added dropwise to a solution of 3 g of Bis₂SnCl₂ (5.9 mmols) in 30 mL THF, cooled at -70°C. The solution was allowed to warm up at room temperature and the solvent was removed under vacuum and replaced with pentane. LiCl was removed by filtration. Several attempts to obtain a crystalline product failed but by eliminating the pentane, a white solid mixture was obtained from which compound **2** was identified through the NMR data. MS(m/z): 796 (M), 681 (M-2 t-Bu+1); 637 (M-Bis); 602 (M-Bis-Cl)

Synthesis of Bis[1-chloro-2-(1,3,5-*tri-tert*-butyl-phenyl)phosphaalkene] methylchlorosilane (3):

To a solution of 2 g Mes*PCCl₂ (5.4 mmols) in 30 mL THF 3.8 mL of n-BuLi 1.6M were added at -90°C. The reaction mixture was stirred for an hour at -85°C and than added dropwise to a solution of 0.40 g Me₃SiCl (2.7 mmols) in 15 ml THF, cooled at -70°C. The solution was allowed to warm up at room temperature and the solvent was removed under vacuum and replaced with pentane. LiCl was removed by filtration. Attempts to obtain crystalline product failed. By removing the pentane a light-yellow solid mixture was obtained from which compound **3** was identified through NMR data.

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BISMUTH(III) COMPLEXES WITH DIORGANODICHALCOGENO-PHOSPHINATO LIGANDS. SYNTHESIS AND SPECTROSCOPIC CHARACTERIZATION

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ABSTRACT. Metathesis reactions between BiCl₃ or OBiCl with alkali metal salts of diorganodichalcogenophosphinato ligands, in stoechiometric ratio, led to tris(diorganophosphinato)bismuth, $Bi(O_2PR_2)_3$ (R = Me, Ph, Cy, Prⁿ, Bz), tris(diorganomonothiophosphinato)bismuth, $Bi(O_2PPh_2)_3$, as well as the diphenyldithiophosphinato derivatives $Bi(S_2PPh_2)_3$ and $OBi(S_2PPh_2)$. The compounds were characterized by multinuclear NMR (¹H, ¹³C, ³¹P), IR and mass spectrometry. The spectroscopic data allowed us to predict a polymeric structure with anisobidentate ligands in case of the dioxo derivatives, while for the other species either monomeric or polymeric structures may be proposed.

Keywords: bismuth complexes, organophosphorus ligands, spectroscopic characterization

Introduction

Diorganophosphorus ligands of type $[R_2PXY]^-$ (R = alkyl or aryl, X, Y = O, S) and their metal complexes have attracted much interest for several years, due both to the diversity of their structural features and their potential applications (*i.e.* biological or catalytic activity). In compounds with such ligands different coordination patterns were observed, *i.e.* monodentate or bidentate (chelating or bridging two or more metal centers) as a consequence of (i) the ability of the ligand to involve one or both chalcogen atoms in interactions with metal centers, and (ii) the tendency of many Main Group or transition metals to increase their coordination number [1-4]. Several solid state structural types were established for different metal complexes by singlecrystal X-ray diffraction studies. Related to the applications of these species, a special interest is concerned to their solution behavior and structure. Even if a large number of metal complexes with $[R_2PXY]^-$ (X, Y = O, S) ligands were already described, only few bismuth complexes containing dithiophosphinato groups were so far studied and structurally characterized by single-crystal X-ray diffraction, *i.e.* Bi(S₂PMe₂)₃ [5], Bi(S₂PEt₂)₃ [6], Bi(S₂PPh₂)₃ [7], and

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MesBi(S₂PPh₂)₂ [8]. Our studies, reported here, were focused on some bismuth(III) inorganic species with dioxo-, monothio- or dithiophosphinato ligands. Spectroscopic data (multinuclear NMR, MS) are also included for $Bi(S_2PPh_2)_3$, only IR data [9] and the crystal structure [7] being previously reported.

Experimental

Solvents were pre-dried over activated 4 Å molecular sieves and then distilled from potassium (THF), magnesium (methanol), calcium chloride (acetonitrile), Na₂CO₃ (acetone) under a continuous stream of argon. BiCl₃ and OBiCl were commercially products. All other starting materials were obtained according to literature methods: R_2PO_2Na (R = Me [10], Bz [11], Cy [12], Ph, Prⁿ [13]), Ph₂PS₂NH₄, Ph₂PSONH₄ [13].

¹H, ¹³C and ³¹P NMR spectra were recorded on a VARIAN GEMINI instrument operating at 300, 75.4 and 121.4 MHz respectively, relative to TMS (¹H and ¹³C) and H₃PO₄ 85 % (³¹P). Mass spectra were obtained using either a VARIAN CH7 (A) MAT instrument (EI) or a JEOL SX-102A spectrometer (FAB). IR spectra were recorded either as KBr pellets or Nujol mulls on a SPECORD IR UR 20 instrument.

Preparation of tris(dimethylphosphinato)bismuth(III), Bi(O₂PMe₂)₃ (1)

A mixture of 0.5 g (1.6 mmol) BiCl₃ and 0.68 g (4.8 mmol) $Me_2PO_2Na \cdot 1.5 H_2O$ in acetonitrile (50 ml) was refluxed for 2 hours under argon atmosphere. The resulted NaCl was filtered off and the solvent was removed in vacuum, leaving the title compound as a white solid. Yield 0.54 g (70%). M.p. 71-73°C. IR:(KBr, cm⁻¹): 1295s, 1255m, 1150vs (asym PO_2); 960vs (sym PO_2); 870vs (P-CH_3); 750m, 700s (P-C). ¹H NMR (CDCl₃): 1.42 (d, 18H, CH_3 , ²J_{PH} 16.7 Hz). ¹³C NMR (CDCl₃): 16.50 (d, ¹J_{PC} 102.5 Hz). ³¹P NMR (CDCl₃): 51.9 (s). FAB+ MS [m/z (%)]: 1371 (5) [Bi₃(O₂PMe₂)₈⁺], 976 (8) [Bi₂(O₂PMe₂)₆⁺], 488 (17) [Bi(O₂PMe₂)₃⁺], 395 (58) [Bi(O₂PMe₂)₂⁺], 302 (10) [Bi(O₂PMe₂)⁺], 93 (100) [O₂PMe₂⁺], 77 (7) [OPMe₂⁺].

Preparation of tris(diphenylphosphinato)bismuth(III), Bi(O₂PPh₂)₃ (2)

A mixture of 0.98 g (3.1 mmol) BiCl₃ and 2.23 g (9.3 mmol) Ph₂PO₂Na·in methanol (100 ml) was stirred under argon atmosphere, for 2 hours, at room temperature. The white precipitate containing NaCl and the desired product was filtered off and washed with water to remove NaCl. The unsoluble white solid was dried to yield 2.07 g (77%) of the title compound. M.p. 237°C. IR (KBr, cm⁻¹): 1435vs (P-Ph); 1135s, 1085vs, 1065m (asym PO₂); 1030m, 1010vs, 990vs (sym PO₂); 745vs, 725vs, 690vs (PC). ¹H NMR (Me₂CO-d₆): 7.52 (m, 18H, C₆H₅-*meta+para*); 7.84 (m, 12H, C₆H₅-*ortho*). For ³¹P NMR, see subsequent discussion.

General method for the NMR tube scale reactions

 $BiCl_3$ and the sodium salt of the appropriate diorganophosphinic acid were mixed in a 1:3 molar ratio in CDCl₃, directly in the NMR tube, at room temperature, according to the data in Table 1, and the NMR spectra were recorded.

BISMUTH(III) COMPLEXES WITH DIORGANODICHALCOGENO-PHOSPHINATO LIGANDS

Tris(di-n-propylphosphinato)bismuth(III), Bi($O_2PPr_2^n$)₃ (**3**): ¹H NMR (CDCl₃): 1.02 (m, 18H, CH₃); 1.64 (m, 24H, CH₂-CH₂). ¹³C NMR (CDCl₃): 15.46 (s, C_{α}); 15.59 (d, C_{β} , ²J_{PC} 19.2 Hz); 31.22 (d, C_{v} , ¹J_{PC} 90.6 Hz). ³¹P NMR (CDCl₃): 58.4 (s).

Tris(*dicyclohexylphosphinato*)*bismuth*(*III*), *Bi*(O_2PCy_2)₃ (**4**): ¹H NMR (CDCl₃): 1.22-1.82 (m, 33H, C₆H₁₁). ¹³C NMR (CDCl₃): 24.61 (s, C₄); 25.92 (s, C₃); 26.19 (d, C₂, ²J_{PC} 13.5 Hz); 34.99 (d, C₁, ¹J_{PC} 90.6 Hz); ³¹P NMR (CDCl₃): 62.3 (s).

Tris(*dibenzylphosphinato*)*bismuth*(*III*), *Bi*(O_2PBz_2)₃ (**5**): ¹H NMR (CDCl₃): 2.84 (s, br, 12H, CH₂); 7.17 (s, br, 30H, C₆H₅). ¹³C NMR (CDCl₃): 36.21 (d, CH₂, ¹J_{PC} 88.6 Hz); 126.68 (s, C_p); 128.49 (s, C_m); 130.01s (C₀); 131.78 (d, C_i, ²J_{PC} 6.6 Hz). ³¹P NMR (CDCl₃): 48.5 (s).

Table 1.

Preparation details for $Bi(O_2PR_2)_3$ (NMR tube scale reactions).

BiCl ₃	NaO ₂ PR ₂	
g (mmol)	g (mmol)	
0.0305 (0.1)	R = Pr ⁿ	0.0499 (0.3)
0.0228 (0.07)	R = Cy	0.0547 (0.21)
0.0228 (0.07)	R = Bz	0.0551 (0.21)

Preparation of tris(diphenylmonothiophosphinato)bismuth(III), Bi(OSPPh₂)₃ (6)

A mixture of 0.31 g (1.0 mmol) BiCl₃ and 2.26 g (3.0 mmol) Ph₂P(S)ONH₄·in acetone (50 ml) was refluxed under argon atmosphere for 2 hours. NH₄Cl was filtered off and the solvent was removed in vacuum. The resulting colorless solid was recrystalized from benzene. Yield 0.5 g (55%). M.p. 180°C. IR (KBr, cm⁻¹): 1437vs (P-Ph); 1128s, 1109s; 1033m, 1013m, 994m (P-O) 656m (P=S); 770s, 748s, 732s, 693s (PC). ¹H-NMR (CDCl₃): 7.36 (m, 12H, C₆H₅-*meta*); 7.46 (m, 6H, C₆H₅-*para*); 7.83 (m, 12H, C₆H₅-*ortho*). ¹³C NMR (CDCl₃): 128.31 (m, C_m, ³J_{PC} 13.9 Hz); 131.45 (m, C₀, ²J_{PC} 12.8 Hz) 132.10 (s, C_p); 134.10 (d, C_i, ¹J_{PC} 113.1 Hz). ³¹P NMR (CDCl₃): 80.9 (s). El MS [*m*/*z* (%)]: 908 (5) [M⁺], 675 (10) [Bi(OSPPh₂)⁺], 442 (17) [Bi(OSPPh₂)⁺], 233 (5) [SOPPh₂⁺], 217 (59) [SPPh₂⁺], 201 (17) [OPPh₂⁺], 154 (100) [Ph₂⁺].

Preparation of tris(diphenyldithiophosphinato)bismuth(III), Bi(S₂PPh₂)₃ (7)

A mixture of 0.63 g (2 mmol) BiCl₃ and 1.6 g (6 mmol) Ph₂PS₂NH₄·in THF (50 ml) was stirred under argon atmosphere, for 3 hours, at room temperature. The solvent was removed in vacuum and the remained solid was treated with chloroform. NH₄Cl was filtered off and the solvent was removed in vacuum to give a yellow crystalline product. Yield 1.37 g (72%). M.p. 127°C. ¹H NMR (CDCl₃): 7.38 (m, 18H, C₆H₅-*meta*+*para*); 7.87 (ddd, 12H, C₆H₅-*ortho*, ³J_{PH} 14.5, ³J_{HH} 8.0, ⁴J_{HH} 1.2 Hz); ¹³C NMR (CDCl₃): 128.19 (d, C_m, ³J_{PC} 13.3 Hz); 130.55 (d, C_o, ²J_{PC} 12.1 Hz); 131.06 (s, C_p): 139.29 (d, C_i, ¹J_{PC} 83 Hz); ³¹P NMR (CDCl₃): 60.1 (s). EI MS [*m*/*z* (%)]:707 (96) [Bi(S₂PPh₂)²], 458 (22) [Bi(S₂PPh₂)⁴], 249 (55) [S₂PPh₂⁺], 217 (100) [SPPh₂⁺], 154 (5) [Ph₂⁺], 77 (5) [Ph⁺].

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Preparation of oxo(diphenyldithiophosphinato)bismuth(III), OBi(S₂PPh₂) (8)

A mixture of 1.3 g (5.0 mmol) OBiCl and 1.335 g (5.0 mmol) Ph₂PS₂NH₄ in acetone (50 ml) was refluxed for 2 hours. NH₄Cl was filtered off and the yellow solution was concentrated in vacuum. The resulting yellow solid was recrystalized from acetic acid. Yield 2.2 g (93%). M.p. 76°C. IR (KBr, cm⁻¹): 1436s (P-Ph); 610s, 660s (asym PS₂); 555vs (sym PS₂). ¹H NMR (CDCl₃): 7.32 (m, 4H, C₆H₅-*meta*); 7.40 (m, 2H, C₆H₅-*para*); 7.78 (dm, 4H, C₆H₅-*ortho*, ²J_{PH} 14.4 Hz). ¹³C NMR (CDCl₃): 128.46 (d, C_m, ³J_{PC} 13.9 Hz), 131.95 (d, C_o, ²J_{PC} 10.8 Hz), 132.23 (s, C_p), 132.88 (d, C_i, ¹J_{PC} 76.1 Hz). ³¹P NMR (CDCl₃): 71.0 (s).

Results and Discussion

Synthesis

Inorganic bismuth derivatives of type $Bi(X_2PR_2)_3$ (X = O, R = Me, Ph, Prⁿ, Cy, Bz; X = S, R = Ph), Bi(OSPPh_2)_3 and OBi(S_2PPh_2) were obtained by salt metathesis reactions between either BiCl₃ or OBiCl and an alkali metal or ammonium salt of the appropriate phosphorus-containing acid, according to equations (1) – (4):

BiCl₃ + 3 NaO₂PR₂
$$\longrightarrow$$
 Bi(O₂PR₂)₃ + 3 NaCl (1)
R = Me (1), R = Ph (2)
R = Prⁿ (3), R = Cy (4)
R = Bz (5)

$$BiCl_3 + 3 NH_4OSPPh_2 \longrightarrow Bi(OSPPh_2)_3 + 3 NH_4CI \quad (2)$$
6

$$BiCl_3 + 3 NH_4S_2PPh_2 \longrightarrow Bi(S_2PPh_2)_3 + 3 NH_4Cl \qquad (3)$$
7

$$OBiCI + NH_4S_2PPh_2 \longrightarrow OBi(S_2PPh_2) + NH_4CI$$
(4)
8

Compounds 1, 2 and 6–8 were isolated as solid products, while compounds 3–5 were generated in solution, directly in the NMR tube, and investigated spectroscopically without further isolation.

Some data concerning the new compounds and related bismuth(III) dithiophosphinates, already described in literature, are given in Table 2.

All compounds were investigated by NMR spectroscopy. Compounds 1, 2 and 6–8 were also characterized by infrared spectroscopy and mass spectrometry (EI or FAB).

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NMR Spectroscopy

180

127

76

242

180 (dec)

Bi(O₂PBz₂)₃ (5)

Bi(OSPPh₂)₃ (6)

Bi(S₂PPh₂)₃ (**7**)

OBi(S₂PPh₂) (8)

MesBi(S₂PPh₂)₂

Bi(S₂PMe₂)₃

Bi(S₂PEt₂)₃

The dioxo derivatives exhibit a poor solubility in CDCl₃. For compound **2** the NMR spectra could be obtained only in presence of more polar solvents, *i.e.* acetone- d_6 or DMSO- d_6 . This suggests a possible polymeric structure for these compounds.

Table 2.

Compound M.p. [°C] ¹P NMR (CDCl₃) Structure data Bi(O₂PMe₂)₃ (1) 71-73 51.9 NMR, MS, IR 12.5^b NMR, MS, IR^a Bi(O₂PPh₂)₃ (2) 237 **NM**R^á Bi(O₂PPrⁿ₂)₃ (**3**) 58.4 **NMR**^a $Bi(O_2PCy_2)_3(4)$ 62.3

Analitical data for the new Bi(III) compounds and related derivatives.

NMR^a

NMR, IR^a

NMR, IR^a

NMR, X-ray [5]

NMR, X-ray [6]

NMR, MS, X-ray [8]

NMR, MS,^a IR [9], X-ray [7]

^a This work; ^b In DMSO-*d*₆.

48.5

80.9

60.1

71.0

60.4 (C₆D₆)

In a mixture of $\text{CDCl}_3 - \text{DMSO-}d_6$, the ³¹P NMR spectrum of **2** exhibits two resonances, at δ 28.4 and 24.8 ppm, respectively. The resonance at lower field (δ 28.4 ppm) is very close to that observed for the free acid in CDCl_3 (δ 28.6 ppm). However, the ³¹P NMR spectrum in pure DMSO- d_6 for compound **2** contains only one resonance (δ 12.5 ppm), different from the values observed in the same solvent for Ph₂P(O)OH, (δ 24.7 ppm) or Ph₂PO₂Na (δ 17.4 ppm), which clearly suggests the formation of a Bi(III) species, probably a DMSO adduct, with a monomeric structure. The strong coordination ability of DMSO to the bismuth atom is well known [14,15].

In order to investigate the influence of the organic group attached to phosphorus upon the solubility of bismuth derivatives, some related species, *i.e.* $Bi(O_2PR_2)_3$ (R = Prⁿ, Cy, Bz), were generated directly in the NMR tube, in CDCl₃. For all dioxo derivatives the ¹H and ¹³C resonances are not significantly shifted compared to the free acids, but in all cases the ¹H resonance for the OH group, characteristic for the free acids, is absent, as well as the corresponding OH stretching vibration in the region 2600 - 1600 cm⁻¹, in the IR spectra (for compounds **1** and **2**). In contrast to the free acids, the proton resonances for the dioxo derivatives exhibit a broad aspect and the proton-proton and phosphorus-proton couplings are not resolved. The ³¹P resonances for the Me and Bz substituted derivatives are enough shifted

to conclude the formation of the desired complexes [*i.e.* δ 48.5 ppm for Bi(O₂PBz₂)₃ in contrast to δ 51.4 ppm for Bz₂P(O)OH, and δ 51.9 ppm for Bi(O₂PMe₂)₃ in contrast to δ 48.6 ppm for Me₂P(O)OH].

The Bi(III) compounds containing monothio or dithio ligands are soluble in common organic solvents and the NMR spectra in CDCl₃ solution are consistent with the formation of the desired compounds (**6–8**). The ¹H and ¹³C NMR spectra exhibit the expected resonances for the organic groups attached to phosphorus, split by the proton-proton and phosphorus-proton couplings, and phosphorus-carbon couplings, respectively. The ³¹P NMR spectra show only one sharp resonance.

Mass Spectra

The FAB+ MS spectrum of compound **1** contains the molecular ion in a relatively high abundance (12%), but peaks with higher mass are also present (*i.e.* m/z 1371 [Bi₃(O₂PMe₂)₈⁺], m/z 976 [Bi₂(O₂PMe₂)₆⁺]), thus consistent with the formation of molecular associations, at least of a trimeric species [Bi(O₂PMe₂)₃]₃.

For the monothio and dithio derivatives, peaks of low intensity, corresponding to the molecular ions, were observed in the EI spectra, along with different other peaks corresponding to ions formed by fragmentation. For compound **7**, even if the single-crystal X-ray studies revealed a dimeric structure [7], no fragments with m/z values higher than corresponding to a monomeric species were observed in the EI MS spectrum, which is consistent with weak intermolecular Bi···S interactions.

Infrared Spectra

The infrared spectra of the dioxo derivatives **1** and **2** are consistent with a coordination of the phosphinato ligands through both oxygen atoms to the metal center (*e.g.* vasym(PO₂) 1135s, 1085vs, 1065m, and $v_{sym}(PO_2)$ 1030m, 1010vs, 990vs for **2** *vs.* vasym(PO₂) 1035m, 1010s, and $v_{sym}(PO_2)$ 1142vs, 1133vs for Ph₂SbO₂PPh₂, for which a polymeric structure, with *O*,*O*'-bridging ligand, was established by single-crystal X-ray diffraction [16]). Taking into account the low solubility of this type of bismuth compounds and the observation in mass spectrum of compound **1** of fragments with a higher mass than expected for a monomeric species, polymeric structures with bridging ligand units might be considered. The difference in solubility might be due to a higher association degree for the phenyl substituted compound.

The strong IR absorptions observed for the compound **6** in the region 1130-1000 cm⁻¹ and at 656 cm⁻¹ are indicative for single P-O and double P=S bonds in the monothiophosphinato ligand [c.f. Ph₂P(S)OH [17]: v(P-O) 892vs, v(P=S) 633vs; Ph₂SbOSPPh₂ [16]: v(PO) 1061vs, 1050vs, v(PS) 593vs; for which a polymeric structure, with O,O'-bridging ligand, was

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established by single-crystal X-ray diffraction). This behaviour is consistent with the primary coordination of the monothio ligands to the metal centre through the oxygen atom, while the sulfur doubly bonded to phosphorus is probably involved in secondary intramolecular bonding, thus resulting in a distorted octahedral Bi $(O,S)_3$ core. Additional increase in coordination number of the metal centre through intermolecular bismuth chalcogen interactions, as observed in the related dimeric antimony derivative, $[Sb(OSPPh_2)_3]_2$ [18], cannot be excluded.

It was suggested that the difference Δ = vasym(PS₂) - vsym(PS₂) might be used for estimation of the coordination pattern of the dithiophosphorus ligands in metal complexes, *i.e.* Δ = 50-70 cm⁻¹, Δ = 70-90 cm⁻¹ and Δ > 95 cm⁻¹ indicates isobidentate, anisobidentate and monodentate coordination, respectively [18]. Indeed, for Bi(S₂PPh₂)₃ (7) the Δ = 75 cm⁻¹ [9] is consistent with the bidentate pattern of the dithio ligand found by single-crystal X-ray diffraction studies [7]. For the bismutyl derivative OBi(S₂PPh₂) (8) the difference Δ of 105 cm⁻¹ would suggest a monodentate coordination of the dithio ligand. However, taking into account that in all bismuth(III) complexes investigated so far by single-crystal X-ray diffraction [5-8] the dithiophosphorus ligands are coordinated through both sulfur atoms thus resulting in higher coordination number of the metal centre, molecular associations in solid state for compound 8 can not be excluded.

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FLAME ATOMIC EMISSION SPECTROMETRY DETERMINATION OF CESIUM IN MINERAL AND WELL WATERS USING A METHANE-AIR FLAME

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ABSTRACT. Effects of flame composition, the observation height and the spectral bandpass of the monochromator on the emission of Cs in methane-air flame were studied and optimized. The best results were obtained using the 852.19 nm Cs line at the observation height of 11 mm, with the flame composition of 1.12 (relative stoichiometric units) and the slit width of 0.8 mm. The effect of Li, Na, K, Rb, Mg, Ca, Sr, Zn, Al and PO₄³⁻ on the emission of Cs was studied too. The detection limit of $3 \pm 1.4 \mu g.L^{-1}$ was obtained, in the presence of 200 mg.L⁻¹ K, at a significance level of 0.05. The Cs content of mineral and well waters was determined without prior concentration by using both the external calibration and standard addition method. The results of these methods show good agreement implying the use of background correction.

Keywords: Cs, FAES, Water analysis

1. Introduction

Cesium is one of the widespread trace elements of the environment, its relative quantity in the Earth crust being of 7.10^{-5} %. It is distributed unequally in different environmental compartments; the mean Cs content of the rocks is of 0.7 mg.kg⁻¹, of the soils is of 3 mg.kg⁻¹. The seawater contains in average only 50 ng.L⁻¹ while the surface waters of 200 ng.L⁻¹, respectively. Higher concentrations of Cs occur in minerals (as pollutite, CsAlSi₂O₆), in certain volcanic rocks (in avogadite as CsBF₄) or can accompany certain minerals, as biotite and leopoldite, Cs content of which could reach 4.5 % of Cs.

Cs has no special physiological role. Plants (as barley, lichens), wild mushrooms and microorganisms (as *Escherichia coli, Anabaena.variabilis, Chlorella salina, Saccaromyces cerevisiae, Streptomyces lividans* etc.) accumulate primarily Cs *via* K⁺ transport in their organisms [1–6]. More harmful, with high biological risks, are the radioisotopes of Cs (¹³⁴Cs and ¹³⁷Cs), the main by-products of nuclear fission, being γ emitters with the half-life of 3 h and 33 years, respectively. Therefore, the main accent falls on the quantification of the Cs radioisotopes, most of the papers in this field workout

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radio-analytical method. Flame atomic absorption spectrometry (FAAS) is the standard method for the determination of the total Cs content of different samples, including natural waters, regardless of their origin. The determinations are carried out at 852.19 nm atomic line; the detection limits are in the order of magnitude of µg.L⁻¹ [7]. Due to the low Cs content of natural waters, the analyte is usually separated and preconcentrated prior measurements by passing through a Dowex 50-X8 resign column, followed by either an ammonium hexacyanocobalt ferrate or potassium hexanitrocobaltate column [8]. Flame atomic emission spectrometry (FAES) is also used for the quantification of Cs. The determinations are carried out either at the Cs 455.54 or the 852.19 nm atomic line; the detection limits being of order of 0.1 µg.L⁻¹. The common feature of these methods consists of the use of high temperature C₂H₂-air or C₂H₂-N₂O flames wherein Cs easily excites and ionizes, due to its low excitation and ionization energies, of 1.386 and 3.89 eV, respectively. The ionization is suppressed by adding KCI solution in 1 % final concentration in the sample [8-10]. From this point of view the cooler flames, as H2-air, propane-butane-air (PB-A), natural gas-air (NG-A) or methane-air (M-A) flames seems to be more advantageous for the direct quantification of Cs in low concentrations. So the lowest detection limits, of 30 fg.mL⁻¹, have been obtained in hydrogen and propane flames using the electron avalanche amplification of the laser enhanced two-step ionization of Cs [11].

The PB–A and NG–A flames are used in low performance commercial flame–photometers for the determination of alkaline metals designated for routine analyses. The M–A flame has similar properties (temperature, burning velocity etc) with the former ones [12, 13]. To our best knowledge the behaviour of Cs in the M-A flame has not yet been studied, nor the determination of Cs content of natural waters using this flame. The aim of this work is to study the behaviour of this element in the M–A flame, to optimize instrumental parameters and to determine the Cs content of natural waters, respectively.

2. Experimental

The optimal flame and instrumental parameters were established as follows: first, the emission spectrum of Cs was identified in the M-A flame; then the flame parameters (flame composition, observation height over the burner head (h)) have been optimized. In the following step the detection limit was determined under optimized conditions. Finally, the Cs content of some natural waters was determined.

2.1. Instrumentation

A HEATH–701 (Heath Co., Benton Harbour, MI, USA) spectrophotometer with a HEATH EU–700 scanning monochromator, a HEATH EU–700–30 type photomultiplier module and a M12FC51 (NARVA,

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Germany) photomultiplier (–1200 V) was used. The photomultiplier signal was introduced via a home made A/V converter/amplifier unit and data acquisition card (Decision–Computer International Co. Ltd., USA) into an IBM Pentium II PC (120 MHz, 32 Mb RAM) and processed using a home written Qbasic software program. The boxcar average technique was used, with data sampling interval of 0.33 ms. One measurement point was the average of 1500 consequent individual readings. The final data represented the mean value of the 50 measurement points. The other instrumental setup was the same as described earlier [14].

2.2. Chemicals

Stock solutions of Cs, Na, K, Li, Rb, Ca, Sr, Zn, Al, Mg and PO₄³⁻ were of 1000 mg.L⁻¹, prepared from RbCl, CsCl, Li₂CO₃, H₃PO₄ (Merck, Darmstadt, Germany), NaCl, KCl (Reactivul, Bucuresti, Romania), CaCO₃, SrCO₃, Mg, Zn and Al (Specpure, Johnson Matthey Chemicals Limited, England), respectively. The calibration solutions were obtained daily by diluting a given volume of stock solution with double distilled water.

2.3. Sampling and Sample Handling

The well water samples were collected in village Săvădisla (county Cluj), located 24 km's north-west from Cluj-Napoca city. The waters were sampled, handled and conserved in accordance with the EPA-recommendations [7]. The determinations were carried out within 72 hours after sampling. The mineral waters (commercial available, "Harghita", "Tuşnad", "Dorna", "Izvorul Minunilor") were analyzed after the sealed bottles were opened and degassed by shaking. All water samples were sprayed directly into the flame-photometer without any other treatment.

2.4. Procedure

The behaviour of Cs in the M-A flame was studied up to h = 18 mm over the burner head in 1 mm steps, at three different flame compositions: 0.88; 1.00; 1.12 (expressed in relative stoichiometric unit - RSU). Four replicate measurements were made; the mean and the standard deviation, the S/N and the S/B ratios were calculated for each h and flame composition. The homogeneity of the means was tested too by the F test at the significance level of 0.05. The sensitivity of the amplifier was set properly for each spectral domain, depending on the intensity of the emission lines and the background observed. For a given set of determinations, the sensitivity was kept constant.

3. Results and Discussion

3.1 The Emission Spectrum of Cs, Determination of the Analytical Emission Line

First the emission spectrum of Cs was determined by recording the spectrum of the flame alone, then that of the flame in the presence of Cs by spraying a solution of 100 mg.L⁻¹ Cs in the flame. The spectra were recorded in the 300–900 nm domain with a scanning rate of the monochromator of 0.02 nm.sec⁻¹. The composition of the flame was kept constant, of 1.12 RSU. Using spectral tables the emission lines with wavelengths of 455.54 nm, 459.32 nm, 852.19 nm and 894.35 nm were identified [10]. These are all atomic lines, the most intensive ones being of 852.19 nm and 455.54 nm, respectively.

The change of the analytical signal versus h at the formerly detected wavelengths was studied further. The concentrations of the calibration solutions were different, in function of the intensity of Cs emission line under study. The background intensity was measured in all cases in the presence of Cs, at the wavelengths of 457.0 nm, 461.0 nm, 854.0 nm and 896.0 nm, respectively. The variation of the net emission signal of different Cs lines versus h over the burner head is represented in Fig.1.



Figure 1. Variation of the net emission signal of the Cs lines vs. h. * not in scale (5 x magnification)

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For the quantitative comparison of the results obtained at different wavelengths only the maximum value of the net analytical signal (I) for each line was selected. The relative intensity (I_{rel}) for each line was calculated too. The reference concentration was of 100 mg.L⁻¹, the sensitivity the highest value used and the reference line was the weakest one. The S/B ratio was determined in the same way, the reference concentration being the highest one used. The results are summarized in Table 1.

Table 1. The relative intensities of the emission lines of Cs in the M-A flame

Wavelength (nm)	l (a.u.)	h (mm)	I _{rel}	S/N	S/B
455.54	39.4	10	75.76	296.5	6.9
459.32	9.6	10	18.46	72.2	2.0
852.19	62.8	11	120.76	468.7	157.0
894.35	0.52	11	1.00	4.2	2.5

The most intensive Cs line is that of 852.19 nm, superimposed on the low value of the background (high S/B ratio). For analytical purposes the 455.54 nm line could be used too, but in less favorable conditions than the former. The intensity of each Cs line varies in the same manner as the translational temperature of the flame, the maximum intensity being at 10–11 mm over the burner head [15] (see Fig.1.). In conclusion the analytical emission line for Cs in the M-A flame is that of 852.19 nm at h = 11 mm.

3.2. Determination of the Optimal Values of the Flame Composition and the Height of Observation Over the Burner Head

The influence of the flame composition, on h, on the analytical signal and S/N ratio for the most sensitive line was studied, at the concentration level of 10 mg.L⁻¹. The data were processed using the MicroCal OriginTM Software package, version 5 (MicroCal Software Inc., MA, USA) and plotted as 2D contour map (Fig.2).

The results show that the maximum intensity is obtained in all cases at the same h = 11 mm over the burner head, regardless of the flame composition. The analytical signal decreases slightly with the decrease of the methane content in the flame. The standard deviations of all means are homogeneous; therefore the magnitude of the S/N ratio is decided by the magnitude of the mean. In conclusion, the optimal conditions for the quantitative determination of Cs in the M–A flame are $\lambda = 852.19$ nm, h = 11 mm, flame composition of 1.12.

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Figure 2. The 2D contour map of the emission of the 852.19 nm Cs line vs. observation height and flame composition. The data labels on the plot indicate the grid matrix values.

3.3. The Influence of the Spectral Bandpass of the Monochromator on the Analytical Signal, S/N and S/B Ratio

The spectral bandpass of the monochromator, determined by its width of the slit (SW), influences in different manner the amplitude and the fluctuations of the emission signal. Therefore the optimal width of the slit for which the S/N ratio is maximum can be determined. The flame and instrumental parameters used were the optimal ones, determined earlier. The influence of the width of slit over I and S/N ratio was studied in the 0.1–1.5 mm domain in steps of 0.1 mm. The analytical signal exhibits a linear variation with the slit width up to 1.5 mm, the I–SW relationship could be approximated with a linear function (I = -43.986 + 0.7608 SW, R² = 0.9994). The standard deviation of the means is homogeneous up to a width of 0.8 mm and then they differ significantly. The highest value for the S/N ratio was found at a value of the width of 0.8 mm. In conclusion, the width of the slit could be increased up to 0.8 mm without decay of the S/N ratio.

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3.4. Interferences

The effect of Li, Na, K, Rb, Mg, Ca, Sr, Zn, Al and PO_4^{3-} ions on the emission signal of Cs of 1 mg.L⁻¹ was investigated, up to a concentration of the interferent of 500 mg.L⁻¹. Some of these elements are potential inorganic interferents, present in natural waters at higher concentrations. The optimal experimental conditions were described in a previous paragraph. The background signal was measured at 854.0 nm in the presence of the interferent. The variation of the emission signal of Cs versus the concentration of the interferents is represented in Fig. 3.



Concentration of the interferent (mg.L⁻¹)

Figure 3. Influence of Li, Na, K, Rb, Ca, Sr, Mg , Zn, Al and PO₄³⁻ on the emission of Cs of 1 mg.L⁻¹

The alkaline metals K and Rb, acting as ionization suppressors, enhance the emission signal of Cs up to an excess of 200 and 100 times, respectively. Both AI and PO_4^{3-} decrease significantly the emission signal of Cs, probably due to a low evaporation rate of Cs from the condensed phase in the M–A flame. The other interferents do not influence the Cs emission, even at higher concentrations.

3.5. Calibration, Determination of the Detection Limit

For the determination of the detection limit the variation of the analytical signal versus concentration was studied. Five calibration curves were plotted, in the $10-10^{-2}$ mg.L⁻¹ Cs concentration range. Each calibration curve was established by using six standard solutions and covers only one

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order of magnitude of concentration. Six replicate measurements were made at each concentration level. There were tested the homogeneity of means and the linearity of the calibration curve, the equation of the regression line, the confidence limits and the coefficient of correlation (R^2), with the least squares method were calculated. The detection limit was calculated using the two-step Neyman-Pearson criteria at the significance level of 0.025 [16,17]. The results are summarized in Table 2.

No. eq.	Concentration range Cs (mg.L ⁻¹)	Slit width (mm)	Equation of the calibration curve	Detection limit (mg.L ⁻¹)
1	1 – 10	0.1	I = - 1.0407 + 1.7398C R ² = 0.9991	0.4 ± 0.14
2*	1 – 10	0.1	l = 0.6355 + 3.0495C R ² = 0.9994	0.2 ± 0.09
3 [@]	1 – 10	0.1	l = 0.4992 + 2.8415C R ² = 0.9992	0.3 ± 0.24
4*	0.1 – 1	0.1	I = -0.0226 + 3.6297C $R^2 = 0.9983$	0.15 ± 0.06
5*	0.01 – 0.1	0.8	I = 0.0287 + 38.727C $R^2 = 0.9964$	0.003 ± 0.0014

Table 2. Calibration curves and detection limits (n = 6)

* with K added, in final concentration of 200 mg.L⁻¹

[@] with Rb added, in final concentration of 100 mg.L⁻¹

Cs exhibits linear dynamic range in the 1–10 mg.L⁻¹ concentration domain in the presence of K of 200 mg.L⁻¹, used as ionization suppressor. Rubidium has the same effect (Table 2, Eq.(3)), but in lesser extent. On the other hand the presence of Rb increases the noise level of the determinations, namely the instability of the flame, as compared with K. The increased slit width up to 0.8 mm allows the signal enhancement and the extension of the usable concentration about one order of magnitude. The detection limit obtained for each concentration range is also summarized in Table 2. The lowest detection limit, of 3 ± 1.4 µg.L⁻¹, obtained with the M–A flame is the same order of magnitude with those obtained with the hotter (C₂H₂–air and C₂H₂–N₂O) flames.

4. Determination of Cs in Water Samples

The Cs content of the water samples was determined by calibration and the standard addition method, the latter being used as reference method for the method validation, in the absence of certified reference material. The determinations were carried out with the SW of 0.8 mm. The other parameters were the optimal ones. Four parallel measurements (n = 4) were made in FLAME ATOMIC EMISSION SPECTROMETRY DETERMINATION OF CESIUM IN MINERAL...

all cases. Two sets of standard calibration solutions were prepared. One set contained the analyte in the presence of K of 200 mg.L⁻¹. In order to ensure the same evaporation conditions with the well water samples, the other set contained K, Na, Ca, Mg, as matrix, in the same concentration level as the well waters (K, Na of 100 mg.L⁻¹ and Ca, Mg of 50 mg.L⁻¹, respectively). The K, Na, Ca, Mg content of these samples has been determined earlier [18, 19]. In the case of standard addition method 100 µL of Cs stock standard was added to the 20 mL of sample. Three additions of standard were made, in 0.025 mg.L⁻¹ concentration steps. Using the data set, the reproducibility of the standard addition method was tested first. The recoveries found were within 88.2-132.3%, the slope of regression lines being close to that obtained by standard calibration. The results obtained with the standard calibration using K of 200 mg.L⁻¹ are systematically lower with 10-66 % than those obtained with the synthetic well water. This fact suggests the presence of the evaporation interference in the flame due to the high salinity of the well waters. For further considerations only the last case values were taken into account. The results of the two methods (Table 3) agree within the error of determinations for the given water. The confidence limits are wide due to the high noise level of the determinations close to the detection limit. The calibration method using synthetic well water could be used for quantification.

Sample		Concentration $(\mu g.L^{-1}, by calibration)$	Concentration (µg.L ⁻¹ , by standard addition)		
Minoral	"Harghita"	11 ± 1.6	12 ± 1.5		
water	"Tuşnad"	8 ± 1.7	7 ± 2.1		
water	"Dorna"	4 ± 1.6	4 ± 1.9		
	no.1	< 3 ± 1.4	< 3 ± 1.4		
	no.2	3 ± 1.7	3 ± 1.4		
Well water	no.3	5 ± 1.6	6 ± 0.9		
Well Water	no.4	5 ± 1.6	7 ± 1.1		
	no. 5	7 ± 1.7	7 ± 1.4		
	no. 6	10 ± 1.5	8 ± 1.9		

 Table 3. Results of analysis of water samples (n = 4)

The mineral water "Harghita" and "Tuşnad" have close Cs content, originating from the same volcanic region of the country (as indicated on the label of the bottles).

The well water samples nos.1–4 have practically the same Cs content, the wells being sunk very close one to the other (about 20 meters).

The other samples (nos.5–6), collected from another site of the village, have significantly higher Cs contents. The potassium and rubidium

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content of these samples are higher too as well as the K/Rb and K/Cs concentration-ratios [14, 19]. These facts show that the wells from different sites are supplied from other ground water sources, being sunk in rocks with different geological structure and chemical composition. The high alkaline metal content of the waters suggests the presence of rock salt bulbs or layers stratified among the sedimentary structures.

In conclusion, the Cs content of mineral and well waters can be determined directly in the M–A flame with acceptable precision by calibration method using synthetic well waters and background correction.

5. Conclusions

In the M–A flame Cs exhibits only an atomic spectrum. The most intense line is at 852.19 nm. The optimal excitation zone is at 11 mm over the burner head, in fuel rich conditions. The observation height for the maximum intensity is independent of the flame composition. In order to increase the S/N ratio, and to make the detection limit lower, the width of the slit of the monochromator could be increased till 0.8 mm without decay of the S/N ratio. The calibration curves for the 852.19 nm Cs line are linear in the 10–0.001 mg.L⁻¹ concentration range; the detection limit obtained is of $3 \pm 1.8 \ \mu g.L^{-1}$ Cs. The ionization of Cs most efficiently is suppressed with 200 mg.L⁻¹ K. The Cs content of mineral and well waters can be determined with acceptable precision using the calibration method and background correction.

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ANALYSIS OF FREE AMINO ACIDS FROM PLANT EXTRACTS BY CHROMATOGRAPHIC METHODS

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Dedicated to Professor Sorin Mager on the occasion of his 75th birthday

ABSTRACT. Three modern techniques: thin-layer chromatography (TLC), high performance liquid chromatography (HPLC), and gas chromatography-mass spectrometry (GC-MS) have been used for separation, identification and quantitative determination of free amino acids from *Equisetum arvense* and *Ocimum basilicum*. The results of this research explain the utilization of *Equisetum arvense* and *Ocimum basilicum* basilicum leaf extracts in phytopharmaceutical and cosmetic products.

Keywords: plant extract; chromatographic methods; separation; identification; quantitative determination

1. Introduction

Natural compounds offer without doubt the richest resources of chemical diversity. In the last decade the pharmaceutical and cosmetic industries have been extensively using medicinal plant. Isolation, identification and quantitative determination of the active compounds from plant extracts are some of the oldest fields for the application of chromatographic methods for studying the structure-activity relationship.

Chromatographic methods allow separation and identification of biological active compounds such as: amino acids, peptides, flavones, sugars, tannins, organic acids etc., their metabolites and intermediaries which do not display biological activity [1].

Thin-layer chromatography (TLC) is advantageous in separation and identification of amino acids from plant extracts because multiple samples can be analyzed simultaneously in a short time, with low detection limits. A variety of adsorbents such as silica gel [2], modified silica gel [3, 4], polyamide [5], alumina and cellulose [6] can be used for one- or two-dimensional separation of

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amino acids. The most frequently used mobile phase systems are n-butanolacetic acid-water, phenol-water, and n-butanol-acetic acid-acetone-water.

Separation of amino acids can be achieved by high performance liquid chromatography (HPLC) on C-18 stationary phase using mixtures of acetonitrile-acetate, phosphate, and citrate or borate buffer as mobile phases [7, 8].

The detection has been done under UV light if the adsorbent had a fluorescent indicator or after derivatization with ninhydrine, dansyl chloride, phtalic anhydride, dimethylamino azobenzene isothiocyanate (DABITC), etc [9, 10].

The best results for the quantitative determination of amino acids have been obtained using the gas chromatography-mass spectrometry (GC-MS) technique [11, 12].

The aim of this paper is the analysis of free amino acids from *Equisetum arvense* and *Ocimum basilicum* leaf extracts using TLC, HPLC, and GC-MS techniques.

Equisetum arvense is an excellent astringent genito-urinary system. It is useful mild diuretic and it is use in the treatment of kidney and bladder problems, cystitis, urethritis, prostate disease and internal bleedings such as urinary bleeds and stomach ulcer [13]. *Equisetum* has been found to ease the pain of rheumatism and stimulate the healing of chilblains. It is restorative to damage pulmonary tissue after pulmonary tuberculosis and other lung disease. In cosmetic, it is good for splitting nails and lifeless hair [14].

Ocimum basilicum is use as a culinary and medicinal herb. It acts principally on the digestive and nervous systems: easing flatulence, stomach cramps, colic and indigestion, poor digestion, nausea, gastro-enteritis, in treatment of feverish (especially colds and influenza), migraine, insomnia, depression and exhaustion [13]. Externally, it is use to treat acne, loss of smell, insect stings, snake bites and skin imperfections [14].

2.Experimental

2.1. Materials

Standard amino acids, $Na_3P(W_3O_{10})_4$ and trifluoroacetic acid were obtained from Merck (Darmstadt, Germany). Acetyl chloride and ion exchange resins Amberlite IR 120H and Dowex 50W-X8 were from Fluka (Buchs, Switzerland). ¹⁵N-glycine (Gly 98.9%) was obtained from ITIM (Cluj-Napoca). Thin-layer chromatographic separation of amino acids was achieved on 20x20 cm, 0.1 mm thick cellulose plates CEL 300-10UV254 (Macherey Nagel). All other chemicals obtained from Comchim Bucharest were analytical grade.

2.2. Extraction of Free Amino Acids

The isolation of amino acids can be done, from dry plants, using different extraction methods, such as: extraction with 5% NaCl solution, 75% ethanol,

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0.25% NaOH, 0.25M HCl, metasiliconic acid or a CH_3COOH -HCl-H₂O (18:1:1 v/v/v) mixture [12].

In our experiments the isolation of amino acids from dry plant has been done using two different extraction methods. For TLC and HPLC experiments 0.5g dry plants were extracted in 10mL 1% HCl solution. Then a 10% $Na_3P(W_3O_{10})_4$ solution was used for removing proteins from the extract by precipitation. After centrifugation the clear solution was passed through an ion exchange Amberlite IR120H column eluted with 40mL ammonia solution. The obtained solution was evaporated to dryness and the residue was redissolved in 1mL aqueous solution 30% (v/v) isopropanol. For GC-MS experiments 0.5g dry plant were extracted in ethanol. The amino acids were purified on a Dowex 50W-X8 exchange resins, in a 2x40 mm column and eluted with 3M ammonia solution.

2.3. TLC

Aliquot $(1\mu L)$ of standard solutions of 17 essential amino acids (1ng/mL) and the extract solutions were applied as spots to the cellulose plates with a micropipette.

The separation and identification of the free amino acids from the standard samples and extracts were achieved by bidimensional TLC (2D-TLC). The compositions of mobile phase were optimized by "Prisma" method. The plate was eluted on the first direction with n-butanol-acetone (35:35, v/v) and acetic acid-water (7:23, v/v) prepared and mixed (1:1, v/v) before elution. The second mobile phase was methanol-water-pyridine (80:20:5, v/v/v). The elutions have been done in unsaturated N-chamber and the elution distance was 18 cm. After the elution the plates were dried in hot air. The detection has been done by spraying the plates with a 0.5% ninhydrine solution in ethanol then dried at 110° C for 10-15 min. The identification of the amino acids was achieved by comparing the R_f values and the colors of the spots.

2.4. HPLC

The HPLC analysis amino acids from standard solution and from plant extracts were converted into their phenyl-thiocarbamilic derivatives.

The standard mixtures, the plant extracts $(10\mu L)$ were analyzed by HPLC using a HPLC apparatus (Merck Hitachi D-2000) with a Spherisol 5 ODS-2 column (250 mm x 4.6 mm i.d.) and a mixture of 0.15M sodium acetate (pH 6.5)-acetonitrile, concentration gradient 5-22%, 1 mL/min, temperature 55^oC, and UV detection at 254 nm. The identification of derivatized amino acids was achieved by comparing the retention time values.

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2.5. GC-MS

The amino acids were transformed into N-trifluoroacetil n-butyl ester to increase their volatility. Asparagine and glutamine were transformed into aspartic acid and glutamic acid, respectively. Hystidine and arginine were difficult to analyze by gas chromatography. [¹⁵N]-glycine was used as the internal standard (10 μ g/mL).

2.5.1. Derivatization

The amino acids were derivatized using two step derivatization procedure. The dry samples were esterified with 0.5mL distilled butanol-acetyl chloride (4:1, v/v) for 1h at 110° C. The excess reagent was removed by bubbling nitrogen through the mixture. The amino group was acetylated with a 200μ L mixture of trifluoroacetic anhydride (TFAA)-methylene chloride (1:1, v/v) at 60° C for 30 min. After cooling, the excess reagent was removed by bubbling nitrogen then 1mL ethyl acetate was added.

2.5.2. GC-MS

A trace DSQ ThermoFinnigan quadrupol mass spectrometer coupled with a Trace GC was used. The amino acids derivatives were separated on Rtx-5MS capillary column, 30m x 0.25mm, 0.25 μ m film-thickness, using a temperature program from 50°C to 300°C (3 min) at 10°C/min. The following conditions were followed: transfer line temperature 250°C, injector temperature 200°C, and ion source temperature 250°C, splitter 10:1. Electron energy was 70eV and emission current 100 μ A. The flow rate of helium was 2mL/min. 40 μ g of each samples was injected into the column.

The identification of amino acids was carried out by comparing the MS spectra with those from Wiley spectra library.

The quantitative determination of glycine was performed by selected ion monitoring (SIM). The following peaks were monitored for quantitative analyses: 154, 155 m/z from glycine.

3.Results and Discussion

3.1. Separation and Identification of Free Amino Acids by TLC

The R_f values of standards amino acids eluted with two mobile phase systems are presented in Table 1. Some of these amino acids were separated using the first mobile phase. The second mobile phase system could separate the amino acids that are unseparated by the first one. Therefore, the bidimensional elution is recommended.

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Table 1.

The R_f values of standard amino acids.

Amino acid	R _{f1}	R _{f2}
	BuOH-Acetone-HAc-H ₂ O	MeOH-H₂O-Py
	(35 : 35 : 7 : 23, v/v)	(80 : 20 : 5, v/v)
Glutamine (Glutamic acid)	0.77	0.68
Alanine	0.44	0.72
Proline	0.42	0.53
Tyrosine	0.33	0.14
Lysine	0.56	0.73
Phenylalanine	0.83	0.82
i-Leucine	0.85	0.92
Histidine	0.72	0.60
Serine	0.53	0.57
Valine	0.51	0.68
Asparagine (aspartic acid)	0.24	0.36
Methionine	0.81	0.78
Glycine	0.38	0.30
Arginine	0.50	0.43
Leucine	0.88	0.89
Threonine	0.51	0.68

The chromatographic results of bidimensional TLC analysis of studied plant extracts (Table 2) show the presence of glutamine, alanine, proline, glycine, lysine, phenylalanine, iso-leucine, asparagine, serine, threonine, methionine, valine and leucine in both *Equisetum arvense* and *Ocimum basilicum* extracts.

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Table 2.

The R_f values of amino acids from plant extracts

Amino acid	Equisetum	Equisetum arvense		asilicum
	R _{f1}	R _{f2}	R _{f1}	R _{f2}
Glutamine	0.75	0.67	0.73	0.66
Alanine	0.43	0.73	0.42	0.74
Proline	0.41	0.55	0.42	0.54
Glycine	0.31	0.13	0.32	0.13
Lysine	0.55	0.74	0.54	0.73
Phenylalanine	0.82	0.84	0.83	0.83
i-Leucine	0.84	0.94	0.84	0.93
Asparagine	0.31	0.29	0.30	0.30
Serine	0.54	0.58	0.55	0.59
Threonine	0.52	0.70	0.51	0.70
Methionine	0.79	0.79	0.80	0.79
Valine	0.50	0.43	0.50	0.42
Leucine	0.86	0.91	0.87	0.90

The identification was made on the basis of R_f values. The intensity of spot colors is only informative because it is different for different amino acids due to different quantity of compounds or probably to differences between the sensitivity of the reaction with ninhydrine.

3.2. Separation and Identification of Free Amino Acids by HPLC

The separation of standard amino acids is presented in Figure 1 and HPLC separations of plant extracts are presented in Figure 2.





Figure 1. The HPLC chromatogram of standard amino acids.

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Figure 2.

The HPLC chromatograms of: a - Ocimum basilicum; b - Equisetum arvense

These figures show that the amino acids present are those identified by TLC: glutamine, alanine, proline, lysine, phenylalanine, iso-leucine, serine, threonine and valine in the *Equisetum arvense* extract and alanine, proline, phenylalanine, iso-leucine, serine, methionine, valine, and leucine in the *Ocimum basilicum* extract.

3.3. Separation, Identification and Quantitative Determination of Free Amino Acids by GC-MS

The results of quantitative determination are presented in Table 3. By GC-MS the presence of the same amino acids are confirmed. Glutamine, alanine, proline, glycine, lysine, phenylalanine, iso-leucine, asparagine, serine, threonine, methionine, valine, leucine, histidine, and tyrosine are the free amino acids from *Equisetum arvense* and *Ocimum basilicum* extracts determined by this method.

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Table 3.

Amino acid	t _R	F	Equisetun	Equisetum arvense		Ocimum basilicum	
			A (mm²)	C (µg/g)	A (mm²)	C (μg/g)	
Alanine	12.75	0.77	1.04	8.65	0.96	8.27	
Glycine	13.40	0.80	0.90	7.56	0.87	7.32	
Threonine	14.81	0.20	0.05	1.65	0.11	3.51	
Serine	15.28	0.20	0.09	3.01	0.38	12.82	
Valine	15.76	0.84	0.89	7.09	1.20	9.56	
Leucine	17.82	0.80	1.04	8.69	0.91	7.58	
i-Leucine	18.11	0.80	0.49	4.12	0.37	3.06	
Tyrosine	19.90	0.95	0.55	3.85	0.27	1.90	
Proline	21.54	0.84	1.15	9.09	1.24	9.81	
Metionine	24.91	0.39	0.08	1.36	0.45	7.69	
Asparagine	27.73	0.88	1.07	8.07	2.35	17.84	
Phenilalanine	27.90	0.82	1.30	10.53	2.38	19.34	
Glutamine	30.90	1.00	2.07	13.77	2.81	18.75	
Lysine	30.96	0.43	0.68	10.47	0.69	10.75	
Histidine	35.93	0.51	0.07	0.98	0.23	3.02	

The GC results of quantitative determination of amino acid from plants extracts

4. Conclusions

The chromatographic methods allow the separation, the identification, and the quantitative determination of free amino acids from *Equisetum* arvense and *Ocimum basilicum* extracts. The important amino acids from *Equisetum arvense* and *Ocimum basilicum* are phenylalanine, asparagine, and lysine, which have an essential role in protein syntheses and in tissue regeneration, even if their concentration are small.

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STUDIES ON THE NATRIUM SULPHATE SALTING-OUT CRYSTALLIZATION II. CRYSTALLIZATION KINETICS

ADINA GHIRISAN, SIMION DRÅGAN, ALEXANDRU POP AND VASILE MICLÄUS^{*}

ABSTRACT. The kinetic of nucleation and growing of the crystal nuclei in the saltingout crystallization of natrium sulphate were studied. The influence of kinetic parameters (energy consumption or *agitator power input*, temperature and growth rate) on the second nucleation was determined. The influence of the hydrodynamic conditions and of the temperature on the average size of the obtained crystals was discussed. A value of activation energy of $5.4 \cdot 10^4$ J/mol for the nucleation process was determined.

Introduction

A crystallization operation consists of three basic steps: supersaturation (or supercooling), formation of crystal nuclei, and the subsequent growth of these nuclei into crystals [1]. Supersaturation may be achieved by cooling, evaporation of the solvent, addition of a precipitating agent, as a result of a chemical reaction, etc.

The conventional process of natrium sulphate crystallization through evaporation has many disadvantages: the high-energy requirement, the complexity of the installation with many steps and the complex process of separation of the obtained Glauber salt [2].

By the salting-out crystallization, by adding organic alcohol (methanol or ethanol) the solubility of the solute in the solvent supersaturated solution was reduced [3]. The equilibrium study of the salting-out crystallization has shown several advantages [3]:

a). The process can be operated at low temperature without output or input of energy in the crystallizer;

b). The construction of the installation is simple with lower cost of equipment and

c). Very pure crystals can be obtained because of high retention for the impurities of the mixed-solvent mother liquor.

To have a complete information about the process evolution in the case of salting-out crystallization of natrium sulphate the equilibrium data obtained in the first part of our study [3] are not enough. It is necessary to know more about the factors that can influence the nucleation and the crystal

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growth kinetics: intensity of mixing between organic and inorganic phases, concentration of the sulphate in aqueous solution, crystal size evolution, etc.

Nucleation

The generation of ultramicroscopic particles in the process of nucleation is the sum of contributions by primary and secondary nucleation. Primary nucleation occurs in the absence of crystals, while secondary nucleation is attributed to the influence of existing crystals.

The rate of primary nucleation has been modelled by the power law expression [4]:

$$B = \frac{dN}{dt} = k_n (C - C^*)^n$$
(1)

where *B* is the number of nuclei formed per unit volume per unit time, N – the number of new nuclei per unit volume, k_n – the rate constant, C – the instantaneous solute concentration, and C^* the solute concentration at saturation. The (*C*-*C**) term is the driving force of the crystallization process, called the *supersaturation*. The exponent n is typically in the range of 3 to 4 [5].

When a supersaturated solution is in contact with particles of the crystallizing compound, secondary nucleation occurs. Experimental data from industrial crystallizers have indicated that secondary nucleation usually predominates. There are two types of secondary nucleation: shear nucleation and contact nucleation [3]. *Shear nucleation* occurs as a result of fluid shear on growing crystal faces, and *contact nucleation* happens because of crystal collision with each other and with the impeller and other vessel internal surfaces. The most widely used relation for the rate of secondary nucleation in crystallization is the following [6]:

$$B = \frac{dN}{dt} = k_{B} \cdot \rho_{L}^{l} \cdot (C - C^{*})^{b}$$
⁽²⁾

where k_B is the rate constant and ρ_L is the suspension density. The exponent "*l*" ranges up 1.5, with 1 being the more probable value, the exponent *b* can be range up to 5 but has a most probable value of 2 [5].

Crystal growth

Crystal growth is the post nucleation process in which molecules are deposited on the surface of existing crystals. If crystal size is characterized by a characteristic dimension, L, it is convenient to express its linear growth rate by the equation [4]:

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$$G = \frac{dL}{dt} = k_g (C - C^*)^g$$
(3)

where *L* is a characteristic dimension of the crystal and k_g is an overall mass transfer coefficient; k_g is in general depended on the crystal size, hydrodynamic conditions, temperature, the nature of the ions and the presence of impurities.

Crystal growth is actually a process that consists of two steps in series. In the fist step the solute molecules or ions must reach the crystal surface by means of diffusion. In the second one, at the surface, the solute must be integrated into the crystal lattice. These two processes may be described as:

Diffusion:	$\mathbf{G} = \mathbf{k}_{\mathrm{d}} (\mathbf{C} - \mathbf{C}_{\mathrm{i}})^{\mathrm{d}}$	(4)

Surface integration: $G = k_r (C_i - C^*)^r$ (5)

where C_l is the concentration at the interface between the liquid and the solid phase, k_d and k_r are the mass transfer coefficients. If surface integration is very fast compared with bulk diffusion, $k_r >> k_d$ and $k_g \approx k_d$.

In conclusion, the formation of a stable crystal nucleus in a homogeneous or a heterogeneous system is not an easy process. Not only must the requisite number of molecules agglomerate, but also they must become oriented into a fixed lattice.

In this research study we have tried to establish the crystal formation rate during salting-out crystallization of natrium sulphate considering the influence of different factors as density of suspension, power input as the specific consumption of energy for mass unit, and crystal growth rate and the efficiency of the separation process.

Experimental

The experimental research was carried out in a batch isothermal stirred vessel of 0.5 I, type MSMPR (Mixed Suspension, Mixed Product Removal), with a device for the measuring and controlling of the temperature. The used agitator is an impeller with diameter blade of 4 cm.

The growth rates were obtained through granulometric analysis. For the granulometric analysis the suspension samples collected in time were filtered, washed with alcohol, dried and then analysed by sieving. The nucleation rates were obtained from the induction time, defined as the time between realisation of the known supersaturation and the appearance of the visible crystal [7]. The numbers of the crystals were determined for a diluted sample using a Coulter-Counter device [7]. ADINA GHIRISAN, SIMION DRĂGAN, ALEXANDRU POP AND VASILE MICLĂUȘ

The intensity of the hydrodynamic conditions was evaluated by the energy consumption or power input.

The following parameters were investigated:

- the residence time in crystallizer 5 – 180 min,

- the consumption energy (agitator power input) 0.18 1.64 W/kg, the temperature 30 70 $^{\circ}\text{C}.$

Results and Discussion The Efficiency of Crystallization

In this paper the efficiency of crystallization is defined as the ratio between the experimental obtained mass and the theoretical mass of the crystals. The data obtained from Fig. 1 have pointed out that the efficiency of the crystallization depends on the temperature and residence time in the crystallizer at constant energy consumption. The diagram from Fig. 1 shows that at a specific energy consumption of 0.18 W/kg a higher than 90 % efficiency can be obtained only by a residence time of 120 minutes and a temperature of 70 °C. To explain this dependence we have performed some kinetic studies.



Fig. 1. Efficiency of crystallization versus temperature at different residence timefor a power input of 0.18 W/kg.

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Crystallization Kinetics

Estimates of nucleation and growth rates during crystallization can be obtained from the population balance analysis by batch experiments [4]. By making a mass balance on the number of crystals within a given size range between *L* and *L* + ΔL , the population balance equation for size independent growth in a perfectly mixed batch crystallizer with negligible breakage and agglomeration is:

$$\frac{\partial(\mathbf{n}\mathbf{V})}{\partial t} + \frac{\partial(\mathbf{n}\mathbf{V}\mathbf{G})}{\partial \mathbf{L}} = 0$$
(6)

where *n* is the population density defined as the number of crystals of specific size per unit volume of crystallizer (number/m³m¹), *L*- size of crystals at time t (m), *G* – overall linear growth rate of crystals (m/s).

The mass balance in the MSMPR crystallizer with ideal mixing can be obtained using the equation data (solubility curve) and definition of the moments of distribution as pointed by Randolph [8]. For an MSMPR crystallizer with ideal mixing, with constant volume, without agglomeration and breakage, which means that the growth rate independent of the size of the crystal is, operating at steady state, a solution of population balance (equation 6) is:

$$\mathbf{n} = \mathbf{n}_0 \exp\left(-\frac{\mathbf{L}}{\mathbf{G} \cdot \boldsymbol{\tau}}\right) \tag{7}$$

When $L \rightarrow 0$ the solution of equation (6) will be:

$$\mathbf{B}_0 = \mathbf{n}_0 \cdot \mathbf{G}_0 \tag{8}$$

The equation (8) suggests that the secondary nucleation rate B_0 (number/m³s¹) is the product of population density n_0 (number of crystals/m⁴) and the crystal growth rate G_0 (m/s), is defined as equation (3) has already shown.

There are many empirical expressions that correlate the second nucleation rate with the crystallization parameters. The mostly used for the contact nucleation is given by the next equation [9]:

$$\mathbf{B}_{0} = \mathbf{k}_{B} \cdot \boldsymbol{\rho}_{L}^{1} \cdot \boldsymbol{\varepsilon}^{\mathrm{r}} \cdot \mathbf{G}_{0}^{\mathrm{p}}$$
(9)

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The analysis of equation (9) shows the dependence of the nucleation process on the suspension density (ρ_L) to a power exponent "*l*", on power input (ϵ) to a power exponent "*r*" and on growth rate (G₀) to a power exponent "*p*".

By operating on MSMPR crystallizer at steady state for different systems Mersmann [10] it has been concluded that the value of the exponent "f" = 1.

Because the effective nucleation rate B_0 is influenced by the crystal mass in crystallizer the density of the suspension will be replaced by the retention degree Φ defined by the equation:

$$\Phi = \frac{\rho}{\rho_s} \tag{10}$$

where ρ is the suspension density and ρ_{S} – the solid density.

So, the new equation of modified nucleation rate is:

$$\frac{\mathbf{B}_{0}}{\Phi} = \mathbf{k}_{B} \cdot \boldsymbol{\rho}_{S} \cdot \boldsymbol{\varepsilon}^{r} \cdot \mathbf{G}_{0}^{p}$$
(11)

For the determination of the power exponents we have assumed that the equation (11) can be simplified to the form:

$$\frac{B_0}{\Phi} = K_1 \cdot G_0^{p}$$
(12)

A straight line on the double log scale can represent this equation for all used consumption energy (Figure 2). The slope of the straight line corresponds to the value of the power exponent "p". The average value is 1.045, which corresponds to the literature values (1 - 3) [11].



Fig 2. Determination of the power exponent "p".

In order to obtain the value of the power exponent "r" the equation (11) was simplified to the form:

$$\frac{B_0}{\Phi} = K_2 \cdot \varepsilon^r \tag{13}$$

From the slope of the curves log B_0/Φ versus log ε an average value of r = 0.61 was obtained (Figure 3), which corresponds to the literature values (0.6 - 0.7) [11]. The results have shown the maintenance of a constant slope at different power input of stirrer.

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Fig. 3. Determination of the power exponent "r".

The influence of the temperature on the nucleation rate is represented by the constant rate for secondary nucleation k_B :

$$k_{B} = \frac{B_{0}}{\rho_{s} \cdot \Phi \cdot \varepsilon^{0.61} \cdot G_{0}^{1.045}}$$
(14)

as the equation of Arrhenius shows:

$$k_{\rm B} = k_0 \cdot e^{\frac{E_{\rm a}}{\rm RT}}$$
(15)

The plot log k_B versus (1/T) leads to a straight line with a constant slope for different linear growth rates (Figure 4). From the slope of the straight line the activation energy $E_a = 5,4 \, 10^4 \, J/mol$ was obtained.



Fig. 4.

The influence of the temperature on the nucleation rate.

Crystal size

The granulometric analysis of the obtained natrium sulphate has shown that the average diameter of the crystals lies in the range 150 - 500 μ m depending on the crystallization conditions. Sometimes the size of the crystal increases up to 1 mm, because of the agglomeration process that can be observed from the polycrystalline form of the particles visible only by microscopic analysis.

The analysis of the dependence of the average diameter of the crystals on the growth rate at a constant residence time can be calculated by the equation (16):

$$\mathbf{x}_{50} = \mathbf{K} \cdot \mathbf{G}_0 \tag{16}$$

where G_0 is defined by the equation (3). The equation (3) has shown that the linear growth rate depends on hydrodynamic conditions and temperature by means of the k_q coefficient.

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In Figure 5 the experimental values of the average size of the crystals x_{50} (mm) are plotted versus power input per unit mass ϵ (W/kg) at different temperatures. The linear dependence of the average diameter of the crystal on the power input at constant temperature was observed. At the same time the dependence of the crystal size on the temperature has shown that with the increase of the temperature, the average size of the crystals decreases for the same consumption of energy. The decrease of the temperature can be explained through the decrease of the crystallization potential (C - C*), supersaturation having a higher influence on the growth rate than constant K.



Fig. 5.

Crystal size versus specific consumption of energy.

Conclusions

The kinetic results have shown that the efficiency of the crystallization can be improved with the increase of the residence time and with the variation of the temperature in crystallizer.

By using the equation of population balance it was made possible to determine the specific influence of each parameter on the crystallization process.

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The experimental measurements have shown that the second nucleation constant k_B decreases with the increase of the temperature. From the Arrhenius equation the activation energy of nucleation $E_a = 5.4$ 10^4 J/mol was determined.

The dependence of the average diameter of the crystals on the power input ϵ was also determined.

The obtained results have shown that the salting-out crystallization leads to superior sizes of the crystal comparing the classical evaporation process and to a rigorous control of the process.

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STUDY ON THE INHIBITING BEHAVIOR OF A NON-TOXIC THIADIAZOLE DERIVATIVE ON BRONZE CORROSION IN AQUEOUS ELECTROLYTES

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ABSTRACT. The efficiency of a new non-toxic thiadiazole derivative (2 mercapto-5-Racetylamino-1, 3, 4-thiadiazole) as bronze corrosion inhibitor in various aqueous electrolytes (sodium chloride, sodium sulphate and sodium hydrogen carbonate) was studied by electrochemical polarization measurements. The calculated corrosion parameters suggest that the thiadiazole derivative reduces the rate of bronze corrosion, acting as a mixed-type inhibitor in all the investigated electrolytes. The best inhibiting efficiency of the thiadiazole derivative was observed in sodium chloride solution.

Keywords: bronze corrosion; corrosion inhibitor; thiadiazole derivatives

1. Introduction

Copper and copper-based alloys are widely used in a great variety of applications, such as heat exchangers, building construction, electronics, coinage, art works etc. [1]. In spite of the fact that copper is a relatively noble metal, it corrodes easily in oxygen-containing electrolytes [2]. The corrosion behavior of copper and its alloys in various environments (acidic, alkaline and neutral) has been explored for several decades and, in all cases the dissolution of copper was balanced by the oxygen reduction [3].

Since copper and its alloys are not stable in oxygen-containing electrolytes, substantial improvement in their passivity is needed [4]. To this regard, a range of organic compounds can offer a good protection against corrosion to the exposed copper and bronze surfaces in different aggressive environments. Thus, it is well-known that, on copper and bronze, heterocyclic compounds containing nitrogen, oxygen or sulfur [5-10] are efficient corrosion inhibitors due to the chelating action of heterocyclic molecules and the formation of a physical blocking barrier on the surface [7]. In particular, benzotriazole and its derivatives are among the most effective inhibitors for

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copper and copper alloys corrosion over a wide temperature and pH range [7, 11-13], but because they are highly toxic [10, 14] they should be replaced with other environmentally friendly inhibitors.

In an attempt to find non-toxic alternatives to the conventional benzotriazole, recent works showed that 2-amino-5-mercapto-1,3,4-thiadiazole is an excellent corrosion inhibitor for copper and bronze in citric acid [15, 16]. In this context, the present work was carried out to investigate the effect of a new thiadiazole derivative, *i.e.* 2 mercapto-5-R-acetylamino-1, 3, 4-thiadiazole on bronze corrosion in different aqueous electrolytes. The efficiency of the inhibitor was studied by electrochemical polarization measurements.

2. Experimental

Materials

Experiments were carried out in different 0.01 M aerated electrolytes (Table 1), at room temperature.

Table 1.

Characteristics of the corrosive media

Electrolyte	NaCl	Na ₂ SO ₄	NaHCO ₃
рН	5.46	5.76	8.77
λ 10⁻³ (S∕cm)	1.243	2.058	1.073

2 mercapto-5-R-acetylamino-1, 3, 4-thiadiazole was synthetized in the laboratories of the Faculty of Pharmacy from Cluj-Napoca, Romania and its molecular structure is presented in figure 1.



Figure 1. Molecular structure of the inhibitor

In all the investigated aqueous electrolytes, the concentration range of the inhibitor was 10^{-5} to 10^{-3} M.

All the other chemicals were of analytical grade and used as received.

Methods

Potentiodynamic polarization measurements were conducted using an electrochemical analyzer (Autolab-PGSTAT 10, EcoChemie, Utrecht, Netherlands) connected to a PC for potential control and data acquisition. STUDY ON THE INHIBITING BEHAVIOR OF A NON-TOXIC THIADIAZOLE DERIVATIVE...

Chronopotentiometry was used for the open-circuit potential (O.C.P) measurements.

The electrochemical experiments were performed in a three-electrode cell with a separate compartment for the reference electrode connected with the main compartment *via* a Luggin capillary. The working electrode was made of bronze (see its composition in Table 2), the reference electrode was a saturated calomel electrode (SCE) and the counter electrode was a platinum foil.

Table 2.

Weight % composition of the bronze electrode

Cu	Sn	Pb	Zn	Sb	Ni	Fe	Mn	As	S
87.975	6.014	4.020	1.172	0.299	0.181	0.11	0.002	0.033	0.19
Si	Р								
0.004	0.004								

Prior to all the polarization runs, the bronze electrode was polished with grit paper of 600, rinsed with distilled water and ethanol, and immersed for 60 minutes in the electrolyte. Anodic and cathodic curves were recorded in a potential range of $E = E_{corr} \pm 200 \text{ mV}$ with a scan rate of 0.25 mV s⁻¹. The rotation speed of the working electrode was fixed at 2000 rpm.

3. Results and Discussion

3.1. Open-Circuit Potential (O.C.P) Measurements

The way in which a metal changes its O.C.P. with time upon immersion in a solution could bring information about the reactions taking place at the metal solution interface. In general, a shift of potential towards positive direction denotes passivation behaviour, whilst a shift in the negative direction signifies activation behaviour.

The open-circuit potentials evolution in time for the bronze electrode, recorded immediately after 60 minutes immersion in different electrolytes are presented in Figure 2.

In the absence of the inhibitor, the O.C.P values for the sulphate and hydrogen carbonate electrolytes gradually increase in the positive direction during the first minutes to reach a stationary state characterized by -0.102 V vs. SCE and -0.069 V vs. SCE, respectively. This passive behaviour could be most probably attributed to the chemisorption of the dissolved oxygen and its reduction together with possible formation of oxide layers and/or hydroxysulfate or hydroxycarbonate on the surface of the electrode [17].

In the chloride medium, the OCP stabilized very quickly after the immersion at -0.379 V vs. SCE. Cuprite (Cu₂O), nantokite (CuCl) and tin species (hydroxides and/or hydroxychlorides) could be produced in this case on the bronze surface [16].

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Disregarding the electrolyte composition, more anodic values of the O.C.P. are observed in the presence of thiadiazole derivative. The positive potential shift could be associated with the formation of protective film at the electrode surface. This behaviour suggests that, probably, 2 mercapto-5-R-acetylamino-1, 3, 4-thiadiazole is adsorbed on the bronze surface, acting mainly as an inhibitor for bronze corrosion in all the investigated aqueous media.



Figure 2.

Open-circuit potentials evolution for the bronze electrode immersed in various 0.01 M aqueous electrolytes without and with different concentrations of 2 mercapto-5-R-acetylamino-1, 3, 4-thiadiazole: (A) NaCl; (B) Na₂SO₄; (C) NaHCO₃.

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3.2. Potentiodynamic Polarization Measurements

The cathodic and anodic polarization curves of bronze electrodes recorded after its immersion during 60 minutes in different electrolytes in the absence and in the presence of 2 mercapto-5-R-acetylamino-1, 3, 4-thiadiazole are shown in Figure 3.



Figure 3. Influence of thiadiazole derivative concentration on the anodic and cathodic polarization curves of bronze in various aqueous 0.01 M electrolytes: (A) NaCl; (B) Na₂SO₄; (C) NaHCO₃.

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As can be observed, the anodic curves in NaCl solution are characterized by the presence of a shoulder, which has been attributed to the formation of CuCl on the metal surface [19], according to the reactions [20]:

$$Cu + Cl^{-} \leftrightarrow CuCl + e^{-}$$
$$CuCl + Cl^{-} \leftrightarrow CuCl_{2}^{-}$$

It is generally accepted that, in the apparent Tafel region, the corrosion process is controlled by the electrodissolution of copper and the diffusion of soluble $CuCl_2^-$ from outer Helmholtz plane into the bulk solution (mixed control) [21].

Disregarding the electrolyte composition, the addition of the inhibitor determines a considerably decrease both in the anodic and cathodic current density values with respect to the blank solutions. This indicates that the tested thiadiazole derivative cannot be designed exclusively as anodic or cathodic inhibitor. It acts as a mixed-type inhibitor that protects the electrodic surface, and hinders both the anodic dissolution of copper and the cathodic oxygen reduction. However, the slight shift of the corrosion potential values in the negative direction observed in the presence of the inhibitor points to a small predominance of the cathodic reaction.

The corrosion parameters were calculated directly from the potentiodynamic polarization measurements on the basis of the Tafel extrapolation, according to the equation:

$$i = i_0 \exp\left[\frac{2,303}{b_a}(E - E_{corr})\right] - \exp\left[-\frac{2,303}{b_c}(E - E_{corr})\right]$$
(1)

where b_a and b_c are anodic and cathodic Tafel slopes, $E_{corr is}$ the corrosion potential and the other parameters have the usual meaning.

The corrosion inhibition efficiency (IE) in the presence of the inhibitor was calculated from the relation:

$$\mathsf{IE} = \frac{i_{corr} - i_{corr}}{i_{corr}} \cdot 100 \,[\%] \tag{2}$$

where i_{corr} and i'_{corr} are the corrosion current densities in the absence and in the presence of the inhibitor, respectively.

The values of the corrosion potential (E_{corr}), corrosion current density (i_{corr}), inhibition efficiency (IE) and of the anodic and cathodic Tafel

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slopes (b_a and b_c) for bronze in different electrolytes as a function of the inhibitor concentration are presented in Table 3.

As it can be seen, the corrosion potentials dependence on the inhibitor concentration is not a monotonic one: at very low concentrations (below 10⁻³ M), the inhibitor shifts the corrosion potential slightly to the cathodic direction, while at higher inhibitor concentrations, the corrosion potential shifts to the anodic direction. These variations of the corrosion potential values are probably due to a competition between the anodic and the cathodic inhibiting reactions and/or to the metal surface condition. Moreover, in the presence of 2 mercapto-5-R-acetylamino-1, 3, 4-thiadiazole both the anodic and cathodic Tafel slopes change with the inhibitor concentration, indicating once more that both reactions are influenced by the inhibitor presence.

Relatively high values of the corrosion inhibition efficiency are observed in all solutions. However, it should be noticed that the thiadiazole derivative has a better inhibition effect on bronze corrosion in chloride electrolyte as compared to sulphate or hydrogen carbonate solutions. For chloride and sulphate electrolytes, the optimum concentration for the investigated thiadiazole derivative with the corresponding inhibiting efficiency was 10^{-4} M. In the case of hydrogen carbonate solution, the optimum concentration of inhibitor was 10^{-3} M for an efficiency of around 82 % (table 3).

Table 3.

Electrolyte	Inhibitor	Ecorr	İ _{corr}	bc	ba	IE
	conc.					
(0.01 M)	(M)	(mV/SCE)	(10 ⁷ Acm ⁻²)	(V dec ⁻¹)	(V dec ⁻¹)	(%)
NaCl	0	-96	41.14	0.115	0.057	-
	10 ⁻⁵	-170	4.28	0.096	0.047	89.60
	10 ⁻⁴	-256	3.43	0.109	0.045	91.66
	10 ⁻³	-98	3.72	0.124	0.109	90.95
Na₂SO₄	0	-48	23.30	0.109	0.037	-
	10 ⁻⁵	-60	7.89	0.098	0.049	66.13
	10 ⁻⁴	-62	4.30	0.110	0.032	81.54
	10 ⁻³	-50	4.38	0.149	0.077	81.20
NaHCO₃	0	-87	7.31	0.122	0.045	-
	10 ⁻⁵	-97	3.02	0.104	0.080	58.68
	10 ⁻⁴	-116	3.53	0.125	0.083	51.71
	10 ⁻³	-110	1.31	0.116	0.073	82.08
1					1	

Corrosion parameters for bronze in different electrolytes in the absence and in the presence of 2 mercapto-5-R-acetylamino-1, 3, 4-thiadiazole at room temperature.

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A possible explanation for the inhibiting action of 2 mercapto-5-Racetylamino-1, 3, 4-thiadiazole on the bronze corrosion could be found in the covering of the electrodic surface with a protective film made of a polymeric Cu(I)-inhibitor complex. According to [15], the efficiency of the organic inhibitors could be associated to the extent to which they adsorb and cover the metal surface. Their adsorption depends on the structure of inhibitors, on the surface charge of the metal, and on the type of the electrolyte. 2 mercapto-5-R-acetylamino-1, 3, 4-thiadiazole acts as a bidentate ligand both through the aminic and the closed ring nitrogen in the complex.

4. Conclusions

The results of electrochemical investigation show that 2 mercapto-5-Racetylamino-1, 3, 4-thiadiazole has fairly good inhibiting properties for bronze corrosion in all the investigated electrolytes. It behaves as a mixed-type inhibitor.

The maximum inhibition efficiency of 2 mercapto-5-R-acetylamino-1, 3, 4-thiadiazole was observed in chloride electrolytes. Its inhibiting action can be attributed mainly to the blockage of the bronze surface by a protective film consisting of a polymeric Cu(I)-inhibitor complex in which the inhibitor acts as a bidentate ligand through the aminic nitrogen atom and the closed ring nitrogen.

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ESTIMATIONS OF SURFACE FREE ENERGIES FOR SOLID METALS

OSSI HOROVITZ¹, ANCA-LIANA POPOVICIU¹, ELENA-MARIA PICA²

ABSTRACT. Experimental methods rarely give reliable values of the free surface energies/surface tensions for solid metals and just few data exist. Therefore correlations between surface and bulk thermodynamic properties are important in order to obtain estimated values for the surface quantities. From experimental values, molar free surface energies are calculated on the basis of molar volumes or atomic radii of metallic elements. The bulk properties considered here are melting point, enthalpy of atomization, free enthalpy of atomization and inner free enthalpy of atomization. Different correlations are investigated and the results are discussed and compared with the predictions of the rule of broken bonds between first neighbour atoms. The deviations of some elements from the regression lines are examined. From the correlations, a set of calculated free surface enthalpies for 69 solid elements is proposed.

Keywords: metals, surface free energy, enthalpy of atomization, correlations

Introduction

Surface free energy is one of the most important properties of solid state and particularly of solid metals. It determines the equilibrium shape of crystals and plays a decisive role in surface phenomena and processes, such as surface faceting, roughening, surface segregation in binary alloys, crystal growth, formation of grain boundaries, growth and stability of thin films, the shape of small crystallites in a supported catalyst, adsorption, sintering. The surface free energies between two interacting surfaces control processes like the stability of aqueous colloidal suspensions, the dynamics of molecular selfassembly, wetting, spreading, deinking, adhesion. Many mineral processing techniques, e.g. froth flotation, selective flocculation, filtration and thickening depend on the solid/liquid interfacial interaction [1].

Surface free energy, γ is defined as the excess Helmholtz energy (free energy), or the excess Gibbs energy (free enthalpy) per unit area:

$$\gamma = F_{\rm s} = G_{\rm s} = \Delta F/A_{\rm s} = \Delta G/A_{\rm s} \tag{1}$$

For liquids it is numerically and dimensionally equal to the *surface tension* (*interfacial tension*), Some authors consider the two notions to be synonymous for solids too and related to the work spent in forming the surface,

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and introduce the *surface stress* or *stretching tension* in order to describe the work spent in stretching the surface [2, 3]. The first is a scalar quantity, the second - a tensor. The difference comes from the reduced mobility of atoms in solids, which prevents them to quickly rearrange to an equilibrium position when a new surface is created, while liquids cannot maintain a strain. However, this mobility is increased near the melting point, where this difference should disappear. As the temperature is approaching the melting point, a roughening of the surface occurs and even a "surface melting" can take place.

The temperature dependence of the surface free energy is related to the surface entropy, $S_s = \Delta S/A_s$, A_s being the surface area, the entropy variation involved in the creation of the unit area of the solid-vapour interface, by the obvious relation:

$$S_{s} = -\left(\frac{\partial G_{s}}{\partial T}\right)_{p} = -\left(\frac{\partial \gamma}{\partial T}\right)_{p}$$
(2)

and the surface enthalpy (or energy) can be obtained from the Gibbs-Helmholtz equation:

$$H_{s} = \gamma - T \frac{\partial \gamma}{\partial T}$$
(3)

So-called "molar" surface quantities are defined as products of the above functions with the "molar" surface, $V_M^{2/3}$ (V_M – the molar volume of the solid), e.g. the "molar" surface free energy $\gamma V_M^{2/3}$. For liquids, a linear decrease of surface tension with temperature increase is generally assumed, i.e. surface entropy is constant, independent of temperature. In the case of solid metals, exact measurements on the temperature dependence of surface free energy are missing, but here also a decrease of γ with increasing temperature is observed.

Since experimental methods rarely give reliable values of the free surface energies for solid metals and just few data exist, there were many attempts to calculate the surface energy. They range from empirical correlations between this quantity and some bulk properties of the metal to the use of *abinitio* techniques, passing through semi-empirical methods, among them many applications of the law of corresponding states and statistical mechanics [4].

Empirical Correlations Between Free Surface Energy and Other Properties of Solid Metals

In order to find correlations between free surface energies and other properties of the metals, experimental values were used (Table 1), taken mostly from the compilation of Kumikov and Khokonov [5]. The authors

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have selected, compared and evaluated experimental values, obtained by various methods. These measurements were made at different, most of

Table 1.

Surface free energies and other properties of solid metals

Element	ν ^{exp}	ν TM	T _m [K]	ΛHa	V[cm ³]	R
	[mJ.m⁻²]	[mJ.m ⁻²]		[kJ/mol]		[pm]
Li		472	454	159.4	13.096	156
Ве	1000	1298	1551	324.3	4.875	113
Na		234	370.9	108.4	23.686	186
Ma		688	922	147.7	13.981	160
AĬ	1140	1020	933.5	326.4	9.985	143
Si	1200	940	1683	455.6	12.051	117
к		129	337	89.2	45.348	231
Ca		425	1112	178.3	25.851	197
Sc			1814	377.8	15.033	161
Ti	1938	1749	1941	469.9	10.546	145
V	2900	2301	2173	514.2	8.782	131
Cr	2090	2006	2130	396.6	7.230	125
Mn	1400	1298	1518	280.7	7.393	137
Fe	2170	2123	1808	416.3	7.095	125
Co	2424	2218	1768	424.7	6.621	125
Ni	1940	2080	1726	429.7	6.594	124
Cu	1520	1566	1356.4	338.3	7.123	128
Zn	868	896	693	130.7	9.158	133
Ga	767	845	302.9	277	11.70	122
Ge	820	748	1210.4	376.6	13.64	123
Rb		104	312.2	80.9	55.856	243
Sr		358	1041	164.4	33.7	215
Y			1795	421.3	19.888	180
Zr	1850	1687	2125	608.8	14.055	161
Nb	2400	2313	2741	725.9	11.059	147
Мо	2630	2510	2883	658.1	9.405	136
Тс			2445	678	8.513	135
Ru	3400	2655	2607	642.7	8.217	132
Rh	2800	2325	2239	556.9	8.298	134
Pd	2000	1743	1825	378.3	8.868	138
Ag	1205	1086	1235	384.6	10.272	144
Cd	675	698	594.1	112	12.995	149
In	633	658	429.3	243.3	15.869	163
Sn	673	661	505	302	16.306	140
Cs		84	301.6	76.1	70.955	265
Ва		326	998	180	39.122	210
La			1193	427	20.731	187
Nd		812	1297	328	20.605	181
Pm			1312	301	22.380	183
Sm			1345	206.7	19.941	180
Eu			1095	178	28.895	204
Gd	900		1586	398	19.936	179
Yb			1097	152	24.790	194

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Lu			1936	428	17.781	174
Hf		1923	2495	669	13.521	154
Ta 2	2480	2493	3269	782	10.900	143
W 2	2900	2765	3683	849.4	9.476	137
Re 2	2200	3133	3453	769.9	8.854	138
Os		2950	3327	790.8	8.492	134
Ir		2655	2637	665.2	8.573	136
Pt 1	950	2203	2045	565.3	9.094	139
Au 1	410	1333	1337.6	369	10.210	144
TI 5	562	550	576.7	182.2	17.710	170
Pb 5	560	540	601	196	18.271	175
Bi 5	501	446	544.5	207.1	21.324	155
Ra			973	161.9	44.403	132
Th			2023	598.3	19.832	181
Pa			1873	606.7		161
U		1780	1405	535.6	12.494	138

 γ^{exp} – experimental values for free surface energy [5], γ^{TM} – Tyson and Miller values [6], T_m – melting point, ΔH_a – enthalpy of atomization, V – molar volume, R – metallic radius

them at relatively high, temperatures, near the melting point of the metal. The preferred values are mostly those obtained by the compensated zero-creep method. The scatter of the values is important, the relative errors of the measurements reach 10-20%, but only 1-3% in the zero-creep method. We selected the most probable values (most of them recommended by the authors) along with some values from other sources. Insofar as possible, we selected values obtained at lower temperature, where the peculiarities of the solid metals are more pronounced.

In literature the "experimental" values of Tyson and Miller [6] are often used. As a matter of fact, these are extrapolated from the values for liquid metals at the melting point, γ_L which are reasonable well known quantities. They inferred that the ratio of free surface energies for solids γ_s and for liquids should be approximately constant for metallic elements:

$$\gamma_{\rm S} / \gamma_{\rm L} = 1.18 \pm 0.03$$
 (4)

The values given by Tyson and Miller are also included in Table 1.

Since surface free energy of a pure solid is determined by the strength of cohesive forces between the adjacent particles, it seems logically to expect correlations between it or other surface properties and some physicochemical bulk properties of the materials, that are also the result of forces of this kind. Of course, this should apply only to pure materials, inasmuch as even the smallest amounts of surface active contaminants have a strong influence on the values of surface tension, while the bulk properties are only

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slightly modified. The bulk properties considered here are melting temperatures and standard enthalpies of atomization.



Figure 1. Surface free energies of solid metals versus melting temperatures



Figure 2. Surface free energies of solid metals versus enthalpies of atomization

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Experimental values for the surface free energies of solid metals were plotted versus melting points T_m in Figure 1. The two properties appear to be related, but there is not a strong linear correlation. Particularly, metals with high melting points (Nb, Mo, Ta, Re, W) don't have correspondingly high γ -values. The same is the case for the semi-metals Ge and Si. On the other hand, the surface free energy for Ga is unexpectedly high for its melting point. The correlation line for 33 experimental values corresponds to the equation:

$$\gamma [mJ/m^2] = (313\pm157) + (0.791\pm0.083)T_m[K], r = 0.864$$
 (5)

Another measure of cohesion forces in the metal lattice could be the standard enthalpy of atomization, ΔH_a (the values are also given in Table 1). As in the case of the temperature dependence, an increase of the free surface energies with increasing ΔH_a -values is evident from Figure 2, but there is no linear correlation, the scatter of the points is even higher than in Fig.1. The equation of the correlation line, passing through the origin, is:

$$\gamma$$
[mJ/m²] = (3.80±0.18) Δ H_a[kJ/mol] r = 0.819, n = 33 (6)

One of the reasons for the poor correlation is that, while surface free energies refer to the surface unit, the enthalpies of atomization are molar quantities. Therefore, "molar" surface tensions γ .V^{2/3} should be used, where V stands for the molar volume of the solid metal, calculated as the ratio of the atomic weight to the density of the solid, M/ ρ_s . Calculated values for the molar volumes are given in Table 1. The plot of this quantity versus melting point is given in Figure 3. The correlation is somewhat better as in Fig.1 (eq. 5), but not satisfactory. The equation of the correlation line is:

$$10^{4}\gamma$$
.V^{2/3} [mJ] = (2160±580) + (3.16±0.31) T_m r = 0.880 n = 33 (7)

The greatest deviations form the fitting line are those for Be and Re (negative deviations) and for V, Ru (positive deviations). Excluding the evidently wrong values for Be and Re, the equation of the new correlation line is:

$$10^{4}\gamma.V^{2/3}$$
 [mJ] = (950±460) + (3.44±0.25) T_m r = 0.931 n = 31 (7a)

and without the uncertain values for V and Ru also:

$$10^{4}\gamma$$
.V^{2/3} [mJ] = (2100±350) + (3.22±0.20) T_m r = 0.953 n = 29 (7b)

i.e. a quite good correlation. 142

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Subsequently "molar" surface tensions were plotted versus enthalpies of atomization (Figure 4). The correlation line is given by the equation:

$$10^{4}\gamma$$
.V^{2/3} [mJ] = (17.12±0.57) Δ H_a[kJ/mol] $r = 0.894$ $n = 33$ (8)



Figure 3. "Molar" surface tensions of solid metals versus melting points



Figure 4. "Molar" surface tensions of solid metals versus enthalpies of atomization

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The most important deviations are caused by the same elements as in the previous representation. Without Be and Re, the equation becomes:

$$10^{4}\gamma.V^{2/3}$$
 [mJ] = (17.70±0.51) ΔH_{a} [kJ/mol] $r = 0.923, n = 31$ (8a)

and letting apart V and Ru too:

The general appearance of the plot is very similar to that on Figure 3, the correlation is quite the same. The introduction of molar surface tensions leads therefore to a better correlation as compared to surface tensions, but the scatter of the points is still important. Some reasons could be the imprecision of experimental values, the different types of crystal lattices in different metals, the fact that the experimental data were measured at temperatures sometimes quite far from the standard temperature, and the nature of molar surface tensions, these being essentially *free* energies, other than the enthalpies of atomization.

Two metals giving here deviations (but not in Fig.2) are Zn and Cd: their $\gamma V^{2'3}$ -values appear to be too high as compared to their enthalpies of vaporization. This could be related to peculiarities of their crystal structure. They have a hexagonal closest packed structure, but the height of their unit cell is some 15% larger than normal for this type of crystals, i.e. the 6 adjacent atoms in the basal plane are nearer to one another as the other 6, in the next planes above or bellow. The interatomic distances are 2.660 and 2.907 Å for Zn, respectively 2.973 and 3.287 for Cd [7]. This could diminish the cohesion energy, i.e the enthalpy of atomization, while in the surface layer, these greater distances probably vanish. Besides, V^{2/3} could give here a too large evaluation for the molar surface.

For comparison, the correlations were tested also with the values proposed by Tyson and Miller (Table 1). The corresponding "molar" quantities $\gamma^{TM}V^{2/3}$ were calculated for 45 elements and plotted versus melting points (Figure 5). The correlation line is described by the equation:

$$10^{4}\gamma^{1M}V^{2/3}$$
 [mJ] = (1350±330) + (3.42±0.18) T_m r = 0.944, n=45 (9)

The corresponding representation against standard enthalpies of atomization is given in Figure 6. The equation of the correlation line is:

$$10^4 \gamma^{\text{TM}} V^{2/3} \text{ [mJ]} = (16.60 \pm 0.31) \Delta H_a \text{[kJ/mol]} r = 0.969, n = 45$$
 (10)

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The much better correlation in both cases is a consequence of the higher precision of experimental data for liquid metals, the values of Tyson and Miller are based on, and of the disappearance of differences contingent on the crystal lattice near the melting point as well. Some larger deviations are those for Be, Ge and Si, accountable by their semi-metallic character and the presence of covalent binding. In Fig. 5 the large deviation for U is striking; caused by the unusual high melting temperature of this metal. In Fig. 6 this point is not longer an outlier; enthalpies of atomisation are better correlated to surface tensions than melting points.



Fig.5. Molar surface tensions (Tyson and Miller) versus melting points



Molar surface tensions (Tyson and Miller) versus enthalpies of atomization

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The points for 2nd group elements in the periodic system (Mg, Ca, Sr, Ba) and for 12th group elements (Zn, Cd) are somewhat apart, over the correlation line; they all correspond to nearly the same value of "molar" surface tension. Such deviations appear also for the melts of these metals, in the correlation of their molar surface tensions with enthalpies of vaporization [8].

The experimental value for berrylium ($\gamma = 1000 \text{ mJ/m}^2$) is evidently to low; it was determined by a single method [5] and is much under the value given by Tyson and Miller (1298). But even with this value, the point is clearly under the correlation line. Mezey and Giber [9] calculated for Be a γ -value of 1117; the value of Tyson and Miller extrapolated for 0 K is 1628, Boer et al. [10] estimated a value as high as 2700, while an *ab initio* calculation [11] gives $\gamma = 1834$.

The point for rhenium in Fig. 3 and 4 is far under the line, while the calculated value of Tyson and Miller gives a point near the line (Fig. 5, 6). The experimental value (2200 mJ/m²) coming from a single determination (FEM) [5] is apparently wrong, the value extrapolated by Tyson and Miller (3133), and that calculated by Mezey [9] (3109) are much higher. Even larger are the values given by Boer [10]: 3600 and calculated ab initio [11]: 4214.

For V and Ru the points in Fig. 3 and 4 are far above the correlation line, while with the values of Tyson and Miller there are not such deviations. For both metals the experimental γ -values [5] are much higher than those given by Tyson and Miller and even than those given by Boer, but are in good agreement with the calculations of Mezey and ab initio (for V), while the results of the ab initio calculation for Ru are still higher. Therefore, our linear regression calculations were conducted as well without these four elements.

As a measure for the area occupied by an atom in the surface layer, we could select also the square of the metallic radius of the element R (Table 1), even though in the surface shell the positions of the atoms are somewhat different to those inside the metal lattice. The differences resulted from the reconstruction of the surface should however affect interatomic distances between different planes rather than distances within the same plane. The product γ .R² as an alternative to the "molar" surface tension, was thus plotted against the enthalpy of atomization in Figure 7, with experimental γ -values. The equation of the correlation line is:

$$10^{24}$$
.γ.R²[J] = (71.9 ± 2.9)∆H_a[kJ] r = 0.86, n = 33 (11)

This correlation is slightly worse than the corresponding one with γ .V^{2/3} (Fig. 4, eq. 8), but the distribution of the points is very similar. The great deviations of the point for Ge and Si from the correlation line are evident. The reason could consist in the use of covalent radii for these elements, while metallic radii were used for all the other elements.

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Excluding again Be, Re, V and Ru, the equation becomes:

$$10^{24}$$
. γ .R²[J] = (71.7 ± 2.8) Δ H_a[kJ] r = 0.88, n = 29 (11a)



Fig.7. γ .R² products against enthalpies of atomization



 γ .R² products against enthalpies of atomization (γ from Tyson and Miller)

without a significant improvement for this once.

The similar plot with the values given by Tyson and Miller (Figure 8) corresponds to the correlation line:

$$10^{24} \gamma^{\text{TM}} \cdot R^2[J] = (67.9 \pm 2.2) \Delta H_a[kJ] r = 0.90, n = 45$$
 (12)

The correlation is also somewhat worse than for "molar" surface tensions (Fig. 6, eq. 10), while the general form is the same. The greater deviations correspond again to elements whose radii don't fit the pattern of metallic radii (Ge, Si).

Many approaches in the evaluation of surface energy make use of the broken-bond rule, according to which the surface energies scale with the number of broken bonds between first neighbours, when bulk atoms become surface atoms [12]. If only interactions between pairs of adjacent atoms are considered, the energy needed to bring atoms from the bulk to the surface should be proportional to the fraction α of broken bonds. Usually the ratio of surface energy (or enthalpy) to the enthalpy of vaporization for liquids or the enthalpy of atomization for solids is considered to be given by α . But above, we had free surface energies correlated to bulk energies. In order to have the same kind of quantities, we could either use instead of the molar surface free energies (surface tensions) the molar surface enthalpies (3) or energies

$$U_{sM} = \gamma_M + T.S_{sM}$$
(13)

or instead of standard enthalpies of atomisation the corresponding free enthalpies

$$\Delta G_a = \Delta H_a - T \Delta S_a \tag{14}$$

 S_{sM} is the molar surface entropy (cf. Eq.(2)). In correlating surface and bulk properties of melted metals, the first way was chosen [8], since the temperature coefficients of the surface tension were reasonable well known and the molar surface entropy is nearly a constant, whatever the metal or the temperature were [13]. However for solid metals the temperature coefficient of surface tension was rarely determined and its constancy is questionable. For liquids, this constancy is related to constant value of the entropies of vaporization (Trouton's rule), while melting entropies are not constant, and atomization entropies should not be expected to be constant. The calculation of surface energies of solid metals would thus be very intricate. We preferred therefore the second way, to estimate free enthalpies of atomization (10).

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The standard atomization entropy for solid metals can be calculated as the difference:

$$\Delta S_{a} = S_{298}^{0} (gas) - S_{298}^{0} (solid)$$
(15)

Standard entropies of solid metals are well known quantities and are given in Table 2. The entropies for gaseous metals can be calculated

Table 2.

$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	i 1
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	
IN IN<	
L1 29.1 138.8 109.7 128.7 -	
Be 9.5 142.0 132.5 284.8 1096 305.92 10.30 20.62 0.087 Na 51.3 153.7 102.4 77.9	
Ma 51.3 153.7 102.4 77.9	
AI 28.3 155.7 127.4 288.4 1180 307.77 17.78 46.22 0.150	
Si 18.8 156.2 137.4 414.7 1200 433.53 14.55 53.26 0.123	
K 64.7 160.3 95.6 60.7	
Ca 41.6 160.6 119.0 142.7	
Sc 34.6 162.1 127.5 339.8	
Ti 30.7 162.8 132.1 430.5 2211 449.72 23.64 89.78 0.200	
V 28.9 163.6 134.7 474.1 2900 493.71 25.00 104.24 0.211	
Cr 23.76 163.9 140.1 354.8 2488 374.98 24.81 78.55 0.209	
Mn 32 164.6 132.6 241.2 1400 261.28 20.33 44.87 0.172	
Fe 27.3 164.8 137.5 375.3 2550 395.52 23.80 79.50 0.201	
Co 30 165.4 135.4 384.3 2859 404.69 24.90 85.11 0.210	
Ni 29.9 165.4 135.5 389.3 2304 409.69 19.77 68.41 0.156	
Cu 33.2 166.4 133.2 298.6 1770 318.78 20.55 55.33 0.174	
Zn 41.6 166.7 125.1 93.4 976 112.96 37.83 36.08 0.319	
Ga 40.9 167.5 126.6 239.3 762 258.20 15.21 33.16 0.128	
Ge 31.1 168.0 136.9 335.8 966 354.40 15.57 46.60 0.131	
Rb 76.8 170.1 93.3 53.1	
Sr 52.3 170.4 118.1 129.2	
Y 44.4 170.6 126.2 383.7	
Zr 39 170.9 131.9 569.5 2215 587.99 22.01 109.27 0.186	
Nb 36.4 171.1 134.7 685.8 2667 704.84 18.78 111.79 0.159	
Mo 28.7 171.5 142.8 615.5 3124 635.02 21.92 117.59 0.185	
Tc 33.5 171.8 138.3 636.8	
Ru 28.5 172.2 143.7 599.9 3400 619.70 22.15 116.91 0.189	
Rh 31.5 172.4 140.9 514.9 2800 534.71 21.46 96.91 0.181	
Pd 37.6 172.8 135.2 337.9 2000 357.54 23.96 72.36 0.202	
Ag 42.6 173.0 130.4 245.7 1393 265.01 24.83 55.58 0.210	
Cd 51.8 173.5 121.7 75.7 745 94.42 43.59 34.76 0.368	
In 57.8 173.7 115.9 208.8 668 226.95 18.58 35.61 0.157	
ISn 51 2 174 2 123 0 265 4 727 283 47 17 36 39 48 0 139	
ICs 852 1756 904 492	
Ba 62.8 176.0 113.2 146.3	

Experimental and calculated properties of solid metals

La	l	56.9	176.1	119.2	391.5					
Nc	ł	71.5	176.6	105.1	296.7					
Pn	n		176.7							
Sn	n	69.6	177.1	107.5	174.7					
Eu	1	77.8	177.2	99.4	148.4					
Go	t	68.1	177.7	109.6	365.3	900	382.96	17.28	55.88	0.146
Yb)	59.9	178.9	119.0	116.5					
Lu		51	179.0	128.0	389.9					
Hf		43.6	179.3	135.7	628.6					
Та	I	41.5	179.4	137.9	740.9	2957	760.01	19.13	122.76	0.162
W		32.76	179.6	146.9	805.6	3198	825.10	17.36	120.95	0.147
Re	9	36.9	179.8	142.9	727.3	2679	746.95	15.35	96.81	0.130
Os	3	32.6	180.1	147.5	746.9					
lr		35.5	180.2	144.7	622.1					
Pt		41.6	180.4	138.8	523.9	2318	543.51	18.58	85.31	0.157
Au	I	47.4	180.5	133.1	329.3	1641	348.62	22.15	65.20	0.187
ΤI		64.2	180.9	116.7	147.4	616	165.32	25.33	35.37	0.214
Pb)	64.8	181.1	116.3	161.3	619	179.17	23.96	36.26	0.202
Bi		56.7	181.2	124.5	170.0	547	187.45	22.43	35.50	0.189
Ra	a	71.1	182.0	110.9	128.9					
Th	1	51.8	182.5	130.7	559.3					
Pa	1	51.9								
U		50.2	182.8	132.6	496.1					

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Standard entropies of solid, $S_{298,s}^{0,0}$, and gas, $S_{298,g}^{0,0}$; ΔS_a - standard entropies of atomization, ΔG_a – free enthalpies of atomization, γ_{298} - γ values extrapolated for 298 K, ΔG_a^{i} – internal free enthalpies of atomization, k = $\gamma_{298} V^{2/3} / \Delta G_a^{i}$, γ_M – molar surface free energy.

by the methods of statistical thermodynamics. Since gaseous metals are monoatomic, the equation of Sackur und Tetrode [14] can be applied:

$$S(T, p) = \frac{5}{2}R\ln T - R\ln p + \frac{5}{2}R + R\ln \frac{(2\pi m)^{3/2}k^{5/2}g_0}{h^3}$$
(16)

The entropies calculated by this formula are given in Table 2, together with the calculated values for the entropy of atomization (15) and the standard free enthalpies of atomization (14)

The plot of molar free surface energy against these calculated free enthalpies of atomization is similar to that in Fig. 4. The correlation line corresponds to the equation:

$$10^{4}\gamma V^{2/3}$$
 [mJ] = (18.53±0.66) ΔG_{a} [kJ/mol] $r = 0.893, n = 33$ (17)

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The correlation is just the same as for enthalpies of atomisation (8), the introduction of free enthalpies did not improve it. The most important deviations are again for Be and Re (wrong experimental data) and for V and Ru (uncertain experimental values). Without the first two elements we obtain:

$$10^{4}\gamma V^{2/3}$$
 [mJ] = (19.20±0.60) ΔG_{a} [kJ/mol] $r = 0.922, n = 31$ (17a)

and excluding all the 4 elements:

$$10^{4}\gamma V^{2/3}$$
 [mJ] = (18.60±0.56) ΔG_{a} [kJ/mol] $r = 0.945, n = 29$ (17b)

The same correlation was verified for the values proposed by Tyson and Miller. The equation of the correlation line is:

$$10^4$$
γTMV^{2/3} [mJ] = (17.92±0.39)∆G_a[kJ/mol] r = 0.969, n = 45 (18)

The image is very similar to the one in Figure 6 and the coefficient of correlation is just the same.

The correlation with free enthalpies of atomization was also tested using R^2 as a measure of atomic surface. The correlation described by the equation:

$$10^{24}\gamma.R^{2}[J] = (77.89 \pm 3.23)\Delta G_{a}[kJ] r = 0.860, n = 33$$
(19)

does not differ from the one using ΔH_a (Fig. 7, eq. 11). The exclusion of the 4 doubtful values (for Be, Re, V. Ru) does not result in an obvious improvement:

$$10^{24}\gamma$$
.R²[J] = (77.85 ± 3.20) Δ G_a[kJ] *r* = 0.881, *n* = 29 (19a)

Even though correlation is not improved by using free enthalpies of atomization instead of enthalpies of atomization, this treatment is theoretically legitimate.

The experimental values of surface free energies for solid metals were determined at very different temperatures. In order to compare them with ΔG_a it would be appropriate to refer them all to the standard temperature (298K). On the basis of the theory of corresponding states, Digilov [15] deduced an equation for the temperature dependence of surface tensions for solid metals, starting with the surface tension of the liquid at the melting point:

$$\gamma_{s}(T) \cong 1,2\gamma_{L} \left[1 - 0,229 \left(\frac{T}{T_{m}} - 1 \right) + 0.01 \left(\frac{T}{T_{m}} - 1 \right)^{2} \right]$$
 (20)

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Actually, this relation should hold only for medium temperatures, about 0,6 T_m , but one may assume that the errors should not be important if extrapolated to the standard temperature. Thus the experimental data were recalculated for 298 K, inasmuch the temperature of their determination was found in literature, by an extension of equation (20):

$$\gamma_{s}(298) \cong \gamma_{s}(T) \frac{1 - 0.229 \left(\frac{298}{T_{m}} - 1\right) + 0.01 \left(\frac{298}{T_{m}} - 1\right)^{2}}{1 - 0.229 \left(\frac{T}{T_{m}} - 1\right) + 0.01 \left(\frac{T}{T_{m}} - 1\right)^{2}}$$
(21)

These values were also enlisted in Table 2; of course, they are somewhat higher than the experimental values, determined at higher temperatures.

The values of free surface energies, recalculated for the standard temperature, were used to calculate the "molar" quantities $\gamma_{298} V^{2/3}$. These were plotted against free enthalpies of atomization (Fig. 9) and the correlation line has the equation:



$$10^{4}\gamma_{298}V^{2/3}$$
 [mJ] = (20.66±0.65) Δ G_a[kJ/mol] r = 0.916, n = 33 (22)

"Molar" free surface enthalpy (standard temperature) against free enthalpy of atomization
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Now the correlation is somewhat improved, but not in such a degree to justify all the amount of calculations necessary. The points for Be and Re remain here also far from the field of the other values, and without them the equation becomes:

$$10^{4}\gamma_{298}V^{2/3}$$
 [mJ] = (21.28±0.61) Δ G_a[kJ/mol] *r* = 0.936, *n* = 31 (22a)

and after exclusion of V and Ru also:

$$10^{4}\gamma_{298}V^{2/3}$$
 [mJ] = (20.92±0.63) Δ G_a[kJ/mol] r = 0.940, n = 29 (22b)

Among the other elements, higher deviations are observed for Si (nonmetal), Zn and Cd (their specific structure was discussed above) and W, whose very high free enthalpy of atomization should demand a higher value for the free surface energy also.

A possible improvement according to Mezey and Giber [9], is to use so-called "internal" free enthalpies of atomisation $\Delta G_a^{\ i}$. These values are enthalpies of atomization corrected for the expansion work of the gas resulted by splitting all the bonds in the metal, from the (very small) volume of the metallic crystal to the volume of the monoatomic ideal gas. This work is missing in the case of the partial splitting of the bonds, associated with the conversion of a bulk atom into a surface atom. Since this work of expansion is negative, internal free enthalpies of atomisation have larger numerical values than ΔG_a :

$$\Delta G_a^i = \Delta G_a + RT \ln \frac{V_G}{V_S} = \Delta G_a + RT \ln \frac{RT}{p_0 V_S}.$$
(23)

The molar volume of the gas was calculated from the law of ideal gases for the standard state, and the molar volume of the solid metal from the atomic mass and density, in the same state. These ΔG_a^i values are given in Table 2. The difference to the uncorrected values of the enthalpy of atomization is not important (some 10%) and it does not affect substantially the sequence of these values.

The plot of "molar" free surface enthalpies at 298K against the corrected ΔG_a^{i} -values is almost the same as in Fig. 9, and the correlation coefficient is also nearly unchanged.

$$10^{4}\gamma_{298}V^{2/3}$$
 [mJ] = (19.90±0.60) ΔG_{a}^{\dagger} [kJ/mol] $r = 0.917, n = 33$ (24)

The corresponding equations without Be and Re:

$$10^{4}\gamma_{298}V^{2/3}$$
 [mJ] = (20.48±0.55) ΔG_{a}^{i} [kJ/mol] r = 0.937, n = 31 (24a)
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and when V and Ru are also excluded:

$$10^{4}\gamma_{298}V^{2/3}$$
 [mJ] = (20.13±0.57) ΔG_{a}^{i} [kJ/mol] $r = 0.941, n = 29$ (24b)

remain also quite unchanged.

It is interesting, that for both, the highest and the lowest values of free enthalpy of atomization, there are important deviations from the line of correlation. A sigmoidal curve seems to fit better the experimental points than a straight line (Fig. 10). Only the points for Be, Si and Re are now far outside



Sigmoidal representation of $\gamma_{298}V^{2/3}$ -values versus internal enthalpy of atomization

the curve. It may be assumed that for high values of the surface energy, a more important surface relaxation and reconstruction process takes place, in order to reduce tensions, so that the surface energy represents a smaller fraction of the bulk energy than the average for all metals. For low energies, the rearrangement of the atoms on the surface occurs to a lower degree and surface energy is a higher fraction of bulk energy.

Estimation of Surface Free Energies for Solid Elements

In order to compare for different metals the ratio of "molar" surface free enthalpy to the enthalpy of atomization, its value: $k = \frac{\gamma_{298}V^{2/3}}{\Delta G_a^i}$ was calculated (Table 2). The values are quite different, between 10.3 for Be and 43.6 for Cd.

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ESTIMATIONS OF SURFACE FREE ENERGIES FOR SOLID METALS

The mean value is 21.57, the standard deviation 6.15, and the standard error of the mean 1.07. By excluding the extreme values for Be, Si (semi-metals), Re (uncertain value), Zn, Cd (exceptional cases), the span of the remaining 28 values is much narrower (from 15.6 to 25.3); the mean value is 21.08, the standard deviation 2.98 and the standard error of the mean: 0.56.

The "molar" surface quantities used above were conventional ones. In order to estimate the actual molar surface free energies, let us consider the atomic volume in the solid, Ω as the molar volume of the solid divided to Avogadro's number, V_S/N_A. The area assigned to an atom in the surface layer is:

$$s = f\Omega^{2/3} = f \frac{V_s^{2/3}}{N_A^{2/3}}$$
(25)

and the area occupied by a mole of atoms of this kind is $A_M = N_A s$. The molar surface free energy is therefore given by:

$$\gamma_{\rm M} = f N_{\rm A}^{1/3} \gamma V^{2/3}$$
 (26)

f is a packing coefficient, a little greater then 1, depending on the crystalline structure. Ignoring this factor, the $\gamma V^{2/3}$ -values should be multiplied by $N_A^{1/3} = 8,445.10^7$ to give molar surface free energies. The ratio of these values to the enthalpies of atomization, α , should represent, according to the model of broken bonds, the fraction of broken bonds with the nearest neighbours, when an bulk atom goes to the surface of the crystal. Actually, for unknown f-values, the ratio of the two energies is equal to α/f . These values are also given in Table 2.

The statistical analysis of the results is similar to that of the *k*-values. The extreme values are 0.087 (Be) and 0.368 (Cd). For the 33 elements, the mean value is 0.182, the standard deviation 0.052, and the standard error of the mean 0.09. Without the values for Be, Si, Re, Zn and Cd, the range of the remaining 28 values is restrained to 0.128... 0.214; the mean value is 0.177, the standard deviation 0.026 and the standard error of the mean 0.005.

The α -value is therefore about 1/5. If we assume, with Mezey and Giber [16], that the effect of the relaxation and reconstruction in the surface layer were equivalent to the recovery of one of the broken bonds, the transition of an atom from the bulk to the surface should thus bring about the splitting of about 1/4 of its bonds to the nearest neighbours.

In the case of melted metals, for 64 elements, the average ratio found between molar surface energy and enthalpy of vaporization was 0.18 [8], i.e. the same as here for α /f. This speaks for the model of the broken bonds, and shows that the situation is similar in the case of metal melts and of polycrystalline solid metals.

Among the metals with different crystal structures, no significant differences were found: the mean α /f-value is 0.182 for the 7 metals with

body centred cubic lattice and 0.181 for the 9 metals with cubic close packed lattice. There are probably too few available data. According to the theoretical estimations of Mezey and Giber [9] the ratio should be in these two cases 0.184 and 0.192 respectively.

The same ratios calculated from the data of Tyson and Miller [6], the range of values is narrower: the k-values go from 11.40 (Si) to 40.86 (Cd). Other values far from the average are those for Ge (12.06), Be (12.20) and Zn (34.72). For the other elements k varies between 15.00 and 30.72 (α /f between (0.127 and 0.260). The mean value is somewhat lower: k = 19.52 and α /f = 0.165 for 46 elements. Without the extreme values for Si, Ge, Be, Zn and Cd, the mean values become 19.186 and respectively 0.162. These lower ratios can be explained by the fact that the values of the surface free energies refer to the melting point of the solid elements, while the internal free enthalpies of atomization were calculated at 298 K. According to Mezey [9], this ratio should not vary with the temperature, but both quantities should refer to the same temperature.

On the basis of the relations found here, free surface energies for 69 elements were estimated, in three variants:

- from $\gamma_M = k\Delta G_a^{I}$, with the mean value k = 21
- from $\gamma_M = k\Delta G_a^i$, taking for k the slope of the straight line in eq. (20b): k = 20,1
- from the γ_M-values read on the sigmoidal representation (Fig. 14). These free surface energies are given as γ₁, γ₂ and γ₃ in Table 3. By comparison with experimental data (if available) and with calculated

values from different authors (cf. Table 1), the most plausible values were selected (as recommended values printed in bold characters in Table 3). Insofar as possible, for elements of the same group in the periodic table the same kind of values was selected.

For non-metals and semi-metals, as Si, Ge, Be, Sb, Te none of the calculated values are suitable. The sigmoidal correlation gives better values for elements of the main groups: group 2 (Mg - Ra), groups 13 (Al - Tl), 14 (Sn - Pb), but also for groups 12 (Zn,Cd) and 15 (V - Ta), or for transition metals with highest values of surface tensions: W, Re. This correlation works also for the bivalent lanthanoides Yb and Eu, whose similarities to elements of the groups 2 and 12 are known, and also for Th and U.

For alkali metals (1st main group) the linear correlation with the lower k-value (γ_2) seems to fit the best; as well as for lanthanoides (except Eu and Yb), elements of group 15 (Sb, Bi) and for Ni and Mo.

The linear correlation with the higher k-value (γ_1) seems to work best for transition metals of the groups 3 (Sc, Y), 4 (Ti - Hf), Mn, groups 8-10 (except Ni), 11 (Cu - Au).

ESTIMATIONS OF SURFACE FREE ENERGIES FOR SOLID METALS

For some elements (Po, At, Ac) there is no possibility to select the best values, since both experimental and calculated values are missing.

Table 3.

Free surface energies (mJ/m^2) of solid metals at standard temperature, estimated from linear correlations for k=21 (γ_1) and for k = 20,1 (γ_2) and from sigmoidal correlation (γ_3). Recommended are the bold values

Element	Li	Be	Na	Mg	Al	Si	К	Са	Sc
γ1	550	2230	240	470	1390	1730	130	380	1230
γ2	530	2140	230	450	1330	1660	120	370	1180
γз	730	2030	470	690	1280	1780	300	470	1162
Element	Ti	V	Cr	Mn	Fe	Со	Ni	Cu	Zn
γ1	1960	2440	2110	1450	2250	2410	2450	1810	540
γ2	1880	2330	2020	1380	2150	2310	2340	1730	520
γ3	2050	2610	2020	1320	2220	2390	2450	1640	900
Element	Ga	Ge	Rb	Sr	Y	Zr	Nb	Мо	Tc
γ1	1050	1300	100	290	1150	2120	2980	2990	3310
γ2	1010	1250	95	280	1100	2030	2850	2860	3170
γ3	960	1220	260	390	1150	2180	2710	2950	3180
Element	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Те
γ1	3200	2740	1750	1180	360	750	930	730	500
γ2	3060	2620	1680	1130	340	720	890	700	480
γ3	3200	2910	1650	1080	700	730	830	690	570
Element	Cs	Ba	La	Ce	Pr	Nd	Sm	Eu	Gd
γ1	80	300	1140	1140	950	880	550	370	1090
γ2	75	283	1090	1090	910	840	530	350	1050
γ3	220	360	1140	1140	880	790	590	440	1060
Element	Tb	Dy	Ho	Er	Tm	Yb	Lu	Hf	Та
γ1	1100	820	860	910	670	330	1260	2390	3250
γ2	1050	780	820	870	640	320	1200	2290	3110
γ ₃	1060	740	770	830	640	470	1260	2330	2750
Element	W	Re	Os	lr	Pt	Au	ΤI	Pb	Bi
γ1	3870	3660	3870	3220	2620	1560	510	540	510
γ2	3700	3510	3700	3080	2510	1490	490	520	490
γ3	3040	3160	3260	3140	2760	1440	610	610	560
Element	Po	At	Ra	Ac	Th	U			
γ1	330	220	240	1010	1650	2010			
γ ₂	320	210	230	970	1580	1920			
γ ₃	500	530	320	990	1710	2150			
									1

Conclusions

The scarcity of direct experimental data for the surface tensions or surface free energies of solid metals make necessary their semi-empirical estimation or the theoretical calculation. Using selected experimental values, their correlation with melting points and standard enthalpies of atomization was verified. "Molar" surface free energies, calculated by means of molar volumes or metallic radii gave acceptable linear correlations with the enthalpies of atomization. With standard free enthalpies of atomization, calculated with the entropies of atomization, obtained as differences between the entropies of gaseous and solid metals, the correlations are almost the same as with standard enthalpies.

In order to improve the calculations, surface free energies at standard temperature were estimated from the experimental values and internal free enthalpies of atomization were calculated, but not significantly better correlations were obtained. The deviations of some elements from the correlation lines can be explained by errors in the experimental data or by peculiarities in their structure.

The results can be interpreted in terms of the theory of broken bonds between neighbouring atoms in the metallic crystals. The correlations found here could be used to estimate values of free surface energies for 69 solid elements.

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ADSORPTION KINETICS OF STEARIC ACID AT THE BENZENE/WATER INTERFACE

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ABSTRACT. The adsorption kinetics of stearic acid from benzene solutions at the benzene/water interface was studied by using the pendant drop method. Kinetic equations for diffusion controlled adsorption and for Langmuir adsorption were tested. A theoretical approach to the dynamic interfacial pressures is developed and a new kinetic equation for the diffusion controlled kinetics is proposed, based on Ward and Tordai's diffusion equation associated with a two dimensional van der Waals state equation. It is shown that this approach describes the dynamic interfacial pressures within the limits of the experimental errors and the new equation is valid over a wide range of time and for different surfactant concentrations in oil solutions. It is also concluded that this new approach allows the calculation of diffusion coefficients, subsurface concentrations and intermolecular interaction parameters within adsorbed layers of biosurfactants at the oil/water interfaces. Obtained results are in substantial agreement with similar earlier reported data for the surfactant adsorption at liquid/liquid interfaces.

Keywords: Diffusion controlled adsorption; Langmuir adsorption kinetics; Dynamic interfacial pressures; Pendant drop method; Stearic acid; Benzene/water interface

Introduction

The adsorption of surfactant molecules at the air/water and oil/water interfaces has attracted a considerable attention in the last several decades (1-28) due to its fundamental and industrial importance (1-4, 25) as well as to its biological significance (29-32). These studies have involved various experimental techniques as well as the formulation of theoretical models for describing the adsorption kinetics of surfactants at liquid interfaces.

Generally, the adsorption of surfactant molecules from a solution to a liquid/gas (1-15) or to a liquid/liquid interface (1, 5, 16-24) occurs in two steps. Surfactant is first transported from the bulk to the subsurface by diffusion (the subsurface is a liquid layer just below the interface, belonging still to the bulk). The second step is the transfer from the subsurface to the interface, implying sometimes a transfer through a potential barrier (22, 25-28). In miscellaneous systems the controlling rate may be either the diffusion (i.e. the adsorption may be diffusion controlled), or the transfer

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(i.e. the adsorption is barrier-controlled). Several theoretical models have been developed for both processes, as well as for mixed adsorption mechanisms in which both steps are considered (1-6).

In the case of adsorption to a liquid/liquid interface (1, 16-24), the situation is analogous to that for the air/water interface if the solute is soluble only in one liquid phase. If the solute is soluble in water and in oil phases, then the adsorption process includes also the solute transfer across the interface into the other liquid bulk phase (16).

For example, the adsorption kinetics of some aliphatic carboxylic acids and aliphatic alcohols (6-9), like 1, 9 nonane dicarboxylic acid and 1, 9 nonane diol (6), was studied at the air/water interfaces and described by Langmuir kinetic equation, considering the adsorption and desorption rate constants. Another example is related to the adsorption of some salts of fatty acids, like sodium laurate (10-13), sodium myristate (10-12), and sodium oleate (14), from aqueous solutions to the air/water interface, which was analyzed by means of diffusion controlled kinetics. However, the very slow adsorption of fatty acids from oil phase to the oil/water interface has received only a limited attention (5).

The main goal of this work is to focus on the experimental dynamic data for the adsorption of stearic acid at the benzene/water interface and to discuss the theoretical aspects of adsorption kinetics at the oil/water interface. The experimental data are recorded in terms of the time dependent interfacial pressures, which are also known as dynamic interfacial pressures. These interfacial pressures are dependent of time and of the surfactant bulk concentration and of the surfactant interfacial adsorption. An equilibrium state is reached when the interfacial pressure is constant (18, 23, 24).

Stearic acid is a fatty acid widely used in model biomembranes (29-32) as well as in Langmuir-Blodgett films due to its increased stability in monolayers at fluid interfaces (33-35). The understanding of the adsorption kinetics of stearic acid at fluid interfaces is important for the description of its dynamic surface properties. This is not only an interesting subject in basic research, but also very important for fabricating stable, homogeneous and ordered molecular films.

Further, we examine the diffusion approach, its simplifying assumptions, and Langmuir adsorption kinetics for the adsorption of stearic acid at the oil/water interface. As a first step, we apply the theory of diffusion controlled kinetics on ideal interfacial layers and the Langmuir adsorption kinetics to experimental data of the adsorption of stearic acid at the oil/water interface.

Furthermore, we extend the diffusion theory for non-ideal adsorbed layers at liquid/liquid interfaces (24) and we develop a theoretical approach based on the Ward and Tordai's diffusion equation associated with a two

dimensional state equation of van der Waals type and a new diffusion equation is derived and used to describe the dynamic interfacial pressures. This approach is developed and an analytical procedure of calculation is proposed to solve the resulting new diffusion controlled equations for the surfactant adsorption at liquid/liquid interfaces.

Finally, a mathematical procedure is formulated to determine the diffusion coefficient, the subsurface concentration and the molecular interaction parameters within the adsorbed layers of surfactant at liquid/liquid interface.

Theoretical Models of Adsorption Kinetics *Diffusion Controlled Kinetic Model*

Diffusion controlled adsorption was analyzed quantitatively by Ward and Tordai (5, 24) using the following equation:

$$\Gamma(t) = 2 \left(\frac{D}{\pi}\right)^{1/2} \left\{ c_0 t^{1/2} - \int_0^{t^{1/2}} c_s(t-\tau) d\tau^{1/2} \right\}$$
[1]

where $\Gamma(t)$ is the dynamic adsorption of the surfactant from an infinite bulk phase at the interface up to the time t, *i.e.*, $\Gamma(t)$ is the amount of the solute which has diffused across the subsurface into the interface, up to the time t; D stands for the bulk diffusion coefficient of the surfactant (defined by Fick's equation); π is 3.1415; c₀ is the surfactant bulk concentration far from the surface; c_s(t) is the subsurface concentration and τ stands for a dummy variable ranging from 0 to t. At zero time, c_s(t) = 0. Later, it begins to increase and c_s(t) becomes equal to c₀, when the adsorption equilibrium is attained.

In Eq. [1] the second term on the right side contains the back diffusion integral taking into account the desorption process. Practically, at the beginning this integral vanishes and the second term may be neglected, obtaining the short time approximation:

$$\Gamma(t) = 2 \left(\frac{\mathbf{D}}{\pi}\right)^{1/2} \cdot \mathbf{c}_0 \mathbf{t}^{1/2}$$
 [2]

Another approximation has been derived for long time by presuming that at long time $c_s(t)$ is almost constant. In this approach one obtains:

$$\Gamma(t) = 2(c_0 - c_s(t)) \left(\frac{D}{\pi}\right)^{1/2} t^{1/2}$$
[3]

Both Eqs. [2] and [3] suggest a proportionality of the adsorption with $t^{1/2}$. Generally, the dynamic interfacial tension, $\sigma(t)$, or the dynamic interfacial pressure, $\Pi(t)$, is measured and used as a tool to investigate the surfactant adsorption at the fluid interfaces.

In order to establish a correlation between the adsorption $\Gamma(t)$ and the dynamic interfacial tension $\sigma(t)$ or the dynamic interfacial pressure $\Pi(t) = \sigma_o - \sigma(t)$, frequently, the surface state equation:

$$\Pi(t) A(t) = kT \text{ or } \Pi(t) = \Gamma(t)kT$$
[4]

is used, which is the two dimensional analog of the perfect gas state equation (22, 27); σ_0 represents the interfacial tension of the pure interface in the absence of the surfactant. It is argued that the state equations apply whether the monolayer is in equilibrium with the subsurface or whether it is not.

Combining Eq. [4] with Eqs. [2] and [3], respectively, one obtains:

$$\Pi(t) = 2kT \left(\frac{D}{\pi}\right)^{1/2} c_0 t^{1/2}$$
 [5]

$$\Pi(t) = 2kT \left(\frac{D}{\pi}\right)^{1/2} (c_0 - c_s(t))t^{1/2}$$
 [6]

Both Eqs. [5] and [6] show a linear relationship between the dynamic interfacial pressure, $\Pi(t)$, and $t^{1/2}$. Details on this approach will be presented further on in this investigation.

Langmuir Kinetic Model

On the other hand, by presuming that the molecular diffusion is very fast and the transfer from the subsurface to the surface is the rate limiting step and it follows a *Langmuir kinetics* (6-9), one obtains:

$$y = \ln \frac{\Delta \sigma_0}{\Delta \sigma} = \ln \frac{\sigma_0 - \sigma_e}{\sigma(t) - \sigma_e} = K t$$
[7]

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where $\sigma(t)$ stands for the actual and σ_e for the equilibrium interfacial tension and σ_0 for the interfacial tension in the absence of the surfactant. Further, the rate constant K is given by:

$$K = \frac{k_1}{\Gamma_{\infty}} c_0 + \frac{k_2}{\Gamma_{\infty}}$$
[8]

where k_1 is the rate constant of adsorption (cm s⁻¹), k_2 stands for the rate constant of desorption (mole cm⁻² s⁻¹) and Γ_{∞} is the maximum adsorption (mole cm⁻²) at the saturation of the liquid/liquid interface with surfactant molecules.

In the present paper the dynamic interfacial pressures, $\Pi(t) = \sigma_o - \sigma(t)$, at the benzene/water interface have been measured for benzene solutions of stearic acid (octadecanoic acid: SA) of various SA concentrations. The validity of the above equations has been tested and an attempt is made to derive a new equation for the diffusion controlled adsorption, taking into account the non-ideal behavior of adsorbed films at liquid interfaces, particularly for high bulk surfactant concentrations. The values of the experimental data performed on the basis of the new equation shows that the latter allows a better description of the adsorption kinetics, than the other equations given above.

Experimental

Stearic acid (octadecanoic acid: SA) was a synthetic commercial product of high purity (minimum 99%) purchased from Sigma. Benzene pro-analysis was purchased from Merck. All chemicals were used without an additional purification. Twice-distilled water of pH 2 was used, containing 0.01 mole/dm³ of hydrochloric acid, as the aqueous phase. Volumetric aqueous solutions of HCl pro-analysis were purchased from Reactivul Bucharest.

Stearic acid at pH 2 forms uncharged adsorbed monolayer, the molecules being completely unionized (29). Due to the very low solubility of stearic acid in water of pH 2, any transport across the benzene/water interface can be neglected. The pH of the aqueous solutions was constant during all experiments and it was measured by an MV-84 type pH-meter by using a glass electrode.

Dynamic surface tensions in the time range from 1 minute up to 120 minutes were measured by pendant drop and by ring method for pure benzene and benzene solutions of various stearic acid concentrations at the interface with aqueous solutions of pH 2.

The pendant drop technique was described by us elsewhere (23). The shape of drops was recorded on a highly quality film in order to determine the characteristic drop diameters. By using an appropriate computer program the dynamic interfacial tension values were finally determined.

Experimental data obtained by the pendant drop technique were compared with the data obtained by ring method described by us previously (36). The agreement between the two methods is excellent and the deviations do not exceed the error of the individual method. The accuracy of interfacial tension measurements was \pm 0.1 mN/m, in agreement with literature data (20, 37). All measurements were performed at constant temperature of 20 \pm 0.1 °C.

Results and Discussions

Dynamic interfacial tension $\sigma(t)$ values obtained by means of the pendant drop method, allowed us to calculate the corresponding $\Pi(t) = \sigma_0 - \sigma(t)$ interfacial pressure values. The $\Pi(t)$ *versus* time (t) curves, characterizing the adsorption process of SA from benzene at the benzene/water (pH 2) interface is presented in Fig.1.





Experimental dynamic interfacial pressures, $\Pi(t) = \sigma_o - \sigma(t)$ in mN/m, of stearic acid (SA) benzene solutions as function of time, t in min, at the benzene/water of pH 2 interface. Figures indicate the bulk concentration C₀ of stearic acid in benzene, in mole dm⁻³. Solid lines calculated by Eq. [18].

As can be seen, the interfacial pressures vary with surfactant bulk concentrations and with time over a wide range of time, from 0 to 90 min. Generally, the equilibrium interfacial pressures (Π_e in mN/m) are recorded at 120 min when the adsorption equilibrium is completely attained and it is demonstrated by the constant value of the interfacial pressures.

The validity of Eq. [7] has been tested by calculating the left hand side (denoted y) of this equation, by using the $\sigma(t)$ and σ_e values (the latter ones being measured at 120 min) and the interfacial tension of the pure benzene/water interface in the absence of surfactant was taken $\sigma_o = 34.7$ mN/m.

Inspecting the experimental data, it is found that y *versus* t data exhibit a quite good linearity, although the straight lines do not pass through the origin of the coordinate system, *i.e.* actually instead of Eq. [7] one has:

$$y = \ln \frac{\sigma_0 - \sigma_e}{\sigma - \sigma_e} = B + K t$$
 [7']

By performing a linear regression, the parameters B and K have been determined. Results are presented, together with the correlation coefficients, r, in Table I. In the last column n indicates the number of experimental points used in the linear regression.

Table I.

The B and K parameters of Eq. [7'].

C ₀ mole dm ⁻³	В	K min ⁻¹	r	n
0.001	0.430	0.0397	0.9991	7
0.00178	0.374	0.0396	0.9977	7
0.0057	0.395	0.0406	0.9990	8
0.012	0.439	0.0415	0.9978	8
0.018	0.568	0.0431	0.9994	9
0.051	0.566	0.0501	0.9949	10

As seen from Table I, the B values are rather far from zero. This means that the basic hypothesis used to derive Eq. [7] is not perfectly valid, presumably the diffusion equilibrium is not yet established, and consequently the boundary condition $c_s(t) = c_0$ is not fulfilled. Nevertheless, from the K values reported in Table I, some conclusions can be drawn. As seen, with increasing c_0 the K values derived also increase, as expected on the basis of Eq. [8] and the dependence of K with c_0 exhibits indeed a quite good linearity. The relative adsorption k_1/Γ_{∞} and desorption k_2/Γ_{∞} rate constants derived by means of linear regression Eq. (8) and the corresponding correlation coefficients are presented in Table II.

Table II.

Relative adsorption, k_1/Γ_{∞} , and desorption, k_2/Γ_{∞} , rate constants derived by Eq. [8] from K values listed in Table I.

Surfactant	k_1/Γ_{∞} mole ⁻¹ dm ³ min ⁻¹	k_2/Γ_∞ min ⁻¹	r	k ₁ /k ₂ mole ⁻¹ dm ³
Stearic acid	0.212	0.0392	0.9990	5.4
Dibucaine	7.25	0.159	0.9914	45.6
Tetracaine	4.99	0.149	0.9988	33.5

Table II contains also the same magnitudes previously obtained for two anesthetics, *viz.* dibucaine and tetracaine, under similar working conditions (24).

The analysis of data in Table II shows the quite good validity of Eq. [8], reflected by the correlation coefficient values. The relative rate constant values are rather reasonable and their k_1/k_2 ratios are in substantial agreement with published data (6-8) for the adsorption of various surfactants at liquid/liquid interfaces. These ratios represent the equilibrium adsorption constants and characterize the interfacial activity of these biocompounds at the oil/water interface. In this respect, the k_1/k_2 value for dibucaine is higher than the corresponding value for tetracaine suggesting a higher interfacial activity of the dibucaine in agreement with the interfacial pressure measurements at the oil/water interface.

Generally, the relative rate constant values (Table 2) for anesthetics are much higher than those corresponding to stearic acid. These values are not easy to compare because the mechanism of adsorption and desorption of these two types of biocompounds (i.e. anesthetics and fatty acids) is completely different and consequently the rate constants are determined by different factors. In spite of the different mechanism for adsorption, both fatty acid and anesthetics strongly adsorb at the oil/water interface and show a certain type of similarity in their adsorption behavior.

In the case of stearic acid, molecules adsorb from oil phase to the oil/water interface immersing their unionized polar groups into the water phase. The driving force for the adsorption of stearic acid molecules is the change in enthalpy due to the hydration of their adsorbed hydrophilic groups, when they immerse into the water phase being accompanied by conformational rearrangements in the hydrocarbon chains. Unlike stearic acid, anesthetics are ionized water soluble compounds. They adsorb from 166

water to the oil/water interface by penetrating their hydrophobic chains into the oil phase. Their adsorption is controlled by the hydrophobic effect, by the electrostatic interactions among the ionized polar groups, and by the hydrophobic interactions among the hydrophobic chains and the oil phase. According to the hydrophobic effect, the hydrocarbon chains in water phase are surrounded with structured water. Therefore, the adsorption of anesthetics is also accompanied by the increase of the entropy of the system because of the destruction of the ordered structure of water molecules formed around the hydrophobic chains in aqueous phase.

Although these results seem to be reasonable, the quite high B values reported in Table I show that Eq. [7], successfully used to describe the adsorption at the air/water interface (6-8), in our case is an approximation and the diffusion might play an important part in the overall adsorption process. Consequently, we further test the validity of Eqs. [5] and [6].

For this purpose, the experimental $\Pi(t)$ values calculated from σ_0 and $\sigma(t)$ values are presented *versus* t^{1/2} in Fig. 2.



Fig. 2.

Experimental dynamic interfacial pressures of stearic acid benzene solutions at the interface with water of pH 2 as function of $t^{1/2}$. (Symbols as in Fig. 1).

If the Eq. [5] or Eq. [6] was valid, the $\Pi(t)$ versus $t^{1/2}$ curves would exhibit a linear portion at least at small t values. This is why the tangencies to the experimental curves for t = 0 were constructed (Fig. 2). Although, the experimental curves exhibit an important negative deviation from these

straight lines, certain linearity may be observed. In order to obtain a clear image, a linear regression analysis has been performed by adding successively the experimental data, one by one, and by calculating every time the parameters of the following equation:

$$\Pi(t) = mt^{1/2} + p$$
 [9]

The general picture is the following, by adding the experimental data in the order of increasing t values, the slope of the straight line (*i.e.,* the m value) diminishes, and the ordinate intercept p increases. In the same time the correlation coefficient diminishes.

Results are presented in Table III, where the limits of m and p values are given. Those obtained by means of the first three experimental points (noted m_i and p_i , respectively), as well as the values obtained with a final number, n_f , of experimental points (noted m_f and p_f , respectively), for which one had still r > 0.99, r being the correlation coefficient. In Table III are also listed the total number, n_{max} , of the experimental points obtained for every bulk concentration of SA.

Being more specific, Eq. [9] is a general form of Eqs [5] and [6].

C ₀ mole dm ⁻³	m _i mN m ⁻¹ min ^{-1/2}	m _f mN m ⁻¹ min ^{-1/2}	p _i mN m⁻¹	p _f mN m⁻¹	n _f	n _{max}
0.001	0.47	0.33	1.16	1.59	10	10
0.00178	0.63	0.48	1.59	1.72	7	9
0.0057	1.47	1.26	2.23	2.85	6	9
0.012	2.27	2.08	1.96	2.48	4	11
0.018	2.14	1.88	4.64	5.92	5	11
0.051	2.68	2.50	6.57	7.06	4	11

Limits of the m (m_i and m_f) and p (p_i and p_f) parameters of Eq. [9] for r > 0.99. For symbols see the text.

Table III.

The analysis, of the data given in Table III, shows that Eqs. [5] and [6] are very rough approximations, because p from Eq. (9) is very far from the expected zero value, especially in the case of higher SA concentrations. Further, the decrease of the m values with increasing time $(m_i > m_f)$ indicates negative deviations from the linearity predicted by Eq. [5] or Eq.

[6], both being consistent with the case of diffusion adsorption for the ideal behavior, given by Eq. (4), of the interfacial film of adsorbed surfactant.

As a first approximation let us presume that at low t values Eq. [5] is valid and consequently in Eq. [9] one has p = 0 and m_i, the coefficient of $t^{1/2}$, is given by the following equation:

$$m_{i} = 2kT \left(\frac{D}{\pi}\right)^{1/2} c_{0}$$

By expressing D, one obtains:

$$D = \frac{\pi m_i^2}{(2kTc_0)^2}$$
[10]

We mention that in Eq. [1] $c_{0}\xspace$ means the bulk concentration expressed in surfactant molecules/volume. Consequently, if one gives mi in dyne cm⁻¹ s^{-1/2} (see Table IV), k in erg/(molecule grade) and c_0 in molecules cm^{-3} , D will be obtained in $cm^2 s^{-1}$. Obviously, if the bulk concentration is given as C_0 mole/L, one has $c_0 = 10^{-3} N_A C_0$, where N_A stands for Avogadro's constant. Diffusion coefficients, D values, calculated by means of Eq. [10] from m_i values, listed in Table III, are given in Table IV.

Table IV.

١	values calculated	by Eq.[12] from th	e slopes m _i take	en from Table III.
C ₀ mole dm ⁻³	m _i dyne cm ⁻¹ s ^{-1/2}	D 10 ¹¹ cm ² s ⁻¹	D ₀ cm ² s ⁻¹	C _s 10 ³ mole dm ⁻³
0.001	0.0607	0.492		0
0.00178	0.0813	0.278		0.44
0.0057	0.1900	0.148	4.92 10 ⁻¹²	2.57
0.012	0.2930	0.0796		7.17
0.018	0.2760	0.0314		13.4
0.051	0.3460	0.00614		45.3

Diffusion coefficients. D and D₀ calculated by Eq. [10], and C₀

As seen, in Table IV, D diminishes with increasing bulk concentration, suggesting the idea that Eq. [6] might be a better approach than Eq. [5]. One

may take, then, from Eq. [6] for m_i the following expression:

$$m_{i} = 2kT \left(\frac{D}{\pi}\right)^{1/2} (c_{0} - c_{s})$$
 [11]

It is reasonable to presume that the subsurface concentration C_s (expressed in mole/L) increases with increasing bulk concentration C_0 . Let us presume as a first approach that $C_s = 0$ for $C_0 = 0.001$ mole/L. In this case a D_0 value can be obtained by means of Eq. [10]. Further, by using this D_0 value and Eq. [11], C_s values may be derived, from the m_i slopes obtained for higher concentrations, *viz.*, by means of the following relation:

$$C_{s} = C_{0} - \frac{\pi^{1/2} m_{i}}{2kT \cdot 10^{-3} N_{A} D_{0}^{1/2}}$$
[12]

The values of subsurface concentrations (C_s , calculated by Eq.[12]) and diffusion coefficient D_0 are also given in Table IV.

As seen, C_s increases with increasing C_0 , as one might expect. However, the diffusion coefficient is with several orders of magnitude less than real ones (D = 5 10^{-6} cm² s⁻¹) assumed for surfactants diffusing through aqueous phase to the air/water interface (7). This means, that Eqs. [5] and [6] are extremely rough approximations to fit these experimental data.

The remarkable low values of the diffusion coefficients are probably due to a slow orientation (7, 38) of the polar head groups of SA molecules at the interface and to a slow conformational transition in the long hydrocarbon chains of SA molecules to finally achieve their correct orientation at the adsorption equilibrium. These effects can also explain the slow interfacial pressure increase to reach its equilibrium value. As an alternative, the long time dependence of the interfacial pressure can be attributed to the interactions among SA molecules at the oil/water interface, where these SA molecules relax and reorient leading to their proper molecular arrangement characteristic for the adsorption equilibrium.

Modeling of the New Diffusion Kinetic Equation

Eqs. [5-6] and [9-12] have been obtained by presuming the state equation [4] to be valid. But interfacial monolayers obey Eq. [4] only at very small Π values, practically for $\Pi \rightarrow 0$. One of the best state equations, proposed for non-ionic monolayers (39) and quoted and used by others (40-42), is of the following shape:

$$\left(\Pi + \frac{\alpha}{A^{3/2}}\right) (A - A_0) = kT$$
[13]

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where A stands for the mean molecular area, A_0 for the own molecular area of the surfactant and α is an interaction parameter. Eq. [13] was found to describe very well the compression isotherm of miscellaneous monolayers (40, 41), especially if the interaction parameter α and the own molecular area A_0 are treated as adjustable parameters.

From state equation [13], Π may be expressed as:

$$\Pi = \frac{kT}{A - A_0} - \frac{\alpha}{A^{3/2}} = \frac{kT\Gamma}{1 - \Gamma A_0} - \alpha \Gamma^{3/2}$$
[14]

Let us denote:

$$\mathbf{x} = 2(\mathbf{c}_0 - \mathbf{c}_s) \left(\frac{\mathbf{D}}{\pi}\right)^{1/2}$$
[15]

Thus, the long time approximation of diffusion, given by Eq. [3], may be written as $\Gamma(t) = xt^{1/2}$. Then, combining Eqs. [3], [13] and [15], one obtains:

$$\Pi = \frac{kTxt^{1/2}}{1 - A_0xt^{1/2}} - \alpha x^{3/2}t^{3/4} = kTx\left[\frac{t^{1/2}}{1 - A_0xt^{1/2}} - \frac{\alpha x^{1/2}}{kT}t^{3/4}\right]$$
[16]

Therefore, one may expect Eq. [16] to describe better the time dependence of Π , than Eq. [6] does.

Introduction of Dimensionless Variables

In order to study the properties and possibilities of Eq. [16], let us introduce a reduced time scale. If in an experiment Π has been followed from t = 0 up to a maximum time t_m, a reduced time (noted τ) can be defined as:

$$\tau = \frac{t}{t_{m}}$$
[17]

Combining Eqs. [16] and [17] one obtains:

$$\Pi = kTxt_{m}^{1/2} \left[\frac{\tau^{1/2}}{1 - A_{0}xt_{m}^{1/2}\tau^{1/2}} - \frac{\alpha x^{1/2}}{kT}t_{m}^{1/4}\tau^{3/4} \right]$$

The Eq. [17] may be written as:

$$\Pi = a \left[\frac{\tau^{1/2}}{1 - b\tau^{1/2}} - c\tau^{3/4} \right] = a\phi(\tau)$$

$$a = 2kT(c_0 - c_s) \left(\frac{Dt_m}{\pi} \right)^{1/2}$$

$$b = 2A_0(c_0 - c_s) \left(\frac{Dt_m}{\pi} \right)^{1/2}$$

$$c = \frac{2^{1/2} \alpha (c_0 - c_s)^{1/2}}{kT} \left(\frac{Dt_m}{\pi} \right)^{1/4}$$

From the expression of the parameters *a*, *b* and *c* one obtains:

$$D = \left[\frac{a}{2kT(c_0 - c_s)}\right]^2 \frac{\pi}{t_m}; \quad A_0 = \frac{kTb}{a}; \quad \alpha = \frac{c}{a^{1/2}}(kT)^{3/2}$$
[19]

Obviously, if t = t_m , from Eq. [17] one has τ = 1 and Eq. [18] becomes:

$$\Pi_{\rm m} = a\phi(1) = a \left[\frac{1}{1-b} - c \right]$$
[20]

Let us introduce also for the interfacial pressure a reduced scale:

$$\Pi^* = \frac{\Pi}{\Pi_m}$$

where Π_m stands for the maximum interfacial pressure, corresponding to t = t_m. By using the Eqs. [18] and [20], this reduced interfacial pressure is expressed as:

$$\Pi^* = \frac{\Pi}{\Pi_m} = \frac{\varphi(\tau)}{\varphi(1)}$$
[21]

Thus one obtains finally for the time dependence of the interfacial

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with:

pressure:

$$\Pi = \frac{\prod_{m}}{\phi(1)}\phi(\tau)$$
 [22]

By comparing Eq. [22] with Eq. [18] it is obvious that:

$$a = \frac{\prod_{m}}{\phi(1)}$$
 [23]

Procedure of Calculations

In order to test the utility of Eq. [22] or of Eq. [18], we studied the influence of the parameters *b* and *c* upon the shape of the Π /a versus τ and Π^* versus τ curves. For this purpose theoretic curves have been constructed. Some examples are given in Figs. 3-6.





Influence of the *c* parameter upon the theoretical $\varphi(\tau)$ versus τ curves, given by Eq. [18], for *b* = 0.1. The *c* values: curve (1) 0; (2) 0.2; (3) 0.4; (4) 0.6; (5) 0.8; (6) 1.0. Dashed line curve: *b* = *c* = 0.

In Fig. 3 the influence of the *c* parameter, upon the shape of the $\Pi/a = \varphi(\tau)$ versus the dimensionless τ curves, is illustrated by taking the constant *b* = 0.1 value. Fig. 4 illustrates the influence of the *b* parameter for the constant *c* = 0.6 value.

Since parameters *b* and *c* influence also the $\varphi(1)$ value (see Eq. [20]), the shift of the calculated curves makes their comparison rather difficult. The influence of these parameters is more clear if the $\Pi^* = \varphi(\tau)/\varphi(1)$ versus τ 173

curves are constructed. These are shown in Figs. 5 and 6.



Fig. 4.

Influence of the *b* parameter upon the theoretical $\varphi(\tau)$ versus τ curves, given by Eq. [18], for c = 0.6. The *b* values: curve (1) 0; (2) 0.05; (3) 0.1; (4) 0.15; (5) 0.2; (6) 0.3; (7) 0.4. Dashed line curve: b = c = 0.



Influence of the *c* parameter upon the theoretical $\prod^* versus \tau$ curves, given by Eq. [21]. The *b* and c values as in Fig. 3.



Influence of the *b* parameter upon the theoretical \prod^{\star} versus τ curves, given by Eq. [21]. The *b* and *c* values as in Fig. 4.

It is worth mentioning that for b = 0.1 and c = 1.0 both $\varphi(\tau)$ (curve 6 in Fig. 3) and Π^{*} (curve 6 in Fig. 5) exhibit a maximum for about $\tau = 0.3$. This would imply the appearance of a maximum interfacial pressure at a certain time, followed by the diminution of Π . This is of course unrealistic. Therefore, it is obvious that not all *b* and *c* pairs can describe the real adsorption kinetics. Further, the curve for b = 0.4, c = 0.6 (curve 7 in Fig. 4) indicates the adsorption rate to decrease down to a minimum value, followed by the acceleration of the adsorption. This also cannot happen in a real adsorption process, since the more occupied the adsorption sites, the less will be the adsorption rate, becoming zero at the adsorption equilibrium.

Although, not all calculated $\varphi(\tau)$ functions correspond to real adsorption behavior of surfactants, the main feature is that in the case of *diffusion controlled adsorption* the own molecular area A₀ and the intermolecular interactions expressed by the parameter α may count for both positive and negative deviations from the t^{1/2} law (see Fig. 2).

Since according to the $t^{1/2}$ law expressed by Eq. [6] one has $\Pi^* = \tau^{1/2}$, the deviation of the reduced surface pressure from the $t^{1/2}$ law may be expressed by the function:

$$\Delta = \frac{\varphi(\tau)}{\varphi(1)} - \tau^{1/2}$$
 [24]

The influence of the *c* and *b* parameters upon the theoretical Δ vs. τ curves is illustrated in Figs. 7 and 8, respectively. It is obvious that both positive and negative Δ values may appear and the deviation from the t^{1/2} law may be very important.





Influence of the *c* parameter upon the theoretical Δ versus τ curves, given by Eq. [24]. The *b* and *c* parameters as in Fig. 3.



Fig. 8.

Influence of the *b* parameter upon the theoretical \triangle versus τ curves, given by Eq. [24]. The *b* and *c* parameters as in Fig. 4.

In real systems, as seen in Fig. 2, with increasing time negative deviations from the t^{1/2} law appear. In a Π^* vs. τ plot this means that the curve is more convex (has a higher curvature) as compared to $\tau^{1/2}$, i.e. Δ has positive values. Consequently, negative Δ values indicate unrealistic theoretical curves. Even the positive values have an upper limit, it is easy to see that the Π vs. t curve passes through a maximum if $\Delta > 1 - \tau^{1/2}$. The dashed line curve in Fig. 7 indicates this upper limit $\Delta = 1 - \tau^{1/2}$.

Since the Δ vs. τ curves exhibit important shifts if *b* or *c* is modified, in order to illustrate the influence of the parameters b and c upon the shape of the curves, we choose a set of *b* – *c* pairs which oblige the Δ vs. τ curves to pass through a common point. If we impose the condition that at a certain τ_0 value the Δ function must be equal to Δ_0 , Eq. [24] becomes:

$$\Delta_{o} = \frac{\phi(\tau_{0})}{\phi(1)} - \phi_{0}^{1/2}$$

From this relation one might express the *c* parameter and one obtains:

$$c = \frac{1}{\Delta_0 + \tau_0^{1/2} - \tau_0^{3/4}} \left[\frac{(\tau_0^{1/2} - \tau_0)b}{(1-b)(1-\tau_0^{1/2}b)} + \frac{\Delta}{1-b} \right]$$
[25]

Table V.

The *b* and *c* pairs, calculated by Eq. [26], ensuring $\Delta = 0.2$ for $\tau = 0.2$.

No.	b	С
1	0	0.575
2	0.02	0.601
3	0.05	0.643
4	0.10	0.721
5	0.15	0.810
6	0.20	0.913
7	0.25	1.033
8	0.30	1.172

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Eq. [25] allows us to calculate for a given $\Delta_0 - \tau_0$ pair the *c* parameter as function of *b*. Thus, for $\Delta_0 = \tau_0 = 0.2$ one obtains:

$$c = \frac{0.71b}{(1-b)(1-0.447b)} + \frac{0.574}{1-b}$$
 [26]

Some illustrative examples of c - b pairs ensuring $\Delta = 0.2$ for $\tau = 0.2$, calculated by means of Eq. [26] are given in Table V.

For cases 1, 3, 4, 6, 7 and 8 whole Δ vs. τ curves are given in Fig. 9.



Some \triangle versus τ curves passing through the point $\triangle = \tau = 0.2$. Figures indicate the *b* and *c* pairs given in Tab. 5.

It is clear that Eq. [18] corresponds to a very flexible function able to describe well a great variety of curves. Since for b = 0.3 the deviation Δ arrives in the negative region for $\tau > 0.85$, Eq. [26] may give b - c pairs corresponding to real experimental curves only if b < 0.3.

Deriving of the Parameters of Eq. [18] from Experimental Data

In order to derive the parameters *a*, *b* and *c* of Eq. [18] we adopted the following procedure of working up the experimental data given in Fig. 1. We have chosen $t_m = 90$ min and calculated for each experimental point the reduced parameters τ and Π^* . Further we calculated, for the found τ values, the theoretical Π^* values by taking certain b - c pairs. In the first approach we took b = 0 and calculated theoretical Π^* values for different *c* ones. In the case of each *c* value the standard deviation (SD) of the experimental Π^* values from the theoretical ones was calculated. By means of a systematic variation of *c* the minimum standard deviation (SD)_m was sought for. Then, the calculations were repeated for other *b* values, too. Thus, a double minimization of the standard deviation has been performed and the b - cpair ensuring the least value for the standard deviation has been taken for the most realistic parameter values.



Deriving of *b* and *c* parameters for the stearic acid benzene solution of $C_0 = 0.051$ mole dm⁻³ at the interface with water phase of pH 2.

This calculation procedure is visualized in Fig. 10 for the stearic acid oil solution of 0.051 mole/L. Curve 1 gives the minimum standard deviation $(SD)_m$ obtained for different b values presumed and curve 2 indicates the corresponding *c* value. Let us take, e.g. *b* = 0.12. Eq. [21] allows us to calculate theoretical Π^* values corresponding to *b* = 0.12 and to different *c* values. In each case, i.e. with each *c* value presumed, the standard deviation:

$$SD = \sqrt{\frac{1}{n} \sum \left(\prod_{t}^{*} - \prod_{e}^{*} \right)^{2}}$$
[27]

is calculated where n stands for the number of points used, and Π_t^* and Π_e^* stand for the theoretical and for the experimental reduced interfacial pressure, respectively. By trying different *c* values, one observed that the standard deviation will be the less for *c* = 0.86, viz. it will be equal to (SD)_m = 0.00928. Thus, we obtained a point on curve 1 and a point on curve 2.

By repeating these calculations for other *b* values, eventually one obtains the whole curve 1 and curve 2. As seen, curve 1 has a minimum (indicated by a downward vertical arrow), corresponding to b = 0.163, this minimum (SD)_m is ensured by a *c* value equal to 0.937 (indicated on the curve 2 by an upward vertical arrow).

By using the *b* and *c* values obtained, $\varphi(1)$ may be easily calculated according to Eq. [20], as well as the corresponding parameter *a* by means of Eq. [23]. Results obtained by means of this procedure are presented in Table VI.

Table VI.

	r	r	1	1	r		r
C ₀	t _m	Πm	φ(1)	а	b	с	(SD) _m
mole dm ⁻³	min	mN m⁻¹					mN m⁻¹
0.001	90	4.5	0.3820	11.78	0.0813	0.706	0.0060
0.00178	90	6.4	0.4096	15.62	0	0.590	0.0046
0.0057	90	12.1	0.3682	32.86	0.0039	0.635	0.0019
0.012	90	15.6	0.3506	44.50	0	0.649	0.0019
0.018	90	18.1	0.3367	53.76	0.0019	0.665	0.0022
0.051	90	22.0	0.2575	85.43	0.163	0.937	0.0054

Parameters of Eq. [18] derived from the experimental dynamic interfacial pressures for SA at the benzene/water of pH 2 interface.

The parameter values given in Table VI allow us to construct by means of Eq. [18] theoretical Π vs. t curves. These curves are given in Fig. 1 as full line curves. Thus, it is demonstrated that Eq. [18] describes very well the experimental Π vs. t data within the experimental errors.

Inspection of Table VI shows that the *a* parameter increases with increasing bulk concentration C_0 of stearic acid. The *c* parameter shows also a clear increasing tendency. The *b* parameter has very small values frequently vanishing ones and cannot be correlated with C_0 .

With respect to the meaning of the parameter *a*, according to Eq. [19] it allows us to calculate the diffusion coefficient D. Let us assume as a first approach that $C_s = 0$ at all bulk concentrations C_0 . By taking into account that $c_0 = C_0 \ 10^{-3} \ N_A$, Eq. [19] becomes:

$$D = \left(\frac{a}{2kT \cdot 10^{-3} \cdot N_A \cdot C_0}\right)^2 \frac{\pi}{t_m}$$
[28]

Diffusion coefficients calculated by means of Eq. [28], and by using the *a* values listed in Table VI, are presented in Table VII.

Table VII.

Diffusion coefficients, D and D₀, calculated by Eq. [28], subsurface concentrations C_s , calculated by Eq. [29], and interaction parameters α calculated by Eq. [19], from the a and c parameters of Eq. [18]. Symbols are defined in the text.

C ₀ mole dm ⁻³	D 10 ¹⁰ cm ² s ⁻¹	D ₀ 10 ¹⁰ cm ² s ⁻¹	C _s 10 ³ mole dm ⁻³	$\alpha \ 10^{22}$ dvne cm ²
0.001	0.3430		0	16.7
0.00178	0.1900		0.45	12.1
0.0057	0.0821	0.343	2.91	9.02
0.012	0.0840		8.22	7.92
0.018	0.0220		13.4	7.38
0.051	0.0069		43.7	8.24

By analyzing the D values listed in Table VII and by comparing them with the values given in Table IV, one can see that the general picture is similar, since D values show a decreasing tendency with increasing bulk concentration of stearic acid and seem to be more realistic, their values being

higher with about one order of magnitude as compared to the previous approach (Eqs. [5] and [6]).

As a second approach we shall presume that $C_s = 0$ only in the case of the surfactant solution of 0.001 mole/L and it increases with increasing bulk concentration, but the diffusion coefficient remains the same D_0 as obtained from the a value for the 0.001 mole/L solution. In this assumption C_s may be calculated from the expression of D given in Eq. [19] and one obtains:

$$C_{s} = C_{0} - \frac{\pi^{1/2} \cdot a}{2kT \cdot 10^{-3} \cdot N_{A} \cdot D_{0}^{1/2} \cdot t_{m}^{1/2}}$$
[29]

The C_s values calculated by Eq. [29] are presented in Table VII and they increase linearly with increasing C₀ as in the case of those C_s values presented in Table IV, but the correlation coefficient is better, r = 0.9994, as compared to r = 0.9983 obtained with C_s values from Table IV. Although the D₀ value (Table VII) is higher with almost one order of magnitude than D₀ value (Table IV), it is far from being realistic. This situation is similar with the adsorption of palmitic acid at the hexane/water interface, for which D = 5.66 10⁻¹³ cm² s⁻¹ was reported (5) and the authors consider that the anomalous D value is due to the existence of an activation barrier for the adsorption between the subsurface and the interface.

Generally, the diffusion coefficient obtained for SA is with at least one order of magnitude smaller than its value previously determined for the two anesthetics, e.g. dibucaine and tetracaine (24, 43). This finding is probably related with the structure of anesthetic molecules, since dibucaine and tetracaine have much shorter hydrocarbon chains and are quite rigid molecules, and their diffusion through the bulk phase and penetration into the oil/water interface might occur easier than for stearic acid molecules.

Eq. [19] allows also the calculation of the interaction parameter α from the parameters *a* and *c* of Eq. [18]. The α values are presented also in Table VII. As seen one does not obtain the same value in the case of differing bulk concentrations, viz. a decreasing tendency is observed with increasing C₀. This indicates, similarly to the too low D₀ value, that the process is not purely diffusion controlled and the potential energetic barrier to adsorption (43, 44) and an interfacial reorientation (45-47) of surfactant molecules could play an important part.

By calculating the mean value of the interaction parameter one obtains $\alpha = 1.02 \ 10^{-21}$ dyne cm². It is worth mentioning that for stearic acid monolayers spread at the air/water interface we obtained 4.3 10^{-21} dyne cm² (42). In the case of the air/water interface in the air phase strong hydrophobic interactions appear between the long hydrocarbon chains, which are almost absent at the

benzene/water interface, when the hydrocarbon chains are solvated with benzene molecules hindering the direct interaction between the neighboring SA molecules. Consequently the α value must be much less as compared to the air/water interface. Thus, the above α value seems to be in substantial agreement with our earlier results (42).

The low values of the diffusion coefficients can be further correlated with the presence of an energy barrier of adsorption due to a slow reorientation (7, 38) of the polar head groups of SA molecules at the interface to finally achieve their proper arrangement or their correct orientation at the adsorption equilibrium. This effect can also explain the slow interfacial pressure increase to reach its equilibrium value.

Therefore, these data represent strong experimental evidence about the non-ideal behavior of adsorbed SA layers at the oil/water interface and about the interfacial reorientation of SA molecules at the oil/water interface, which apparently operates for the adsorption of fatty acids from oil solution to an oil/water interface. It seems that the adsorption of SA from benzene to the benzene/water interface is of this type in agreement with the adsorption of palmitic acid at the hexane/water interface reported earlier (5). Hence, the adsorption process is controlled by the molecular diffusion in the oil phase associated with the surface reorientation and the molecular interactions of stearic acid molecules at the oil/water interface.

Conclusions

A theoretical approach to the dynamic interfacial pressures is developed and a new kinetic Eq. [18] for the diffusion controlled kinetics is proposed, based on Ward and Tordai's diffusion equation associated with a two dimensional van der Waals state equation. This approach is consistent with the case of the non-ideal behavior of the adsorbed films widely used in the kinetic of adsorption of surfactants from concentrated bulk solutions to the liquid interfaces. This provides a possibility for one to determine the diffusion coefficients, subsurface concentrations and intermolecular interaction parameters within adsorbed films of surfactants at the liquid/liquid interfaces. The theory is applied to describe and fit the experimental data of the dynamic interfacial pressures for the adsorption of stearic acid (SA) at the benzene/water interface. Obtained results are in general agreement with similar earlier reported data for the adsorption of palmitic acid (5) and the adsorption of chlorophyll (22) at the different oil/water interfaces.

The adsorption of stearic acid (SA) at the benzene/water interface seems to be a process in which the rate is controlled both by the molecular diffusion from the bulk to the subsurface and by the molecular interactions in adsorbed SA layer at the liquid/liquid interface since both Eqs. [7] and

[18] yield interesting pieces of information, but none of them gives a perfect description of the phenomenon.

The low D values for SA further suggest the presence of an energy barrier of adsorption between the subsurface and the oil/water interface. This barrier might be attributed to the steric restraints on the stearic acid molecules, in the proximity of the oil/water interface, and therefore, the molecules need to achieve a correct orientation to adsorb at the interface. This barrier of adsorption can be also due to a slow reorientation of the polar head groups of SA molecules at the oil/water interface. This will cause an increase in the timescale for the dynamic interfacial pressure to reach out its equilibrium value.

Eq. [18] proposed in the present paper is better than Eq. [6] and it allows a more correct description of the experimental data of the dynamic interfacial pressures for a large interval of surfactant concentrations in oil solutions and over a period of time that can range up to two hours, depending on the stearic acid bulk concentration. In this period of time, the dynamic interfacial tension, $\sigma(t)$, will decay to reach its equilibrium value and, consequently, the dynamic interfacial pressure, $\Pi(t)$, will increase to attain its equilibrium value.

In future studies we intend to extend this investigation by using the diffusion adsorption theory associated with the surface reorientation of the fatty acid molecules at the oil/water interface.

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ERRATUM

KINETICS AND MECHANISM OF *P*-XYLENE OXIDATION BY CE(IV) IN AQUEOUS ACID MEDIUM. LFER AS AN ARGUMENT TO THE OXIDATION MECHANISM

GABRIELA-CRISTINA BUCŞA AND IOAN BÂLDEA

(Article appeared in Studia Univ. Babeş-Bolyai. Chem. L (1), 2005, 157-168)

Some errors were discovered in the article mentioned concerning the mechanism of the title reaction (p.165) and the rate law from the mechanism (eqn. 13). The correct mechanism and rate law are as follows:

$$Ce^{4+} + H_{2}O \iff CeOH^{3+} + H^{+} \qquad k_{1}, k_{-1}, \quad K_{h}$$

$$Ce(OH)^{3+} + CH_{3}C_{6}H_{4}CH_{3} \iff Adduct^{3+} \qquad k_{2}, k_{-2}, \quad K_{f}$$

$$Adduct^{3+} + H^{+} \xrightarrow{r.d.s.} [CH_{3}C_{6}H_{4}(H)CH_{2}^{o}]^{+} + Ce^{3+} + H_{2}O \qquad k_{3}$$

$$[CH_{3}C_{6}H_{4}(H)CH_{2}^{o}]^{+} + Ce(OH)^{3+} \xrightarrow{fast} Ce^{3+} + CH_{3}C_{6}H_{4}CH_{2}^{+}H_{2}O$$

$$CH_{3}C_{6}H_{4}CH_{2}^{+} + H_{2}O \xrightarrow{fast} CH_{3}C_{6}H_{4}CH_{2}OH + H^{+}$$

$$r = \frac{d[Ce(IV)]_{t}}{d_{t}} = \frac{k_{3}K_{2}[p - xylene][H^{+}]}{1 + K_{p}^{-1}[H^{+}] + K_{2}[p - xylene]} [Ce(IV)]_{t} (13)$$

The rate law has the same form as the published one and the conclusions remain the same.

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STUDIA UNIVERSITATIS BABES - BOLYAI, CHEMIA, LI, 2, 2006

Dedicated to professor Gh. Marcu at his 80th anniversary

ELECTROCHEMICAL CHARACTERIZATION OF SANDWICH-TYPE PHOSPHOTUNGSTOCERATE (IV) AND ARSENOTUNGSTOCERATE (IV) IN AQUEOUS AND NON-AQUEOUS SOLUTION

ADRIAN-RAUL TOMSA^{ab}, AGLAIA KOUTSODIMOU,^b POLYCARPOS FALARAS,^b MARIANA RUSU^c*

ABSTRACT. The electrochemical properties of two sandwich-type polyoxotungstocerate (IV) $[(CeO)_3(H_2O)_2(PW_9O_{34})_2]^{12}$ and $[(CeO)_3(H_2O)_2(ASW_9O_{34})_2]^{12}$ have been studied by cyclic voltammetry in aqueous and acetonitrile solutions. Voltammograms recorded in aqueous solutions showed the existence of two waves for the Ce(III)/Ce(IV) redox couple. These could be explained by a nonequivalence of the Ce atoms in the polyoxotungstate molecule, where one Ce atom is six coordinated and the other two, are seven coordinated. The presence of only one peak pair for Ce(III)/Ce(IV) suggests the equivalence of these metal ions in the polyoxometalate complexes when isolated as tetrabutylammonium salts. When increasing amounts of LiCIO₄ have been added to the acetonitrile solutions of the polyoxotungstocerate while keeping the ionic strength constant with 1M [Bu₄N]CIO₄ a dramatic change of the electrochemical behavior was observed, which was assigned to the association of Li⁺ ions with the reduced forms of the polyoxometalates.

Introduction

The interest regarding the polyoxometalates chemistry has increased enormously in last decades when a very large number of different structures as well as new potential applications of such compounds have been reported [1-8]. On this context, it is recognized that the different coordination geometries of the lanthanide cations and the vacant sites afforded by the polyoxometalates usually result in large oxometalate clusters [4-24], showing very interesting electroluminiscence and photoluminiscence activity [4, 11-14, 16, 22] as well as enhanced efficiency against HIV [25, 26].

One example of such compound, in which lanthanides act as linkers between trilacunary Keggin polyoxometalates was described for the first time by Knoth et al. in 1986 [27]. Its structure consists on a belt of three Ce alternating with three oxygens, sandwiched between two trilacunary 9-phosphotungstate units. In the belt, two of the cerium (IV) ions are 7 coordinated, and the third one is only 6 coordinated (Fig. 1).

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Fig.1. The polyhedral representation of the sandwich polyoxotungstocerates (WO₆ white octahedral; Ce – black spheres; P – grey spheres; O- white spheres).

Herein we report a detailed investigation of electrochemical behavior of $[(CeO)_3(H_2O)_2(PW_9O_{34})_2]^{12-}$ and its arsenic (V) derivative, in aqueous and non-aqueous solutions.

Experimental

1. Methods

Inductively Coupled Plasma Spectroscopy (I.C.P.) was used for the elemental analysis of tin and tungsten, and flamephotometry for sodium and potassium. The water content was thermogravimetrically determined, by means of a Paulik Erdely OD-102 type derivatograph at a temperature range of 20-600°C, using a heating rate of 5°C min⁻¹.

A Nicolet 5DX spectrophotometer was used to record the IR spectra of samples pelleted in KBr.

Cyclic voltammetry measurements were performed using a Vinci Technologies P/G STAT Z1 potentiostat, operating in conjunction with a 33120 A Hewlett Packard function /arbitrary waveform generator and with a Linseis LY 18100 X-Y recorder. A conventional (single-compartment) three electrode electrochemical cell was used, with a Metrohm planar platinum counter electrode and a reference electrode. For the aqueous solutions a Metrohm saturated calomel (Hg/Hg₂Cl₂) (SCE) was used as reference electrode, while for non-aqueous solution, a Metrohm Ag/AgCl electrode (a silver wire covered by AgCl, immersed in a saturated LiCl solution in acetonitrile, which was separated from the studied solution by a salt bridge of 0.1 M [Bu₄N]ClO₄ in acetonitrile). Pyrolytic graphite (Union Carbide, Φ ~ 3mm) was used as a working electrode (WE). This was polished with emery paper 4000 and cleaned in an ultrasonicator, prior to use.

For aqueous solutions, the WE was polarized in a 10^{-3} M solution of the complex under study in phosphate buffer (pH = 4.0 - 5.5) with a scan rate between 50 and 400 mV s⁻¹.

For non-aqueous solutions, the WE was polarized in 10^{-3} M acetonitrile solution of the complex in the presence of 0.1 M [Bu₄N]ClO₄.

Solutions were deoxygenated using purified Ar gas (99.9%). While recording the voltammograms, Ar was passed over the solution surface.

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2. Materials

Reagent grade chemicals were used and all syntheses and studies were carried out in distilled water.

The potassium salts $K_{10}H_2[(CeO)_3(H_2O)_2(PW_9O_{34})_2]\cdot 18H_2O$ and $K_{10}H_2[(CeO)_3(H_2O)_2(AsW_9O_{34})_2]\cdot 21H_2O$ have been prepared by the reported procedures [27, 28] and their purity have been proved by chemical analysis and IR spectra.

Synthesis of [Bu₄N]₁₀H₂[(CeO)₃(PW₉O₃₄)₂]

To a solution of 1.0 g $K_{10}H_2[(CeO)_3(H_2O)_2(PW_9O_{34})_2]$ 18H₂O (0.176 mmol) in 30 mL hot water at pH= 4.5-5.0, 2.0 g [Bu₄N]Br (6.204 mmol) were added under vigorous stirring. A pale yellow precipitate was formed, which was removed by vacuum filtration on G₃ frite, washed with 10 mL distilled water, dried under suction for 2 h and then over P₂O₅ for 24h. Yield: 1.23 g (95.45%).

Calcd. for: $[(C_4H_9)_4N]_{10}H_2[(CeO)_3(PW_9O_{34})_2]$ P: 0.84; W: 45.00; Ce: 5.71. Found: P: 0.77; W: 45.22; Ce: 5.62.

Synthesis of $[Bu_4N]_{10}H_2[(CeO)_3(AsW_9O_{34})_2]$

To a solution of 1.0 g $K_{10}H_2[(CeO)_3(H_2O)_2(AsW_9O_{34})_2]$ 21 H_2O (0.171 mmol) in 50 mL hot water at pH= 4.5-5.0, 2.0 g [Bu₄N]Br (6.204 mmol) were added under vigorous stirring. A pale yellow precipitate was formed. which was removed by vacuum filtration on G₃ frite, washed with 10 mL distilled water, dried under suction for 2 h and then over P_2O_5 for 24h. Yield: 1.22 g (95.84%).

Calcd. for: [(C₄H₉)₄N]₁₀H₂[(CeO)₃(AsW₉O₃₄)₂] As: 2.01; W: 44.46; Ce: 5.65. Found: As: 1.89; W: 44.61; Ce: 5.58.

Results and Discussions

The cyclic voltammograms of the $[(CeO)_3(H_2O)_2(XW_9O_{34})_2]^{12}$ (X= P, As) complexes recorded in aqueous solutions at pH= 4.0 are depicted in Fig. 2.

Both complexes showed two successive redox peak pairs in the negative potential range, which were attributed to two single electron tungsten centered processes [29]. In this range, the cyclic voltammograms were very similar to that of the parent anion, indicating a lack of influence by the Ce(IV) present in the polyoxometalate framework. However a negative shift of the peaks provides proof for the coordination of the Ce(IV) to the trilacunary polyanions.

Two additional successive redox peak pairs appear in the positive range, which were assigned to two single electron cerium-centered processes [5]. The existence of two waves for the Ce(III)/Ce(IV) redox couple could be explained by the nonequivalence of the Ce atoms in the polyoxotungstate framework, where one Ce atom is six coordinated and the other two are seven coordinated [30].

The formal standard potentials $\epsilon^{0'}$ (estimated as the average of anodic and cathodic peak potentials) of the tungstophosphate complex are negatively shifted by comparing with its tungstoarsenate analog (Table 1) which is in agreement with previous reports [31].

The positive shift of the formal standard potentials $\epsilon^{0'}$ observed in Ce(IV) complexes when compared to the Ce(III) analogs [30], could be due to the lower negative charge of the Ce(IV) polyoxometalate complexes.

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Fig. 2. Cyclic voltammograms of $K_{10}H_2[(CeO)_3(OH_2)_2(PW_9O_{34})_2]$ ·18H₂O (a) and $K_{10}H_2[(CeO)_3(OH_2)_2(AsW_9O_{34})_2]$ ·21H₂O (b) recorded in aqueous solutions (pH= 4.0, v= 50 mV s⁻¹).

Table 1.

CVUa	ata ioi the $[(CeO)_3$	$(\Pi_2 \cup)_2 (\land V V_9 \cup_{34})_2$	in aqueous soit	luons.
Process	E _{p,a} (V)	$E_{p,c}(V)$	(V)	$E_{p}(V)$
	K ₁₀ H ₂ [(C	CeO)3(OH2)2(PW9O34)2]	·18H₂O	
ľ	0.575	0.510	0.542	0.065
'	0.399	0.379	0.389	0.020
	-0.800	-0.852	-0.826	0.052
	-0.921	-0.999	-0.960	0.078
	K ₁₀ H ₂ [(C	eO)3(OH2)2(AsW9O34)2]·21H₂O	
ľ	0.812	0.738	0.775	0.074
'	0.631	0.610	0.620	0.021
	-0.619	-0.710	-0.664	0.091
	-0.801	-0.836	-0.818	0.035

CV data for the $[(CeO)_3(H_2O)_2(XW_9O_{34})_2]^{12-}$ in aqueous solutions.

^a E_{pa} , oxidation potential; E_{pc} , reduction potential; $\varepsilon^{0'}$ formal standard potential estimated as the average of anodic and cathodic peak potentials; ΔE_p the difference between the redox peak potentials. Experimental conditions: H₃PO₄-Na₂HPO₄ buffer solution; pH 4.0, 25°C; graphite working electrode; scan rate 100 mV·s⁻¹; c= 10⁻³M, potentials recorded *vs* SCE.

The linear relationships between peak currents and the square root of the scan rates for redox waves I (Fig. 3) suggest the diffusion control of the electrochemical processes [32].

Changes in the pH resulted only in minor variations of the formal standard potential ϵ^{0} , which imply that protons were not involved in the redox processes [30].

In order to investigate the influence of the different cations on the electrochemical behavior of the polyoxotungstocerate (IV) complexes, their tetrabutylammonium salts have been studied in acetonitrile solutions.

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Fig. 3. Dependence of the cathodic peak I current on the scan rate for: $K_{10}H_2[(CeO)_3(OH_2)_2(PW_9O_{34})_2]$ ·18H₂O (a) and $K_{10}H_2[(CeO)_3(OH_2)_2(AsW_9O_{34})_2]$ ·21H₂O (b) (pH=4.0).



Fig. 4. Dependence of the formal standard potentials on the pH for: $K_{10}H_2[(CeO)_3(OH_2)_2(PW_9O_{34})_2] \cdot 18H_2O(a)$ and $K_{10}H_2[(CeO)_3(OH_2)_2(AsW_9O_{34})_2] \cdot 21H_2O(b)$ (v = 100 mV s⁻¹).

When 0.1 M [Bu₄N]ClO₄ was used as supporting electrolyte, only one pair of waves were observed in the negative potential range for 9-tungstophosphate derivative, while the 9-tungstoarsenate analog exhibited two pair of peaks (Fig. 5). These were assigned to W(VI)/W(V) processes in the polyoxometalate framework. Both complexes, in acetonitrile solution showed only one peak pair in the positive potential range, due to the Ce(III)/Ce(IV) processes, suggesting equivalence of all the cerium ions present in the structure.

When increasing LiClO₄ quantities were added to the acetonitrile solutions of the polyoxometalates, while keeping the ionic strength constant at 0.1 by adding [Bu₄N]ClO₄, dramatic changes were observed in the voltammograms (Fig. 6). At high Li⁺ concentrations, two additional peaks (II and III) appeared in the negative potential range, while the peaks I and I' shifted to more positive values.

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Fig. 5. Cyclic voltammograms of $[Bu_4N]_{10}H_2[(CeO)_3(PW_9O_{34})_2]$ (a) and $[Bu_4N]_{10}H_2[(CeO)_3(AsW_9O_{34})_2]$ (b) (0.1 M $[Bu_4N]CIO_4$ in acetonitrile, v= 100 mV s⁻¹).

The positive shift of the peaks observed upon increasing the Li^{+} concentration could be attributed to the association of Li^{+} ions with the reduced forms of the polyoxometalates [33, 34].

Table 2.

• •		/3(****9@34/2]	~.	
Process	E _{p,a} (V)	E _{p,c} (V)	(V)	$E_{p}(V)$
	[Bu ₄ l	N]10H2[(CeO)3(PW9C)	D ₃₄) ₂]	
'	0.212	0.268	0.265	0.056
	-0.582	-0.470	-0.526	0.112
	[Bu ₄ N	I]10H2[(CeO)3(AsW9	O ₃₄) ₂]	
'	0.461	0.584	0.522	0.123
	-0.744	-0.593	-0.668	0.151
I	-1.009	-0.865	-0.937	0.144

CV data for the $[(CeO)_3(XW_9O_{34})_2]^{12-}$ in acetonitrile solutions.

 E_{pa} , oxidation potential; E_{pc} , reduction potential; $\varepsilon^{0'}$ formal standard potential estimated as the average of anodic and cathodic peak potentials, ΔE_{p} the difference between the redox peak potentials. Experimental conditions: 0.1 M [Bu₄N]ClO₄ acetonitrile solution; 25°C; graphite working electrode; scan rate 100 mV·s⁻¹; c= 10⁻³M, potentials recorded vs Ag/AgCl.



Fig. 6. Cyclic voltammograms of $[Bu_4N]_{10}H_2[(CeO)_3(PW_9O_{34})_2]$ (a) and $[Bu_4N]_{10}H_2[(CeO)_3(AsW_9O_{34})_2]$ (b) at different concentrations of LiClO₄ (0.1 M [Bu₄N]ClO₄ in acetonitrile, v= 100 mV s⁻¹).

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Fig. 7. Dependence of the formal standard potentials on the Li⁺ concentration for: $[Bu_4N]_{10}H_2[(CeO)_3(PW_9O_{34})_2]$ (a) and $[Bu_4N]_{10}H_2[(CeO)_3(AsW_9O_{34})_2]$ (b) (v= 100 mV s⁻¹).

The liniar relationships between the Log([Li⁺]) and the formal standard potential $\epsilon^{0'}$, with slopes of 0.078 (I'), 0.064 (I) and 0.096 (II) for the phosphotungstate derivative, and 0.087 (I'), 0.056 (I) and 0.075 (II) for its arsenotungstate analog, suggest that one Li⁺ cation is envolved in each redox process, according to the following equations:

 $\begin{array}{l} \textbf{I}^{\prime}: [(Ce^{IV}O)_{3}(XW_{9}O_{34})_{2}]^{12-} + e^{-} + Li^{+} \leftrightarrow Li[(Ce^{III}O)(Ce^{IV}O)_{3}(XW_{9}O_{34})_{2}]^{12-} \\ \textbf{I}: Li[(Ce^{III}O)(Ce^{IV}O)_{3}(XW_{9}O_{34})_{2}]^{12-} + e^{-} + Li^{+} \leftrightarrow Li_{2}[(Ce^{III}O)(Ce^{IV}O)_{3}(XW_{9}O_{34})_{2}]^{12-} \\ \textbf{II}: Li_{2}[(Ce^{III}O)(Ce^{IV}O)_{3}(XW_{9}O_{34})_{2}]^{12-} + e^{-} + Li^{+} \leftrightarrow Li_{3}[(Ce^{III}O)(Ce^{IV}O)_{3}(XW_{9}O_{34})_{2}]^{12-} \end{array}$

Conclusions

The results of the performed investigations revealed that the electrochemical properties of the sandwich-type polyoxotungstocerate (IV) $[(CeO)_3(H_2O)_2(PW_9O_{34})_2]^{12}$ and $[(CeO)_3(H_2O)_2(AsW_9O_{34})_2]^{12}$ are different in aqueous and non-aqueous solution, suggesting different coordination of the cerium (IV) in aqueous and non-aqueous media. The addition of increasing amounts of the Li⁺ cations results in a positive shift of the peaks which could be explained by the association of lithium ions with the reduced form of the polyoxotungstocerate.

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SYNTHESIS, SPECTRAL AND THERMAL STUDIES OF {[2-[(2,6-DICHLOROPHENYL)AMINO]PHENYL} ACETATE OF RUTHENIUM (III) AND RHODIUM (III)

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ABSTRACT. The reactions of RuCl₃ and RhCl₃ with deprotonated diclofenac (L) were studied in aqueous solution. Coordination compounds of the formulae [Ru(L)₃ (H₂O)₃]3H₂O and [Rh(L)₃ (H₂O)₃]2H₂O were isolated as solid products and characterized by elemental analyses and spectroscopic (IR, UV-VIS) and thermal studies. In all studied compounds, {2-[(2,6-dichlorophenyl)amino]-phenyl}acetate acts as monodentate ligand with coordinate involving the carboxylate oxygen atom.

INTRODUCTION

Diclofenac, sodium {2-[(2,6-dichlorophenyl)amino]phenyl}acetate is a potent non-steroidal antiinflammatory drug (NSAID), therapeutically used in inflammatory and painful diseases of rheumatic and non-rheumatic origin. The anti-inflammatory activity of diclofenac and most of its other pharmacological effects are thought to be related to the inhibition of the conversion of arachidonic acid to prostaglandins, which are the mediators of the inflammatory processes [1,2]. Like other NSAIDs, diclofenac is highly (>95%) protein bound. [1]

The interaction of metal ions with drugs administered for therapeutic reasons is a subject of considerable interest. It is known that the drugs act via chelation or by inhibiting the activity of metalloenzymes, but for most of the drugs such as diclofenac little is known about how metal binding inflences their activity.

As part of our research on understanding drug-metal interaction, we have studied the coordination ability of diclofenac with trasition metal ions.

EXPERIMENTAL

Spectrophotometric titration

Electronic absorption spectroscopy is frequentely used to study metal binding to the ligand. [3-8]

Spectrophotometric titration of ruthenium (III) and rhodium (III) with sodium {[2-(2,6 dichlorophenyl) amino] phenyl}acetate is presented.

Preparation of compounds

The coordination compounds were prepared by mixing hot aqueous solutions of the ligand diclofenac (sodium salt tetrahydrate) (pH~5,5-6,5) and aqueous solutions of metal salts (3:1 ligand to metal molar ratio). The reaction mixture was stirred for 5h at 323K. The resulting powders were filtered, washed with hot water to remove Na⁺ and Cl⁻ ions, dried at 303K to a constant mass. The yields and the elemental analyses are presented in Table 1.

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Table 1.

Compound	GM	Yield	Elemental analysis, [%] found, (calcd.)				
		[%]	С	Н	Ν	CI	М
[Na(H ₂ O) ₄] ⁺ L ⁻ 1	390,07		43,15 (43,06)	4,64 (4,61)	3,55 (3,59)	17,40 (18,17)	5,57 (5,89)
[Ru(L)₃(H₂O)₃]·3H₂O 2	1093,87	91,6	46,15 (46,07)	3,96 (3,84)	3,50 (3,84)	19,40 (19,44)	9,61 (9,24)
[Rh(L) ₃ (H ₂ O) ₃]·2H ₂ O 3	1077,42	92,4	46,59 (46,77)	4,10 (3,90)	3,52 (3,90)	19,21 (19,64)	9,50 (9,55)

Analytical data of compounds.

Where $L = (C_{14}H_{12}O_2NCI_2)^{-}$.

Measurements

The carbon, hydrogen and nitrogen content in the coordination compounds were determined by elemental analyses using a Perkin Elmer CAN 2000 analyzer. The chlorine content was measured by the Schöninger method. The Ru(III) and Rh(III), contents was determined by AAS method using an Atomic Absorbtion Spectrometer AAS-3 (Carl Zeiss, Jena). The experimental results are concordant with the calculated data (Table 1).

Thermal analysis were performed with an OD-103 MOM Derivatograph using a sample weight of $100\pm1-2$ mg, at a heating rate of 10 K min⁻¹, with Al₂O₃ as reference material in static air atmosphere.

IR spectra were recorded over the range of 4000-400 cm⁻¹ using a FT-IR JASCO 600 spectrophotometer, in KBr pellets. UV-VIS spectra were recorded with an UV-VIS Specord Spectrophotometer Carl Zeiss Jena [1].

RESULTS AND DISCUSSION

The coordination compounds: $[Ru(L)_3(H_2O)_3] \cdot 3H_2O$ (2), and $[Rh(L)_3(H_2O)_3] \cdot 2H_2O$ (3), were formed according to egs:

 $MCI_3 + 3NaL + nH_2O \rightarrow [M(L)_3(H_2O)_3]^{-} nH_2O + 3NaCI$

were M=Ru(III), Rh(III) and n=3,2.

The compounds are brown, microcrystaline powdery solids, that appear to be air- and moisture-stable. They are soluble in methanol, ethanol, acetone, benzene, DMF, DMSO and insoluble in water. The colour of the coordination compounds is typical of the particular ion salts, i.e. Ru (III) and Rh (III), having their origin in the lowest energy of $\underline{\mathbf{d}} - \underline{\mathbf{d}}$ electronic transitions of the central ions [7]. The structure of coordination compounds 2 and 3 is supported by thermal analysis, IR and UV-VIS spectroscopic data.

UV-VIS spectra

The sodium diclofenac, is a colorless compound with an intense ultraviolet absorbtion band in the 214nm region, due to the allowed intraligand π - π transitions and an additional band in the 270 nm region, assigned to the n- π transition [24,25].

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The binding of metal ions to the carboxylate group of the diclofenac leads to the production of two new absoption bands at ca 236 nm and 285 nm in the UV-difference spectra.

Table 2.

Electronic spectral data (nm) of the ligand and their coordination compounds in aqueous solution

Compound	λ (nm (ε)			
1	214 (6000)	270 (8000)		
2	230 (6500)	288 (8200)	500 (800)	604 (780)
3	239 (6550)	285 (8250)	515 (805)	608 (790)

The isobestic point situated at 236 nm (Fig.1) point out the formation of the metal coordination compound.

IR –spectra

The vibration modes of coordinated ligands in the complexes have been investigated by means of infrared absorbtion spectra. The most important infrared frequencies attributed to the vibrations of the compounds 1 - 3 are reported in Table 3.

The absorption bands v_{OH} and v_{HOH} which occur in the range 3482 - 3495 cm⁻¹, confirm the presence of water of crystallization. The absorbtion bands $vM-O_{H_2O}$ which

occur in the range 425 – 418 cm⁻¹ confirm the presence of coordinated water in the complexes[12]. The presence of water as water of crystallization and as coordinated water in the compounds is confirmed by the thermal decomposition data.

As the carboxylic hydrogen is more acidic than the amino hydrogen, the deprotonation occurs in the carboxylate group. This is characteristic bands for the secondary amino groups and for the coordinated carboxylate group [5,13–15]. The strong band at 3388 cm⁻¹, which appears in diclofenac, is assigned to the stretching motion and the broad band at 3260 cm⁻¹ is taken to represent the v_{NH...O} mode, due to intramolecular hydrogen bonding [5].The absence of large systematic shifts of the v_{NH} and δ_{NH} bands in the spectra of the coordination compounds **2,3** compared with those of the ligand indicates that there is not interaction between the NH group and the metal ions.





Fig.1. Titration curves of ruthenium (III) chloratum with Sodium diclofenac

Table 3.

Frequences of characteristic absorbtion bands in IR spectra of sodium {2–(2,6diclorophenyl)amino]phenyl}acetate tetrahydrate and its coordination compounds (cm⁻¹).

Compoun d	ν ΟΗ; ν ΟΗΟ	vas COO ⁻	V _{sCOO}	Δν _{COO}	$vM - O_{H_2O}$	vM − O _{coo} -
1	3560 br	1572 s	1402 s	170	410 mw	-
2	3482 br	1670 s	1346 s	324	425 mw	403 m
3	3495 br	1683 s	1350 s	333	418 w	-402 m
* s = strong, m = medium, w = weak, br = broad						

The IR spectra bring evidence of the carboxylate ions modus/type to metal ions such as unidentate, bidentate (chelating) or bridging and there is an evidence of that fact in the IR spectrum. The analysis of COO⁻ group bands frequencies allowed the determination of parameter $\Delta COO^- = v COO^-$ (as) - $v COO^-$ (s).

The separations between v_{as} and v_s , Δv , in a bidentate (chelate) complex will be significantly smaller than in the free ion; in the bridging complex the Δv value is that of the to the free ion, while in the monodentate complex it will be higher[13,14]. For complexes **2** and **3** the difference $\Delta v > 170 \text{ cm}^{-1}$ is higher than that of the free ion (sodium diclofenac salt) and is assigned to the stretching mode

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for the monodentate carboxylate ligation. Calculated from the examined spectra, the values of Δv of the coordination compounds **2** and **3** occur in the range domain 324 - 333 cm⁻¹. These values of two bands (COO⁻ deformation) in coordination compounds **2** and **3** are in good agreement with the literature data for unidentate bonded acetates structures [24]. The absorbtion bands which occur in the range 402– 405 cm⁻¹ v(M-O) confirm the coordination of diclofenac to metallic ions through the oxigen atom of the carboxylate group.

Thermal behavior

The thermal behavior of the ligand and of the synthesized coordination compounds **2** and **3** is presented in Table 4. Thermal stability domains, decomposition phenomena (endo and exo effects in the DTA curves and mass losses, calculated from the TG and DGT curves) and their assignment are presented.

Between 293-473 K an endo peak at 403 K in the TDA curve of $[Na(H_2O)_4]^+L^$ indicates the loss of four water molecules. X-ray scattering studies[17] show that in various criystalline salts, Na⁺ forms tetrahedral $[Na(H_2O)_4]^+$ ions, with the 4H₂O molecules directly coordinated. These results are in agreement with structure and thermal behaviour of sodium diclofenac salts studied by [18-21].

Table 4.

Compound	Temp. Range, K	DTA p Endo	eak, K Exo	TG o Calc.	lata, % Exp	Assignment
1	293 - 473	403	-	18,45	18,21	4H ₂ O
	473 – 1273	555	-	-	-	Melting
		-	558	11,28	11,15	CO_2
		-	673	55,22	55,46	Pyrolysis of organic rest
		1048	-	14,98	14,71	NaCl residue
2	293 – 473	371	-	2,73	2,66	3H ₂ O
		450	-	2,73	2,68	3H ₂ O
	473 – 1273	-	511	2,22	2,31	CO ₂
		-	685	83,08	83,03	Pyrolysis of organic rest
		-	1053	15,22	15,75	Ru ₂ O ₃ residue
3	293 – 473	378	-	3,34	3,20	2H ₂ O
		435	-	5,02	5,08	3 [.] H ₂ O
	473 – 1273	-	508	4,08	4,03	CO_2
		-	692	79,01	79,22	Pyrolysis of organic rest
		-	1060	11,78	11,34	Rh ₂ O ₃ residue

Thermal data of the synthesized coordination compounds

The anhydrous sodium diclofenac, NaL is stable up to 555 K where an endo peak marks its melting. The decomposing starts with the shoulder at 558 K which was assigned to the lose of one carboxylate group. In the temperature range 473-1273 K an exothermic peak situated at 673 K marks the pyrolysis of the organic residue. At 1048 K the formation of NaCl residue is observed. The coordination compounds synthesized, **2** and **3** are stable in air and can be stored without change. When heated in air, the coordination compounds decompose in various ways (Table 4). According to literature data[12,22-25] water released below 423 K can be considered as crystallization water, whereas that eliminated above 423 K, as chemically bounded to the central ion through weak coordination bonds.

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The decomposition of the coordination compounds **2** and **3** occurs in the 435-450K temperature range, when the coordination water separate from each molecule. The stepped dehydration reaction is observed, the two endo peaks at 371K, 378K and 450K, 435K corresponding to the loss of three and two molecules of hydrating water and three molecules of coordination water, respectively.



Fig.2. TG, DTG and DTA curves of the coordination compound 2

The thermal decomposition with the shoulder at 511K and 508 K indicate the destabilization of the ligand due to the weakening of the carboxylate bound as a consequence of M-O bond formation in the coordination compounds.

An exothermic peak situated at 685K and 692K, marks the pyrolysis of the organic residue. In the 720K-1273K temperature range, the formation of intermediated is observed, followed by the thermal decomposition to oxides: Ru₂O₃, Rh₂O₃ respectively.

The thermal analyses results of the investigated compounds confirm the atom ratio M:L:O = 1:3:3.

CONCLUSIONS

The coordination compounds 2 and 3 are stable in air and soluble in methanol, ethanol and benzene. Heating the compounds first results in a release of crystallization water molecules in complexes 2 and 3. The decompozition of the compounds 2 and 3 occurs with the loss of the coordination water and is continued with thermal decompozition of the ligand (L). The results reveal that metallic

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oxides are left as residues. Infrared data are in accordance with the literature data for unidentate bonded acetates structures of complexes **2** and **3** [23,24]. Diclofenac is coordinated to metal (III) through the oxygen atom of its carboxylate group.

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THE USE OF ORGANODITHIOPHOSPHORIC ACIDS FOR SEPARATION AND DETERMINATION OF SOME METAL IONS FROM RESIDUAL WATERS

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ABSTRACT. The U(VI), Cu(II) and Ni(II) separation was performed by thin layer chromatography on silica gel H, using o,m,p-xylene, methyl-ethyl-ketone (MEK) and N,N-dimethylformamide (DMF) (16: 2: 1, v/v) mixture as mobile phase. The di(n-pentyl)dithiophosphoric acid was used as complexing agent introduced in the mobile phase. This method was applied for quantitative determination of studied metal ions from waste waters.

Key words: thin layer chromatography, metal ions, di(n-pentyl)dithio- phosphoric, quantitative determination

INTRODUCTION

In the last years many studies are devoted to the identification, separation and determination of trace metal ions in particular samples, because of their strong environment impact. Uranium, natural occurring actinide, is found at a trace level in the environment or associated with other metal ions in complex mixtures, nuclear fission products, and geological materials. Many methods have been proposed for the separation and determination of uranium and other metals. One of these techniques, more suitable due to the simplicity in handling radioactive elements is chromatography. In particular, extraction chromatography which combines the selectivity of organic extractants with multistage feature of chromatographic process has been extensively applied in the analysis of the radioactive materials [1-3].

In recent years thin layer chromatography (TLC), widely applied in pharmaceutical and food industries, biochemistry, and has been used in metal ions analysis because of their simplicity, low cost of qualitative analysis and short time of separation. TLC technique has been employed using different complexants fixed on silica gel or another supports or included in organic solvents used as eluents [4-9].

Organophosphorus compounds have been intensively investigated as complexants in separation of metal ions [8, 9]. Their analogues when one or more oxygen atoms have been replaced by sulphur atoms have been used in lesser extent.

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Dialkyldithiophosphoric acids are well known as good complexing agents for metal ions [10-12]. The metal chelates formed with organodithiophosphate anions are soluble in organic solvent which make these ligands useful in liquid-liquid extraction [13-18].

Dialkyldithiophosphoric complexants containing different alkyl chains were also investigated as extractants in the chromatographic technique. In our earlier work we have been investigated the extraction of U(VI), Th(IV) and lanthanides with various dialkyldithiophosphoric acids [19-23] and later we extended our investigations on the separation of metal ions by electrophoresis [24] and TLC [25-29].

The work discussed in this paper continues our investigations on the separation of the metal ions by TLC using dialkyldithiophosphoric acids. The work is undertaken to obtain information about TLC behavior of U(VI), Cu(II) and Ni(II) on silica gel H using di(n-pentyl)dithiophosphoric acid (HDPDTP) in mobile phase, in order to establish the optimum conditions for separation and quantitative determination of these elements in different water samples.

EXPERIMENTAL DATA

Apparatus

Normal chromatographic chambers, Brand micropipettes (5 μ L), densitometer (DESAGA CD 60), Varian Tectron A 66 Spectrometer and Spekol Zeiss Yena were used.

Materials

TLC plates coated with silica gel H ("Raluca Ripan" Institute for Research in Chemistry), Arsenazo III and uranium nitrates (Sigma-Aldrich Chemie, Steinheim, Germany) were used. Copper and nickel nitrates and all other reagents were of analytical grade supplied by "Chimopar" Bucharest, Romania. HDPDTP were synthesized according to published procedure [30] and its purity was higher than 95%. Test solutions of metal nitrates were standardized gravimetrically and concentrations were obtained by successive dilutions.

Procedure

Silica gel H was used as stationary phase for separation of U(VI), Cu(II), Ni(II). The mixture of o,m,p-xylene, methyl-ethyl-ketone (MEK) and N,N-dimethylformamide (DMF) (16 : 2 : 1, v/v), containing HDPDTP was chosen as mobile phase after preliminary experiments. Standard solutions of metal ions were spotted on the plates using Brand micropipettes (5 μ L). The plates were developed on 10 cm distance in unsaturated normal chromatographic chamber. After development the plate were dried for 15 min in a hood and visualized with 0.05% Arsenazo III aqueous solution for U(VI) and 0.1% rubeanic acid in ethanol for Cu(II) and Ni(II). All operations were performed at the room temperature.

Calibration curves for U(VI), Cu(II) and Ni(II) determination were plotted using spots area determined by two methods, for chromatographic system containing HDPDTP in o,m,p-xylene – MEK – DMF mixture as mobile phase.

Unknown concentration samples were collected from Roşia Montana (Alba), Baia Mare (Maramureş) and Ştei (Bihor) mining exploitations.

RESULTS AND DISCUSSION

The study of the chromatographic behavior of U(VI), Cu(II) and Ni(II) on silica gel H using HDPDTP as complexing agent in organic mobile phase show that R_F values depend on the HDPDTP concentration. From Figure 1 it can be observed the

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increasing of the retention factor until almost constant value with the increase of complexing agent concentration especially for U(VI).



Figure 1. The influence of the HDPDTP concentration in mobile phase upon the R_F values of the metal ions studied

It can be seen also that the separation of U(VI) from Cu(II) – Ni(II) and the separation of Cu(II) – Ni(II) from each other can be achieved in all range of concentration investigated, 0.02 - 0.07M HDPDTP. The best separation with well defined zones is obtained with 0.05M HDPDTP in a mixture of o,m,p-xylene – MEK - DMF (16 : 2 : 1, v/v). In consequence this mobile phase system was used for TLC quantitative determination of U(VI), Cu(II) and Ni(II).

The calibration curve for Cu(II) and Ni(II) determination were obtained by two methods: a) densitometric determination using DESAGA CD 60 densitometer at 505 nm for Cu(II) and 600 nm for Ni(II), and b) scanning TLC plates with a classic scanner linked to a computer. UTHSCSA Image Tool PC based software was used for area determination. Detection limits were determined using the software Statistical Method in Analytical Chemistry [31]. In both cases the shape of the curve is similar.

Figures 2a and 2b show the calibration curves for Cu(II) and Ni(II), respectively.

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Good linearity is obtained in the concentration range $0.20 - 1.28 \ \mu g/\mu L$ for Cu(II) and $0.3 - 1.2 \ \mu g/\mu L$ for Ni(II). The statistic parameters a (intercept), b (slope of the regression lines, y = ax + b), and r (regression factor) are included. The regression factor, detection limit (LD) and quantitative limit (QL) are presented in Table 1.

Regression factor, DL and QL values for Cu(II) and Ni(II)

Table 1

Cation	Method	r	LD	LQ
			(µg/µL)	(µg/µL)
	Photodensitometry	0.9931	0.132	0.244
Cu(II)	UTHSCSA Image Tool software	0.9872	0.176	0.322
	Photodensitometry	0.9864	0.170	0.310
Ni(II)	UTHSCSA Image Tool software	0.9929	0.118	0.215

The calibration curve for U(VI) was obtained only by scanning TLC plates with classic scanner, because of the strong signal noise. Figure 3 show that good linearity is obtained in the concentration range $0.24 - 24 \ \mu g \ / \mu L$ and r is 0.9973. DL and QL determined by SMAC software are 1.102 and 2.130 respectively.

Calibration curves (Fig. 1-3) were used for the determination of studied metal ions from mining exploitation waters and the results obtained are presented in Table 2. Cu(II), Ni(II) and U(VI) were also determined by other methods: Flame Atomic Absorbtion Spectrometry (FAAS) (Cu(II), Ni(II)) and spectrophotometric method (U(VI)) [32-34].

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Figure 3. Calibration curve for U(VI); Stationary phase: silica gel H; mobile phase: o,m,p-xylene – MEC – DMF (16 : 2 : 1, v/v) with HDPDTP 0.05 M

Table 2

Analysis of Cu(II). Ni(II) and U(VI) content from water samples

Observed	Sample origin	Method/Found concentration (mg / L)		
cation		FAAS	TLC	e _r (%)
Cu(II)	Ore processing waste water	1.50	1.49	-0.67
Ni(II)	(Baia Mare)	2.09	2.12	+1.44
	Natural running mine water (Băita)	Spectrophot	TLC	e _r (%)
0(01)		1.50	1.48	-1.33

It can be seen a good agreement between TLC data and those obtained by FAAS and spectrophotometric method. Determination by TLC analysis requires preconcentration of the sample, but TLC method has some advantages: short time of analysis, inexpensive and easy getting equipment, rapid qualitative results with no necessary treatment of the sample.

CONCLUSIONS

It was found that U(VI), Cu(II) and Ni(II) can be separated by TLC using HDPDTP as complexing agent in the mobile phase, a mixture of organic solvents: o,m,p-xylene – MEK – DMF (16 : 2 :1, v/v). The mobile phase containing 0.05 M HDPDTP was chosen for TLC quantitative determination of the studied metal ions. Calibration curves plotted for Cu(II), Ni(II) and U(VI) were used for the analysis of unknown samples collected from waste water (Cu(II), Ni(II)) and mine water (U(VI)). The results obtained by TLC method are in a good agreement with those obtained by FAAS and spectrophotometric method.

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Dedicated to professor Gh. Marcu at his 80th anniversary

A QUANTUM CHEMICAL CONFORMATIONAL ANALYSIS OF P-TERT-BUTYL/PENTYL/OCTYL-CALIX[8]ARENES

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ABSTRACT. A conformational analysis of *p*-tert-butyl/tert-pentyl/ tert-octyl-calix[8]arenes has been performed by semiempirical calculations. The results are in agreement with the available experimental data, as predicted for the pleated-loop conformation. The relative stability of the conformers depends on the substituents located in the *para* position, however the energy difference between the most stable and the next conformations is low.

Keywords: *p-tert*-butyl-calix[8]arene, *p-tert*-pentyl-calix[8]arene, *p-tert*-octyl-calix[8]arene, conformations, PM3 semiempiric.

Introduction

Calixarenes are [1n]-metacyclophanes, obtained from the condensation of formaldehyde with *p*-alkylphenols, thus resulting structures in which phenol fragments are connected by methylene bridges¹. They have received a special interest as their calix crater is suitable for accommodating guest molecules.

As the calix[8]arenes have many torsion possibilities they are much more flexible than smaller-cycle representatives: there are sixteen up-down conformations plus others in which phenolic fragments adopt an out arrangement².

The goal of this research is to identify and characterize the most stable *para*-substituted calix[8]arene conformations by means of semiempirical methods.

Computational procedure

The starting structures have been built using the graphical interface of Spartan 04³ and preoptimized by molecular mechanics. Optimizations were continued at the semiempirical level employing the PM3 model of Stewart^{4,5} using Gaussian 98 package of programs⁶. Frequency analysis followed all optimizations in order to establish the nature of the stationary points found, so that all the structures reported in this study are genuine minima on the potential energy surface at this level of theory, without any imaginary frequencies. Only the lowest energy conformers (within 15 kcal/mol) have been retained for discussion.

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Results and discussion

The solid state structure of *p*-tert-butyl-calix[8]arene, determined by X-ray diffractometry^{7.8} consists of four pleated fragments joined together. Indeed the global minimum structure found for the *p*-tert-butyl-calix[8]arene is the conformer having this pleated-loop shape, of an average D_{4d} symmetry, an arrangement that favors an internal circular hydrogen bonding stabilizing this conformation⁹ (figure 1). The calculated average O·O distance of 2.74 Å is very close to the 2.71 Å average determined by Gutsche for an empty *p*-tert-butyl-calix[8]arene and the 2.70 Å average reported by Schatz et.al.¹⁰ for the calix:chloroform 1:8 clathrate, thus proving the strong hydrogen bonds maintaining this conformation. The computed value of interplanar angle between two adjacent phenolic units is 111.6° comparable with the ~112° average reported for the chloroform clathrate.



Figure 1. The internal circular hydrogen bonding in the calix[8]arene pleated loop conformer.

The next structure (*tert*-butyl-calix[8] **2** in figure 2) only 0.8 kcal/mole higher in energy may be described as consisting of two trimeric craters with the remaining two phenolic units ensuring a good linking through a "pleated motif" the result being again a stabilising internal circular hydrogen bonding.

The following two structures (*tert*-butyl-calix[8] **3** and **4**) consist of two calix[4] units joined together in an anti C₂ arrangement. Both calix[4] fragments adopt a cone conformation, which has already been proved to be the most stable one among the four possible conformations for a calix[4]arene^{11,12}. Conformations **1**, **2**, **3** and **4** were identified in the case of rare earth (Eu and Lu) dinuclear calix[8]arenes complexes¹³.

Conformer no. **5** at +4.18 kcal/mole above the global minimum is also based on two calix[4]arene fragments but this time having the same orientation although with a certain degree of crater distortion as the two connecting phenols have a parallel orientation. This conformer has been identified in the crystal structure of a tungsten complex trapping a WO group inside each of the calix[4] units¹⁴. This structure is also related to conformer no. **7**, the so-called 'pinched' conformer¹⁵, whose calix[4] cones are more circular due to the lower distance between the connecting groups, but this definitely makes it less stable so its energy is higher than that of the other one.

This conformation was also found experimentally in the case of oligonuclear europium(III) derivatives of *p-tert*-butyl-calix[8]arene crystallized with DMSO^{16,17}.

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The presence of pseudoconic calix[4] units should not be surprising: researchers noticed the remarkable similarity between the temperature dependent ¹H-NMR spectra of *p-tert*-butyl-calix[8]arene and *p-tert*-butyl calix[4]arene¹⁵.

Tert-butyl-calix[8] **6** originates from the lowest energy pleated loop conformer but in this case one of the pleatings has an anti orientation as compared to all the other ones leading to a non-planar structure which makes hydrogen bonding more difficult although the circular shape is well-maintained.

Tert-butyl-calix[8] **8** is also based on two calix conic units joined together, but this time a calix[3] with a calix[5]. The calix[3] unit is slightly tensioned making this conformer less stable but the general orientation of the hydroxyl groups at the lower rim is almost circular, stabilizing this conformation.

Structure *tert*-butyl-calix[8] **9** may be depicted as a distorted two calix[4] structure with the units' connecting fragments adopting an almost out orientation, while the last remaining structure *tert*-butyl-calix[8] **10** may be described as reunion of a large opened calix[4]arene unit connected to a pleated motif by two phenolic fragments having an out orientation.

Interestingly, the global minimum conformer for the *tert*-pentyl-calix[8]arene is the one based on two calix[4]arene cone units joined together in an anti orientation and not the pleated loop conformer found for the *tert*-butyl-calix[8]arene. This arrangement provides enough space for the bulky upper rim substituents and also preserves the stabilizing hydrogen bonding at the lower rim. It is related to *tert*-pentyl-calix[8] **4** which has a similar conformation but with larger calix cavities.

The pleated loop conformer is the next structure at 4.3 kcal/mole above the global minimum, and at the same energy a structure based on two calix[3] units plus a pleated connection fragment is located. All the other structures found for *tert*-pentyl-calix[8]arene have practically the same geometries as the conformers already reported for the *tert*-butyl species respecting the same stability order.

For each class of compounds the corresponding structures are ordered upon their stability, with the corresponding relative energy towards the global minimum conformer indicated (in kcal/mole).

The lowest energy conformer found for the *tert*-octyl-calix[8]arene is the one based on two calix[3] fragments and a V-shaped cleft. Although it shouldn't be so stable from the calix crater point of view, watching the structure not from above but from the calix "plane" side, offers the image of a half-circle geometry, this arrangement providing the necessary space for these large substituents.

The following structure is based on two calix[4] *anti*-oriented units with the calix craters wide opened so that an internal hydrogen bonding can be established between these two units. The space requirement for the bulky octyl fragments is also satisfied, leading to a conformation of only 0.8 kcal/mole above the global minimum structure.

Other two conformers (*tert*-octyl-calix[8] **3** and **7** at 3.0 and 6.8 kcal/mole, respectively) are also based on calix[4] fragments but as they are not so opened the upper rim ligands are more sterically hindered so that these conformers are less stable. The pleated loop conformer is only the sixth one in terms of energy. This is due to the fact that the pleated structure although stabilized in the interior, can not easily accommodate voluminous ligands at the external V-shaped cleft area.

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tert-butyl-calix[8] 4 (+3.3)





tert-butyl-calix[8] 8 (+6.9)



tert-butyl-calix[8] 10 (+10.5)



tert-butyl-calix[8] 1 (0.0)



tert-butyl-calix[8] 3 (+2.3)



tert-butyl-calix[8] 5 (+4.2)



tert-butyl-calix[8] 7 (+6.0)



Figure 2. The ten lowest energy conformers found for *p-tert*-butyl-calix[8]arene.

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tert-pentyl-calix[8] 2 (+4.3)



tert-pentyl-calix[8] 4 (+5.8)



tert-pentyl-calix[8] 6 (+7.3)



tert-pentyl-calix[8] 8 (+10.8)



tert-pentyl-calix[8] 10 (+15.2)

Figure 3. The ten lowest energy conformers found for *p-tert*-pentyl-calix[8]arene.

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tert-octyl-calix[8] 9 (+7.8)

tert-octyl-calix[8] 10 (+10.5)

Figure 4. The ten lowest energy conformers found for *p*-tert-octyl-calix[8]arene.

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Conclusions

Thirty conformers of *p-tert*-butyl/pentyl/octyl-calix[8]arenes have been investigated by PM3 semiempirical method. Their stability is influenced by the hydrogen bonding established at the lower rim and also depends on the volume of the para substituents: thus the most stable geometry for the *p-tert*-butyl-calix[8]arene corresponds to a pleated-loop shape, as exepected from the experimental data. For the *p-tert*-pentyl analogue however the lowest energy conformer is the one consisting of two calix[4] pseudoconic units with an opposed orientation. Their stability is in agreement with the NMR data, as the calix[8]arenes' spectra are similar to those of calix[4]arenes. The most stable conformer for the *p-tert*-octyl-calix[8]arene is the one composed of three fragments including two calix[3] units, an arrangement providing the necessary space for the bulky *tert*-octyl substituents.

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Dedicated to professor Gh. Marcu at his 80th anniversary

SYNTHESIS OF NEW BROMO-STANNANES: TOWARD UNSATURATED TIN DERIVATIVES

PETRONELA MARIA PETRAR, GABRIELA NEMEŞ, IOAN SILAGHI-DUMITRESCU^{*}, LUMINITA SILAGHI-DUMITRESCU

ABSTRACT. Two novel organometallic tin derivatives, bis-dibromomethyl-bis-(2,4,6-tri*iso*propyl-phenyl)-stannane **1** and tribromomethyl--bis-(2,4,6tri*iso*propyl-phenyl)-bromo-stannane **2** have been obtained by the reaction of Tip₂SnF₂ (Tip = 2,4,6-*tri-i*Pr-C₆H₂) with bromoform in the presence of *n*BuLi. The compounds were characterized through NMR spectroscopy and the solid state structure of **2** was determined by X-ray diffraction. Compound **2** is a potential precursor of novel stannaalkenes and 1,3-distannaallenes.

Keywords: stannaalkenes, stannaallenes, double bonded derivatives.

Introduction

Chemists have always shown a big interest in the similarity of carbon with other elements from the same group. The quest for analogue derivatives of alkenes, allenes and even cumulenes containing heavier group 14 elements like silicon, germanium or tin has resulted in an increasing number of publications over the last decades [1]. Alkene derivatives of the type >E=C< (E = heavy group 14 elements) are known [2] and in the last five years, heteronuclear unsaturated compounds of the type >E=E'< have been obtained [3].

Stannaalkenes seem to be more difficult to synthesize than their Si and Ge analogues, as shown by the number of such derivatives known to date: five silenes, eight germenes and only four stannenes [2a] are described in the literature. Allenic compounds >Sn=C=C< are also unknown. Only one tin derivative containing cumulated double bonds, a stannaketenimine, has been reported [4].

In this paper, we describe the synthesis and characterization of two potential precursors of stanna-alkenes and -allenes, starting from a sterrically hindered organometallic tin derivative, Tip_2SnF_2 .

For the synthesis of double bonded tin derivatives, the choice of the starting material is important, because of the protective role played by the organic groups on the tin atom in the Sn=C bond. Bulky radicals, like Tip $(2,4,6-iPr-C_6H_2)$, afford sterrically stabilization of the very reactive tin-carbon bond. The efficiency of Tip as a protection group has already been proved [5]. Scheme 1 summarizes a possible synthetic route which might lead to stanaallenes and stannaalkenes, starting from Tip₂SnF₂:

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Scheme 1

The reaction of REX₂ derivatives (E = group 15 elements) with chloroform or bromoform in the presence of lithium derivatives always leads to the formation of RE=CX₂ compounds [6]. The reaction of supermesithyldichlorophosphane Mes*PCl₂ (Mes* = 2,4,6-*t*Bu-C₄H₂) with haloforms results in the formation of phosphaalkenes Mes*P=CX₂, by means of a lithium carbenoid. Replacement of a halogen atom is possible in the case of bromine and iodine, as shown in Scheme 2.

Mes*PCl₂ + HCX₃ + 2
$$n$$
BuLi
Mes*P=CX₂ X = Cl, Br
Mes*P=CHX X = Br, I

Scheme 2

Similar behavior was recorded for Mes*AsF₂ [7].

It was therefore interesting to explore the reactivity of halostannanes towards bromoform, in the hope of obtaining the corresponding stanaalkenes RR'Sn=CBr₂. However, the behavior of dihalosubstituted group 14 derivatives in such reactions is quite different from that of similar group 15 compounds. Thus, when Tip₂SnF₂ was reacted with one equivalent of CHBr₃ followed by 2 equivalents of *n*BuLi, several unidentified products were obtained with a very poor yield. In order to improve the yield of the formation of the lithium carbenoids CLiHBr₂ and CLiBr₃, the reaction of the dihalostannane with two equivalents of bromoform was attempted.

Results and discussion

The reaction of Tip_2SnF_2 with two equivalents of CHBr₂ and *n*BuLi was performed at -90°C, in THF and leads to the formation of two main compounds **1** and **2** in a 4:1 ratio (Scheme 3).

$$Tip_2SnF_2 + 2 nBuLi + 2 CHBr_3 \xrightarrow{THF} Tip_2Sn \begin{pmatrix} CHBr_2 \\ CHBr_2 \end{pmatrix} + \begin{array}{c} Tip_2Sn - CBr_3 \\ I \\ Br \\ Br \\ 2 \\ \end{array}$$

Scheme 3

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As reported in the literature [6], both $CLiHBr_2$ and $CLiBr_3$ may be formed by the reaction of lithium derivatives with bromoform. The action of the $CLiHBr_2$ carbenoid on Tip_2SnF_2 leads to compound **1**, while $Tip_2Sn(F)CBr_3$ is formed with $CLiBr_3$. A fluorine-bromine exchange is supposed to take place between $Tip_2Sn(F)CBr_3$ and the LiBr formed *in situ*, leading to compound **2**.

Compounds 1 and 2 have been characterized by multinuclear NMR spectroscopy. The relevant NMR data for 1 are given in Table 1.

NMR data for Tip₂Sn(CHBr)₂

Table 1

¹¹⁹Sn NMR ¹³C NMR ¹H NMR 1.50 ppm (d, ³J_{HH} = 6.4 Hz, 23.9 ppm (CH3, orto-iPr) -134.2 ppm 24H, C<u>H</u>₃, orto-*i*Pr) $(^{2}J_{SnBr} = 11.8 \text{ Hz})$ 24.7 ppm ($\overline{C}H_3$, para-*i*Pr) 1.18 ppm (d, ${}^{3}J_{HH} = 6.4$ Hz, 32.5 ppm (CH orto-*i*Pr) 12H, CH_{3} , para- *i*Pr) 2.58 (m, ³J_{HH} = 6.4Hz, 4H, CH, orto-*i*Pr) 2.82 (m, ³J_{HH} = 6.4 Hz, 2H, 34.3 ppm (CH, para-IPr) 40.1 ppm (CHBr₂) 122.6 (meta-<u>C</u>H Tip), 150.9 and 151.6 ppm (orto CH, para-iPr) and para-C Tip). 5.80 (s, 2H, CHBr₂)

The ¹¹⁹Sn NMR spectrum of compound **1** shows the multiplet resulted from the coupling with the four bromine atoms (Figure 1). Such signals are usually broad and the coupling constant is not observed; to our best knowledge, it is the first time that coupling of tin with such an important number of bromine atoms resulted in an observed hyperfine splitting of the signal and not just in its widening.

6.96 (s, 4H, meta-Ć<u>H</u> Tip).



Figure 1. ¹¹⁹Sn NMR signal for Tip₂Sn(CHBr₂)₂

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The ¹H-NMR chemical shifts for compound **2** are within the expected range; the position of all signals is given in the Experimental section. Two different signals were recorded for the two methyl groups in the *o-iso*propyl groups (1.050 ppm, 1.080 ppm respectively). This indicates that the free rotation around the Sn-C bond is significantly hindered, no doubt due to the presence of the sterricaly challenging groups on the tin atom.

Compound **2**, Tip₂Sn(Br)CBr₃, was characterized in solid state through Xray diffraction on single crystal. An ORTEP rendering of the molecular structure is shown in Figure 2. Some geometrical data are also given in Table 2. The presence of the heavy bromine atoms makes the refinement of the structure more difficult. Two equivalent positions atom are possible for the CBr₃ groups, as well as for the Br1 atom, with respect to an axis containing the central tin and acting as a two-fold symmetry axis for the Tip groups. The Sn-Br distance (2.714 Å) is remarkably larger than those found in similar compounds (2.50 - 2.60 Å) [8, 9]. As expected, the geometry about the tin atom is distorted tetrahedral, with wider angles with atoms belonging to the CBr₃ group.



Figure 2. Molecular structure of Tip₂Sn(Br)CBr₃ (hydrogen atoms were omitted for clarity)

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Table 2

Geometrical data for Tip₂Sn(Br)CBr₃

	Bond lengths (Å)	Bond a	ngles (°)
Sn-C1	2.227	Br1-Sn-C1	94.19
Sn-C2	2.156	C2-Sn-C3	110.81
Sn-Br1	2.714	C1-Sn-C3	122.70
C1-Br2	1.932	Br2-C1-Sn	104.52
C1-Br3	1.940	Br1-C1-Br2	114.98

Conclusions and perspectives

Two novel organometallic tin derivatives have been obtained. Their structure in solution was elucidated by NMR spectroscopy. The solid-state structure of $Tip_2Sn(Br)CBr_3$ was also determined.

Even though the reaction of Tip_2SnF_2 with bromoform did not lead to the expected stannaalkene, it afforded a potential precursor of double bonded derivatives of tin. Scheme 4 shows the possible use of $Tip_2Sn(Br)CBr_3$ in the synthesis of stannaalkenes and 1,3-distannaallenes.



Scheme 4

The distaannallene $Tip_2Sn=C=SnTip_2$ would be the first compound of this type obtained to date. The proposed synthetic routes are currently under experimental investigation.

Acknowledgment. Dr. Jean Escudie from Paul Sabatier University Toulouse, is thanked for his hospitality and for useful discussion on this manuscript, as well as Dr. Heinz Gornitzka for collecting the X-Ray diffraction data. We thank the Ministry of Education and Research Bucharest (CEEX project) and CNCSIS Agency for partial financial support of this work.

Experimental procedure

All manipulations were carried out under N₂ or Ar using standard Schlenk techniques with solvents freshly distilled from sodium benzophenone. Mass spectra were measured on a Hewlett-Packard 5989 A spectrometer by El at 70 eV. Melting points were determined on a Leitz microscope heating stage 250. NMR Spectra were recorded on a Bruckner Varian 200 MHz for ¹H and ³¹P and 300 MHz for ¹³C and ¹¹⁹Sn nucleus. CDCl₃ was used as a solvent.

Crystal Data for **2** were collected at room temperatures using an oil-coated shock-cooled crystal on a Bruker-AXS CCD 1000 diffractometer with MoK α radiation (λ =

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0.71073 Å). The structure was solved by direct methods (SHELXS-97) [10] and all non hydrogen atoms were refined anisotropically using the least-squares method on F^2 [11].

Synthesis of bis-dibromomethyl-bis-(2,4,6-triisopropyl-phenyl)-stannane **1** and tribromomethyl--bis-(2,4,6-triisopropyl-phenyl)-bromo-stannane **2**:

A mixture of 2 g (0.0035 mols) Tip₂SnF₂ and 1.77 g CHBr₃ (0.007 mols) in 70 ml THF was cooled at -90°C and 4.5 ml *n*BuLi 1.6 M (0.0072 mols) were added dropwise. The reaction mixture was stirred for half an hour at -90°C and then it was allowed to warm up at room temperature. The solvent was removed under vacuum and replaced with 20 ml pentane. Lithium salts were removed by filtration and the filtrate was stored at -25°C. After 24 hours, compound **1** was isolated by filtration as a white powder with a 45% yield. The remaining solution was further concentrated under vacuum and stored at -25°C. Transparent crystals of **2** were isolated after a few hours with a 11% yield.

1: NMR data are given in Table 1;

MS (Z/e): 697 (M-CHBr₂), m.p. = 177°C

2: ¹H-NMR: 1.050 ppm (d, ³J_{HH}=6.2 Hz, 6H, C<u>H₃</u>, orto-*i*Pr); 1.080 ppm (d, ³J_{HH}=5.8 Hz, 6H, C<u>H₃</u>, orto-*i*Pr); 1.194 ppm (d, ³J_{HH}=7 Hz, 6H, C<u>H₃</u>, para-*i*Pr); 2.567 ppm (m, 1H, C<u>H</u>, para-*i*Pr); 3.35 ppm (m, 2H, C<u>H</u>, orto-*i*Pr), 7.080 ppm (s, 4H, meta-C<u>H</u> Tip).

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NEW ORGANOTIN(IV) AND – LEAD(IV) *N,N*-DIMETHYLDITHIOCARBAMATES: SYNTHESIS, SOLUTION NMR CHARACTERIZATION AND SINGLE-CRYSTAL X- RAY STRUCTURE OF Ph₂MCI(S₂CNMe₂) (M = Sn, Pb) AND Ph₂Sn(S₂CNMe₂)₂

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ABSTRACT. New *N*,*N*-dimethyldithiocarbamato derivatives of organotin(IV) and – lead(IV), R₂MCI(S₂CNMe₂) (R = Bu, M = Sn; R = Ph, M = Sn, Pb), R₂M(S₂CNMe₂)₂ (R = Bu, Ph, M = Sn) and R₃M(S₂CNMe₂) (R = Ph, M = Sn, Pb; R = Me, M = Pb), were prepared and characterized by multinuclear (¹H, ¹³C, 2D) NMR spectroscopy in solution. The molecular structures of Ph₂SnCl(S₂CNMe₂) (**3**), Ph₂Sn(S₂CNMe₂)₂ (**4**) and Ph₂PbCl(S₂CNMe₂) (**6**) were established by single-crystal X-ray diffraction. In all cases the dithio ligand acts as an asymmetric monometallic biconnective moiety. The mixed chloro-dithiocarbamato derivatives **3** and **6** exhibit distorted trigonal bipyramidal C₂MCIS₂ (**4**) a distorted octahedral environment is achieved, with *cis* organic groups attached to tin atom.

Key-words: organotin, organolead, dithiocarbamato ligands, solution NMR studies, X-ray diffraction

INTRODUCTION

Main group dithiocarbamato complexes find various applications in materials and separation science, and have potential use as chemotherapeutics, pesticides, and fungicides. In addition, the dithiocarbamato moiety is a highly versatile ligand towards main group metals, they can stabilize a variety of oxidation states and coordination geometries and small modifications to the ligand can often lead to significant changes in the structure of the complexes formed. The literature on the structural aspects of tin and lead dithiocarbamates was recently reviewed [1], and a search of the Cambridge Structure Database revealed many gaps concerning the structure investigations of organotin(IV) dimethyldithiocarbamates, while no structure of an organolead(IV) analog was so far established by single-crystal X-ray diffractometry.

We report here on the synthesis and solution behavior of several diorganoand triorganometal(IV) (M = Sn, Pb) dimethyldithiocarbamates, the molecular structures of $Ph_2SnCl_x(S_2CNMe_2)_{2x}$ (x = 0, 1) as well as the first molecular structure of a diorganolead(IV) dithiocarbamate, *i.e.* $Ph_2PbCl(S_2CNMe_2)$.

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EXPERIMENTAL

The starting materials, *i.e.* organotin(IV)- and organolead(IV)- chlorides and $Me_2NCS_2Na\cdot 2H_2O$, were commercially available (Aldrich) and were used without further purification. Room-temperature ¹H and ¹³C NMR spectra, including 2D experiments, were recorded in dried CDCl₃ on a BRUKER AVANCE DRX 400 instrument operating at 400.16 and 100.62 MHz, respectively. The chemical shifts are reported in ppm relative to the residual peak of solvent (ref. CDCl₃: ¹H 7.26 ppm, ¹³C 77.0 ppm).

Preparation of $Bu_2SnCl(S_2CNMe_2)$ (1)

Me₂NCS₂Na·2H₂O (0.460 g, 2.567 mmol) was added to a solution of Bu₂SnCl₂ (0.782 g, 2.574 mmol) in CH₂Cl₂. A white precipitate of NaCl formed in short time. After removing the NaCl, the solvent was evaporated and the title compound was isolated as viscous yellowish oil. ¹H NMR: δ 0.93t [6H, CH₃-(CH₂)₃-Sn, ³J_{HH} 5.8 Hz], 1.41m,br [4H, CH₃-CH₂-(CH₂)₂-Sn], 1.83m,br [8H, CH₃-CH₂-(CH₂)₂-Sn], 3.40s (6H, CH₃-N). ¹³C NMR: δ 13.58s (C_δ), 26.21s (C_γ, ³J_{SnC} 98 Hz), 27.75s (C_β, ²J_{SnC} 34 Hz), 29.08s (C_α, ¹J_{SnC} 501 / 522 Hz), 45.10s (CH₃-N), 198.04 (N-CS₂).

Preparation of $Bu_2Sn(S_2CNMe_2)_2$ (2)

Me₂NCS₂Na·2H₂O (0.922 g, 5.146 mmol) was added to a solution of Bu₂SnCl₂ (0.782 g, 2.574 mmol) in CH₂Cl₂. A white precipitate of NaCl formed in short time. After removing the NaCl, the solvent was evaporated and the title compound was isolated as yellowish solid. Yield: 0.95 g (78%), m.p. 137-140°C. ¹H NMR: δ 0.91t [6H, CH₃-(CH₂)₃-Sn, ³J_{HH} 7.3 Hz], 1.41tq [4H, CH₃-CH₂-(CH₂)₂-Sn, ³J_{HH} 7.2 Hz], 1.88m [4H, CH₃-CH₂-CH₂-CH₂-Sn], 2.01m [4H, CH₃-(CH₂)₂-CH₂-Sn], 3.42s (12H, CH₃-N). ¹³C NMR: δ 13.70s (C_δ), 26.36s (C_γ, ³J_{SnC} 124 Hz), 28.46s (C_β), 34.18s (C_α, ¹J_{SnC} 576 / 598 Hz), 44.58s (CH₃-N), 200.93 (N-CS₂).

The following compounds were prepared similarly:

 $\begin{array}{l} Ph_2SnCl(S_2CNMe_2) \quad (\textbf{3}), \mbox{ from } Ph_2SnCl_2 \quad (1.605 \mbox{ g}, \mbox{ 4.668 mmol}) \mbox{ and } \\ Me_2NCS_2Na\cdot 2H_2O \quad (0.837 \mbox{ g}, \mbox{ 4.668 mmol}) \mbox{ in } CH_2Cl_2. \mbox{ Yield: } 1.30 \mbox{ g} \mbox{ (65\%)}, \mbox{ m.p. } 240-241^{\circ}C. \mbox{ }^1H \mbox{ NMR: } \delta \mbox{ 3.40s} \mbox{ (6H, } CH_3-N), \mbox{ 7.47m} \mbox{ (6H, } C_6H_5-meta+para), \mbox{ 8.07d} \mbox{ (4H, } C_6H_5-ortho, \mbox{ }^3J_{HH} \mbox{ 7.8, } ^3J_{SnH} \mbox{ 88 Hz}). \mbox{ }^{13}C \mbox{ NMR: } \delta \mbox{ 45.83s} \mbox{ (CH}_3-N), \mbox{ 128.77s} \mbox{ (C}_m, \mbox{ }^3J_{SnC} \mbox{ 88 Hz}), \mbox{ 130.16s} \mbox{ (C}_p, \mbox{ }^4J_{Sn-C} \mbox{ 18 Hz}), \mbox{ 135.66s} \mbox{ (C}_o, \mbox{ }^2J_{SnC} \mbox{ 64 Hz}), \mbox{ 141.85s} \mbox{ (C}_i), \mbox{ 196.81} \mbox{ (N-CS}_2). \end{array}$

 $Ph_2Sn(S_2CNMe_2)_2$ (4), from Ph_2SnCl_2 (1.605 g, 4.668 mmol) and $Me_2NCS_2Na\cdot 2H_2O$ (1.673 g, 9.336 mmol) in CH_2Cl_2 . Yield: 1.68 g (70%), m.p. 160-165°C. ¹H NMR: δ 3.30s (12H, CH_3 -N), 7.42m (6H, C_6H_5 -meta+para), 8.06m (4H, C_6H_5 -ortho, ³J_{SnH} 88 Hz). ¹³C NMR: δ 45.80s (CH_3 -N), 128.75s (C_m , ³J_{SnC} 88 Hz), 130.15s (C_p), 135.64s (C_0 , ²J_{SnC} 63 Hz), 141.83s (C_i), 196.75 (N-CS₂).

*Ph*₃*Sn*(*S*₂*CNMe*₂) (**5**), from Ph₃*SnCl* (0.82 g, 2.127 mmol) and Me₂NCS₂Na·2H₂O (0.38 g, 2.120 mmol) in CH₂Cl₂. Yield: 0.31 g (31%), m.p. 275-280°C. ¹H NMR: δ 3.46s (6H, *CH*₃-N), 7.42m (9H, *C*₆*H*₅-*meta*+*para*), 7.76m (6H, *C*₆*H*₅-*ortho*, ³J_{SnH} 61 Hz). ¹³C NMR: δ 45.95s (CH₃-N), 128.47s (*C*_m, ³J_{SnC} 61 Hz), 129.08s (*C*_p, ⁴J_{Sn-C} 13 Hz), 136.67s (*C*_o, ²J_{SnC} 46 Hz), 142.16s (*C*_i, ¹J_{SnC} 579 / 606 Hz), 196.76 (N-CS₂).

*Ph*₂*PbCl*(S_2CNMe_2) (**6**), from Ph₂PbCl₂ (1.561 g, 3.61 mmol) and Me₂NCS₂Na·2H₂O (0.647 g, 3.61 mmol) in CH₂Cl₂. Yield: 1.57 g (84%), m.p. 190-192°C. ¹H NMR: δ 3.35s (6H, CH₃-N), 7.41t (2H, C₆H₅-para, ³J_{HH} 7.4 Hz), 7.56dd (4H, C₆H₅-meta, ³J_{HH} 7.6 Hz), 8.26d (4H, C₆H₅-ortho, ³J_{HH} 7.9, ³J_{PbH} 175 Hz). ¹³C NMR: δ 45.29s (CH₃-N), 129.93s (C_p), 130.28s (C_m, ³J_{PbC} 43 Hz), 134.83s (C_o, ²J_{PbC} 115 Hz), 137.92s (C_i), 200.87 (N-CS₂).

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*Ph*₃*Pb*(*S*₂*CNMe*₂) (**7**), from Ph₃*PbCl* (0.793 g, 1.674 mmol) and Me₂NCS₂Na·2H₂O (0.30 g, 1.674 mmol) in CH₂Cl₂. Yield: 0.53 g (57%), m.p. 199-200°C. ¹H NMR: δ 3.48s (6H, *CH*₃-N), 7.38t (3H, *C*₆*H*₅-*para*, ³*J*_{HH} 7.4 Hz), 7.50dd (6H, *C*₆*H*₅-*meta*, ³*J*_{HH} 7.4 Hz), 7.86d (4H, *C*₆*H*₅-*ortho*, ³*J*_{HH} 6.8, ³*J*_{PbH} 105 Hz). ¹³C NMR: δ 46.21s (*CH*₃-N), 128.96s (*C*_p, ⁴*J*_{PbC} 22 Hz), 129.79s (*C*_m, ³*J*_{PbC} 100 Hz), 136.89s (*C*_o, ²*J*_{PbC} 83 Hz), 158.18s (*C*_i, ¹*J*_{PbC} 563 Hz), 190.14 (N-CS₂).

 $Me_3Pb(S_2CNMe_2)$ (**8**), from Me_3PbCI (0.70 g, 2.433 mmol) and $Me_2NCS_2Na\cdot 2H_2O$ (0.438 g, 2.444 mmol) in CH_2CI_2 . Yield: 0.54 g (60%), m.p. 235-237°C. ¹H NMR: δ 1.44s (9H, CH_3 -Pb, ²J_{PbH} 64 Hz), 3.47s (6H, CH_3 -N). ¹³C NMR: δ 13.30s (CH_3 -Pb, ¹J_{PbC} 282 Hz), 45.30s (CH_3 -N), 201.24 (N-CS₂).

X-ray Crystallographic Study

Data were collected with a SMART APEX diffractometer (*National Center* for X-Ray Diffractometry, "Babes-Boyai" University, Cluj-Napoca, Romania) at 297 K. In all cases a graphite monochromator was used to produce a wavelength (Mo- $K\alpha$) of 0.71073 Å. The crystal structure measurement and refinement data for compounds **3**, **4** and **6** are given in Table 1. Absorption corrections were applied using the multi-scan (Bruker SAINT) method [2]. The structures were solved by direct methods (full-matrix least-squares on F²). All non hydrogen atoms were refined with anisotropic thermal parameters. For structure solving and refinement a software package SHELX-97 was used [3]. The drawings were created using the Diamond program by Crystal Impact GbR [4].

CCDC-601481 (3), CCDC-601483 (4) and CCDC-601482 (6) contain the supplementary crystallographic data for this paper. These data can be obtained free of charge at www.ccdc.cam.ac.uk/conts/retrieving.html [or from the Cambridge Crystallographic Data Centre, 12, Union Road, Cambridge CB2 1EZ, UK; fax: (internat.) +44-1223/336-033; E-mail: deposit@ccdc.cam.ac.uk].

Crystallographic data for compounds **3**, **4** and **6**.

	3	4	6			
Empirical formula	C ₁₅ H ₁₆ CINS ₂ Sn	$C_{18}H_{22}N_2S_4Sn$	C ₁₅ H ₁₆ CINPbS ₂			
Formula mass	428.55	513.31	517.05			
Crystal system	Triclinic	Tetragonal	Triclinic			
Spạce group	P-1	l4(1)/a	P-1			
a [Å]	9.1505(13)	16.532(2)	6.68(3)			
b [Å]	9.7273(14)	16.532(2)	9.36(4)			
c [Å]	11.8051(17)	15.533(4)	14.97(6)			
α [°]	79.367(2)	90	79.06(6)			
β [[°]]	67.558(2)	90	81.35(7)			
γ[[°]]	62.500(2)	90	69.98(6)			
V [Å ³]	861.4(2)	4245.1(13)	859(6)			
Z	2	8	2			
D _{calcd.} [g/cm ³]	1.652	1.606	1.999			
F(000)	424	2064	488			
Crystal size [mm]	0.22x0.16x0.16	0.26x0.13x0.10	0.23x0.20x0.15			
$\mu(Mo-K\alpha) [mm^{-1}]$	1.870	1.601	10.209			
$\hat{\theta}$ range \int_{0}^{0}	1.87-26.37	1.80-26.36	2.34-25.35			
No. of reflections collected	9227	16801	8144			
No. of independent	3507	2171	3107			
reflections	(R _{int} = 0.0265)	(R _{int} = 0.0565)	$(R_{int} = 0.0376)$			

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No. of parameters	183	116	183
$R_1 [I > 2\sigma(I)]$	0.0334	0.0560	0.0287
wR ₂	0.0723	0.0635	0.0675
GOF on F^2	1.144	1.271	1.150
Largest difference electron density [e/Å ³]	0.714/-0.784	0.803/0.946	1.006/-1.244

RESULTS AND DISCUSSION

Svnthesis

The new compounds were obtained by reacting R_2MCI_2 and R_3MCI with sodium dimethyldithiocarbamate in 1:1 and 1:2 molar ratios, in CH_2CI_2 , at room temperature, according to the following equations:



Most of the organotin(IV)- and organolead(IV) dithiocarbamates were isolated in good yield as yellowish solids. Compound **1** was obtained as viscous yellowish oil. They are air-stable compounds, soluble in common organic solvents. All compounds were investigated by NMR spectroscopy (¹H, ¹³C, 2D) in CDCl₃ solutions and for compounds **3**, **4** and **6** the molecular structures were established using single-crystal X-ray diffraction.

NMR Spectroscopy

For both the organoSn(IV) and -Pb(IV) complexes the spectra contain characteristic resonance signals for equivalent organic groups bonded to the metal centre as well as for the dimethyldithiocarbamato ligand, thus providing evidences for the identity of the obtained compounds. The ¹H resonances assigned to the *ortho* protons in the phenyl groups attached to the metal and to the methyl protons in the trimethyllead(IV) derivative, as well as most of the ¹³C resonances for the organic groups bound to tin or lead, show satellites due to metal-proton or metal-carbon couplings.

The coordination geometry of organotin(IV) derivatives is reflected in the NMR parameters [5]. Thus, based on the reported dependence on the ${}^{1}J({}^{119}Sn^{13}C)$ coupling constants in butyltin(IV) derivatives [] ${}^{1}J({}^{119}Sn^{13}C)$] = 9.996-746] [6], the estimated values for the C-Sn-C angle (θ) are 126.9° [${}^{1}J({}^{119}Sn^{13}C$ 522 Hz] in 1 and 134.5° [${}^{1}J({}^{119}Sn^{13}C$ 598 Hz] in 2. This suggests a distorted trigonal bipyramidal structure for the chloro(dithiocarbamato)tin(IV) derivative 1 and a skew-trapezoidal bipyramid (or highly distorted octahedron) for the bis(dithiocarbamato)tin(IV) derivative 2. Similar coordination geometries were previously reported for several related organotin(IV) derivatives [1].

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The magnitude of the lead-proton and lead-carbon coupling constants is consistent with the presence of diphenyllead(IV) moiety in compound **6** $[{}^{3}J({}^{207}Pb^{1}H)$ 175 Hz, ${}^{2}J({}^{207}Pb^{13}C)$ 115 Hz] and triorganolead(IV) fragments in compounds **7** [R = Ph, ${}^{3}J({}^{207}Pb^{1}H)$ 105 Hz, ${}^{2}J({}^{207}Pb^{13}C)$ 183 Hz, ${}^{1}J({}^{207}Pb^{13}C)$ 563 Hz] and **8** [R = Me, ${}^{3}J({}^{207}Pb^{1}H)$ 64 Hz, ${}^{1}J({}^{207}Pb^{13}C)$ 282 Hz], respectively [7,8]. The ¹H and ¹³C NMR spectra of compounds **1** – **8** are very similar with respect

The ¹H and ¹³C NMR spectra of compounds **1** – **8** are very similar with respect to the dimethyldithiocarbamato moiety. The presence of sharp singlets for the methyl protons in the ¹H NMR spectra and for the methyl carbons in the ¹³C NMR spectra, respectively, indicates the equivalence of the methyl groups on nitrogen. Taking into account the restricted rotation about the (S₂)C-N partial double bond of the dithiocarbamato group, this behavior is consistent with a fast fluxional behavior on the NMR chemical shift time scale which alternatively brings the two sulfur atoms in the same position of the primary coordination sphere of the metal centre.

Single-crystal X-ray diffraction studies

Single-crystals suitable for X-ray diffraction studies were obtained by slow diffusion from a mixture of chloroform and hexane (1/4, v/v) for compounds $Ph_2SnCl(S_2CNMe_2)$ (3) and $Ph_2PbCl(S_2CNMe_2)$ (6). The crystal of 3 consists of discrete monomers separated by normal van der Waals distances, while in the crystal of 6 dimeric associations might be considered (see subsequent discussion). The ORTEP diagrams of the molecular structures of compounds 3 and 6, with the atom numbering scheme, are shown in Figures 1 and 2. Selected interatomic distances and angles are listed in Table 2.

The molecular structure of compounds **3** and **6** exhibits some common structural features. The dimethyldithiocarbamato moiety behaves as *asymmetric monometallic biconnective units*, with short [Sn(1)-S(1) 2.4718(10) Å in **3**, and Pb(1)-S(1) 2.607(8) Å in **6**] and long [Sn(1)-S(2) 2.6834(10) Å in **3**, and Pb(1)-S(2) 2.821(8) Å in **6**] metal-sulfur distances.

The length of the Sn(1)-S(1) primary bond in **3** is typical for a covalent bond, while the secondary Sn(1)-S(2) bond is stronger than that observed in the related Me₂SnCl(S₂CNMe₂) [Sn-S 2.48(1) Å, Sn···S 2.79(1) Å] [9] or (PhCH₂)₂SnCl(S₂CNMe₂) [Sn-S 2.464(2) Å, Sn···S 2.707(2) Å] [10].



Figure 1. ORTEP representation at 30% probability and atom numbering scheme for Ph₂SnCl(S₂CNMe₂) (3).

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Figure 2. ORTEP representation at 30% probability and atom numbering scheme for $Ph_2PbCl(S_2CNMe_2)$ (6).

Tab	ble	2.
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Interatomic bond distances (Å) and angles (°) for compounds 3 and 6 .							
3	3 6						
Sn(1)-C(4)	2.137(3)	Pb(1)-C(4)	2.195(8)				
Sn(1)-C(10)	2.140(3)	Pb(1)-C(10)	2.232(10)				
Sn(1)-Cl(1)	2.4486(9)	Pb(1)-Cl(1)	2.617(10)				
Sn(1)-S(1)	2.4718(10)	Pb(1)-S(1)	2.607(8)				
Sn(1)-S(2)	2.6834(10)	Pb(1)-S(2)	2.821(8)				
C(1)-S(1)	1.739(4)	C(1)-S(1)	1.754(9)				
C(1)-S(2)	1.713(4)	C(1)-S(2)	1.704(8)				
C(1)-N(1)	1.317(4)	C(1)-N(1)	1.339(10)				
C(2)-N(1)	1.457(5)	C(2)-N(1)	1.466(11)				
C(3)-N(1)	1.460(5)	C(3)-N(1)	1.467(11)				
C(4)-Sn(1)-C(10)	116.93(13)	C(4)-Pb(1)-C(10)	146.5(3)				
S(1)-Sn(1)-C(4)	132.35(10)	S(1)-Pb(1)-C(4)	109.0(3)				
S(1)-Sn(1)-C(10)	109.10(9)	S(1)-Pb(1)-C(10)	103.8(3)				
Cl(1)-Sn(1)-C(4)	96.77(10)	Cl(1)-Pb(1)-C(4)	95.0(3)				
Cl(1)-Sn(1)-C(10)	98.25(10)	Cl(1)-Pb(1)-C(10)	93.09(18)				
Cl(1)-Sn(1)-S(1)	87.99(4)	Cl(1)-Pb(1)-S(1)	87.4(2)				
S(2)-Sn(1)-C(4)	92.91(10)	S(2)-Pb(1)-C(4)	93.7(3)				
S(2)-Sn(1)-C(10)	96.47(10)	S(2)-Pb(1)-C(10)	93.2(2)				
S(2)-Sn(1)-S(1)	69.70(3)	S(2)-Pb(1)-S(1)	66.4(3)				
Cl(1)-Sn(1)-S(2)	156.36(4)	CI(1)-Pb(1)-S(2)	153.84(9)				
Sn(1)-S(1)-C(1)	89.42(12)	Pb(1)-S(1)-C(1)	90.2(3)				
Sn(1)-S(2)-C(1)	83.21(12)	Pb(1)-S(2)-C(1)	84.3(4)				

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Table 2.

(continued).

		(continucu).		
N(1)-C(1)-S(1)	120.4(3)	N(1)-C(1)-S(1)	119.1(5)	
N(1)-C(1)-S(2)	122.1(3)	N(1)-C(1)-S(2)	122.1(6)	
S(1)-C(1)-S(2)	117.5(2)	S(1)-C(1)-S(2)	118.8(5)	
C(1)-N(1)-C(2)	122.8(3)	C(1)-N(1)-C(2)	121.3(7)	
C(1)-N(1)-C(3)	121.5(4)	C(1)-N(1)-C(3)	122.4(7)	
C(2)-N(1)-C(3)	115.7(3)	C(2)-N(1)-C(3)	116.2(7)	

The Pb(1)-S(1) primary bond [2.607(8) Å] in **6** is considerably stronger, while the secondary Pb(1)-S(2) bond [2.821(8) Å] is similar with respect to those found for lead(II) dithiocarbamates e.g. Pb(S₂CNMe₂)₂ [Pb-S 2.779(5) Å; Pb···S 2.873(6) Å] [11], Pb(S₂CNEt₂)₂ [Pb-S 2.744(9), 2.786(9) Å; Pb···S 2.885(11), 2.940(10) Å] [12], or Pb(S₂CNPr₂)₂ [Pb-S 2.673(4), 2.681(4) Å; Pb···S 2.843(5), 2.859(5) Å] [13].

The tin-chlorine distance in **3** [Sn(1)-Cl(1) 2.4486(9) Å] compares well with those observed in Me₂SnCl(S₂CNMe₂) [Sn-Cl 2.465(9) Å] [9] or (PhCH₂)₂SnCl(S₂CNMe₂) [Sn-Cl 2.482(2) Å] [10]. The lead-chlorine bond in **6** [Pb(1)-Cl(1) 2.617(10) Å] is shorter than that observed in the polymeric Ph₂PbCl₂ starting material [Pb-Cl 2.795(1) Å in the Pb-Cl-Pb bridges] [14], but of the same magnitude as found in the related Ph₂PbCl[S(S)CPPh₂(O)] [Pb-Cl 2.62(1) Å] [15], respectively.

The resulting coordination geometry at the metal centre is best described as highly distorted trigonal bipyramidal, with the dithiocarbamato ligand spanning axial–equatorial positions. The long metal-S(2) bond is axial, in *trans* to the chlorine atom [Cl(1)-Sn(1)-S(2) 156.36(4)° in **3**, and Cl(1)-Pb(1)-S(2) 153.84(9)° in **6**]. The equatorial plane is described by the C(4), C(10) and S(1) atoms, with the metal atom displaced from this plane towards the chlorine atom with 0.163 and 0.104 Å in **3** and **6**, respectively. The S(1)Sn(1)S(2) plane is almost perpendicular to the C(4)C(10)S(1) basal plane (dihedral angle 84.1° and 89.3° in **3** and **6**, respectively). The distortion is mainly due to the small bite of the ligand [S(1)...S(2) 2.951(2) and 2.98(1) Å in **3** and **6**, respectively].

The main difference between these two molecular structures is the opened equatorial C(4)-Pb(1)-C(10) angle $[146.5(3)^{\circ}]$ in **6** compared to the corresponding C(4)-Sn(1)-C(10) angle $[116.93(13)^{\circ}]$ in **3**. A closer check of the crystal structure of the lead derivative revealed the presence of a weak intermolecular π -Pb–phenyl interaction [Pb(1)···C_{6.7.8} 3.77(1)-4.23(2) Å] which involves one of the aromatic rings attached to the metal centre. The vector of this π -Pb–phenyl interaction bisects the C(4)-Pb(1)-C(10) angle, in *trans* to the S(1) atom, and might be the cause of enlargement of this equatorial angle. Based on these interactions dimer associations might be considered to be formed, in which the coordination geometry around the lead can be described as distorted octahedral (Figure 3). Similar intra- or intermolecular π -Pb–phenyl interactions were previously described in the monomeric Pb[(SPPh₂)₂N]₂ [Pb(1)···C_{1.26} 3.424-3.579 Å] [17].

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Figure 3. Dimeric association in the crystal of Ph₂PbCl(S₂CNMe₂) (6).

The molecular structure of $Ph_2Sn(S_2CNMe_2)_2$ (4) was also established by single-crystals X-ray diffraction studies [the crystals were obtained from a chloroform and hexane (1/4, v/v) mixture]. The crystal of 4 consists of discrete monomers separated by normal van der Waals distances. The ORTEP diagram of the molecular structure of compound 4, with the atom numbering scheme, is shown in Figure 4. Selected interatomic distances and angles are listed in Table 3.



Figure 4. ORTEP representation at 30% probability and atom numbering scheme for Ph₂Sn(S₂CNMe₂)₂ (4). Table 3.

Interatomic bond distances (Å) and angles (°) for compound 4 .							
Sn(1)-C(4)	2.165(5)	C(4)-Sn(1)-C(4a)	98.2(3)				
Sn(1)-S(1)	2.5860(14)	S(1)-Sn(1)-S(2)	68.34(5)				
Sn(1)-S(2)	2.6632(15)	S(1)-Sn(1)-S(1a)	152.83(7)				
		S(1)-Sn(1)-S(2a)	90.90(5)				
C(1)-S(1)	1.728(5)	S(2)-Sn(1)-S(1a)	90.90(5)				
C(1)-S(2)	1.702(5)	S(2)-Sn(1)-S(2a)	82.45(7)				
C(1)-N(1)	1.329(7)	S(1a)-Sn(1)-S(2a)	68.34(5)				
C(2)-N(1)	1.467(7)						

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C(3)-N(1)	1.450(8)	C(4)-Sn(1)-S(1) C(4)-Sn(1)-S(2)	93.81(14) 161 15(14)
N(1)-C(1)-S(1)	120.2(4)	C(4) Sn(1) S(2) C(4) Sn(1) S(1a) C(4) Sn(1) S(2a)	103.94(14)
S(1)-C(1)-S(2)	118.6(3)	C(4)-SII(1)-S(2a)	92.10(14)
C(1)-N(1)-C(2)	121.4(5)	C(4a)-Sn(1)-S(1) C(4a)-Sn(1)-S(2)	103.94(14) 92.18(14)
C(1)-N(1)-C(3) C(2)-N(1)-C(3)	122.1(5) 116.5(5)	C(4a)-Sn(1)-S(1a) C(4a)-Sn(1)-S(2a)	93.81(14) 161.15(14)
- () () - (-)		Sn(1)-S(1)-C(1)	87 39(18)
		Sn(1)-S(2)-C(1)	85.44(18)

As in compound **3**, both dimethyldithiocarbamato moieties in a molecular unit of **4** act as *asymmetric monometallic biconnective units*, with short [Sn(1)-S(1) 2.5860(14) Å] and long [Sn(1)-S(2) 2.6632(15) Å] tin-sulfur distances. However, the Sn(1)-S(1) primary bonds are much longer than in **3** [2.4718(10) Å] and thus the extent of asymmetry is considerably reduced ($\Delta = [Sn(1)-S(2)] - [Sn(1)-S(1)]$ is 0.077 Å in **4** *vs*. 0.212 Å in **3**).

The resulting coordination geometry around the tin atom is highly distorted octahedral, with *cis* phenyl groups $[C(4)-Sn(1)-C(4a) 98.2(3)^{\circ}]$ and axial shorter tinsulfur bonds $[S(1)-Sn(1)-S(1a) 152.83(7)^{\circ}]$. The distortion is mainly due to the small bite of the ligand [S(1)-Sn(2) 2.949(2)] and is reflected by the dihedral angles: $C(4)Sn(1)C(4a) / S(1)Sn(1)S(1a) 83.2^{\circ}$; $S(1)Sn(1)S(2) / S(1a)Sn(1)S(2a) 72.5^{\circ}$. Similar coordination geometries were reported for other related diphenyltin(IV) derivatives, *i.e.* Ph₂Sn(S₂CNEt₂)₂ (regardless the monoclinic [18,19] or tetragonal [20] forms) or Ph₂Sn(S₂CNCy₂)₂ [21]. By contrast, for the Me₂Sn(S₂CNMe₂)₂ the geometry at tin was found to be a highly distorted skew-trapezoidal bipyramid, with angular C-Sn-C angle [136.45(1)^{\circ}] and a planar equatorial SnS₄ system [22].

With respect to the dimethyldithiocarbamato ligand in compounds **3**, **4** and **6** some common features should be noted. The MS_2CNC_2 skeleton is basically planar. Although, consistent with the *asymmetric monometallic biconnective* pattern, short and long metal-sulfur distances are established, in the dithio ligand unit the sulfur-carbon distances are almost equivalent on the basis of the 3 σ criteria [C(1)-S(1) 1.739(4) Å in **3**, 1.754(9) Å in **6** and 1.728(5) Å in **4** *versus* C(1)-S(2) 1.713(4) Å in **3**, 1.704(8) Å in **6** and 1.702(5) Å in **4**, respectively]. The carbon-nitrogen bond distance and the planarity of the NC₃ fragment reflect the double bond character and the *sp*² hybridization of the nitrogen atom, respectively.

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Dedicated to professor Gh. Marcu at his 80th anniversary

CHEMICALLY MODIFIED BASIC ALUMINA N FOR TLC. SYNTHESIS, CHARACTERIZATION AND APPLICATIONS

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ABSTRACT. The synthesis of chemically modified basic alumina N has been performed by organosilylation reaction of hydroxylated alumina surface with trifunctional modifiers: *n*-octadecyltrichlorosilane and 3-mercaptopropyltrimethoxysilane.

These chemically modified aluminas were characterized by elemental analysis, specific surface area, FTIR and ¹³C-CP/MAS NMR spectroscopy, mass spectrometry and thermoanalytical methods.

Chromatographic behaviour of unmodified and chemically modified aluminas were tested by the separation and identification of some dyes and benzo[a]pyrene derivatives.

Keywords: Chemically modified basic alumina N, Thin layer chromatography, FTIR spectroscopy, ¹³C-CP/MAS NMR spectroscopy, Mass spectrometry, Thermal analysis.

The progress of chemically modified stationary phases with applicability in liquid chromatography have been permitted the use of the inorganic oxides, such as alumina, titania and zirconia, as supports due to their specific properties or as an alternative for silica gel. Alumina is a stationary phase widely used in liquid chromatography due to its special chromatographic properties and of inherent higher pH stability [1-3].

The sort of alumina frequently used in thin layer chromatography (TLC) is γ -Al₂O₃ which has on its surface three types of active centers: oxygen anions, surface hydroxyl groups and aluminium cations [4].

In order to increase the selectivity and the efficiency of the chromatographic separation, the chemical modification with a high variety of organic compounds attached to the hydroxylated surface of adsorbent is a very useful method [1,4,5].

The characterization and the applications of reversed phases based on alumina are presented in literature [2,3,6,7].

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Infrared studies on chromatographic aluminas have been mainly focused on the hydroxyl groups and Al–O–Al linkages created during the dehydration process [3]. Diffuse reflectance infrared Fourier transform (DRIFT) spectra have been used to the characterization of chemically modified aluminas [8-10]. The study of chemically modified stationary phases by mass spectrometry is not frequently met in literature [11-15].

The TG (thermogravimetry), DTG (derivative thermogravimetry) and DTA (differential thermal analysis) thermoanalytical methods are not usually used for the study of modified stationary phases. However, these methods are suitable for the characterization of chromatographic adsorbents. The shape of thermoanalytical curves gives information about the mass losses of the material and the effects that occur [7,16,17].

EXPERIMENTAL

REAGENTS:

Basic alumina N was supplied by Macherey-Nagel (Düren, Germany).

n-Octadecyltrichlorosilane, 3-mercaptopropyltrimethoxysilane, hexane and the dyes: Toluidine blue, Bromothymol blue, Sudan black B, Sudan III, were purchased from Merck (Darmstadt, Germany).

Xylene, dichloromethane, diethyl ether, acetone, benzene, ethanol, methanol, carbon tetrachloride were purchased from Chimopar Bucharest (Romania).

The benzo[a]pyrene derivatives: benzo[a]pyrene-*trans*-7,8-diol-9,10-epoxy, benzo[a]pyrene-*trans*-9,10-dihydrodiol, benzo[a]pyrene-*trans*-7,8-dihydrodiol, benzo[a]pyrene-8-hydroxy, benzo[a]pyrene-7-hydroxy were purchased from Fluka (Switzerland).

SYNTHESIS OF CHEMICALLY MODIFIED BASIC ALUMINA N:

The chemically modified aluminas were prepared by the silulation reaction of basic alumina N with the modifiers: n-octadecyltrichlorosilane and 3-mercaptopropyltrimethoxysilane in the mass ratio of 2.5:1 (w/w) [7].

The obtained product was filtered and washed successively with xylene, dichloromethane and diethyl ether until the complete removal of the modifier traces.

CHARACTERIZATION OF CHEMICALLY MODIFIED BASIC ALUMINA N (Abbreviations of the studied samples are given in TABLE 1) has been performed by:

The *elemental analysis* (carbon, hydrogen and sulphur) of samples was performed by means of VARIO EL Elemental Analysis System.

The *specific surface area* of samples was determined using the BET method (krypton adsorption at the temperature of liquid nitrogen).

The *infrared spectra* of alumina samples were registered on a JASCO-610-FTIR spectrometer, using the KBr pellet technique. To improve the sensitivity of the IR method the difference and second derivative spectra were also analyzed.

¹³C *CP/MAS NMR spectra* were recorded on a BRUKEŔ DSX 200 spectrometer with samples of 200-300 mg in a double bearing 7 mm rotors of ZrO₂. Magic Angle Spinning (MAS) was carried out at 3500 Hz. The spectra were recorded with a proton pulse of 6.5 μ s, a recycle delay of 1 s and a contact time of 1 ms.

The mass spectra (EI-MS) were registered on a FINIGAN MAT 311 spectrometer with double focusing in the following conditions: 70 eV electron energy, 100 μ A emission current, 180°C ion source temperature. The sample has been introduced directly in the ion source and heated up to 320°C in 10⁻⁶ torr vacuum. The mass spectra of all samples have been continuously registered.

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The *thermal analysis* was performed by means of an OD 102 Derivatograph (MOM, Hungary) in the conditions: 200 mg sample quantity, 10°C/min rate of increase of oven temperature and TG - 100 mg; DTA - 1/5; DTG - 1/5 sensitivities respectively. The measurements were conducted in air atmosphere in the temperature range of 20-1000°C.

The chromatographic testing was performed on 10x14 cm glass plates prepared both from unmodified and from chemically modified alumina N. The plates were coated with 0.25 mm layers by the application of an ethanolic slurry of a mixture of the stationary phase and an organic binder [6,7].

The standard solutions (0.1%) of dyes singly or as a mixture prepared in ethanol, both singly and as a mixture, were applied as 5 μ L spots on the chromatographic plates by means of Brand micropipettes. Standard solutions (0.05%) of benzo[a]pyrene derivatives were prepared in benzene and spotted on the chromatographic plates in the same mode. Ascending development to a distance of 10 cm was performed at room temperature in an unsaturated Stahl chamber for all studied compounds. The dyes were developed using the dichloromethane - diethyl ether - acetone (20 + 70 + 10, v/v) mixture as mobile phase. The benzo[a]pyrene derivatives were developed using the benzene - acetone (70 + 30, v/v) mixture as mobile phase and the visualization was performed at 365 nm by a Camag lamp.

RESULTS AND DISCUSSIONS

The ideal representation of the *synthetized chemically modified aluminas* is shown in FIGURE 1.



Figure 1. Chemically modified alumina: a, alumina-C18; b, alumina-SH

The obtained chemically modified aluminas were characterized by elemental analysis, specific surface area, coverage density, FTIR spectroscopy, ¹³C CP/MAS NMR spectroscopy, mass spectrometry, thermal analysis and chromatographic behaviour by TLC separation of some dyes on unmodified and chemically modified alumina layers.

Elemental analysis data (combustion method), *specific surface area* (BET method) and *coverage density* (calculus) for unmodified and chemically modified aluminas, are presented in TABLE 1.

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TABLE 1

Alumina	Abbreviation	C (%)	H (%)	S (%)	S _{BET} (m²/g)	α (μmol/m²)
basic alumina N	alumina	-	-	-	114.2	-
<i>n</i> -octadecyl basic alumina N	alumina-C18	3.22	1.66	-	89.0	1.37
3-mercaptopropyl basic alumina N	alumina-SH	2.78	0.67	1.27	96.9	7.6

Coverage density of unmodified and chemically modified aluminas (% C; % H; %S; specific surface area – S_{BET} ; coverage density– α)

The data in this table show the presence of carbon, hydrogen and sulphur and the decreasing of specific surface area of the chemically modified aluminas.

Using *FTIR spectroscopy*, the presence of the modifier on alumina surface is well evidenced by the new bands observed in the 2800–3000 cm⁻¹ range.

Due to the low concentration of the organic part of modifier on the surface, the intensity of the new bands is weak (FIGURE 2).

The bands with maximum at ~ 2923 cm⁻¹ and at ~ 2853 cm⁻¹ are assigned to stretching vibrations of *n*-alkyl chain: $v_{as}(CH_2)$ and $v_{sym}(CH_2)$ respectively.



Figure 2. FTIR spectra of: a, alumina-C18; b, alumina-SH, c, alumina

In the difference FTIR spectrum (FIGURE 3) of unmodified and chemically modified alumina N the band observed at ~ 1468 cm⁻¹ is due to the δ (CH₂) vibration. The bands observed in the range of 900-1200 cm⁻¹ are assigned to v(Al-O) and v(Si-O) vibrations, showing the formation of Al-O-Si bridges.

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FIGURE 3. Difference FTIR spectrum of alumina-C18 and alumina

To improve the sensitivity of the infrared method, the second derivative spectra were also analysed in order to show that there are differences in the envelope of the infrared spectrum of unmodified and chemically modified alumina N (see FIGURE 4 for 3-mercaptopropyl alumina N).



Figure 4. Second derivative spectrum of alumina-SH (-) and alumina (...)

The ¹³C-CP/MAS NMR spectroscopy was used to study the conformational properties of immobilized ligands.

The observed signals in the NMR spectrum (FIGURE 5) were assigned to the corresponding carbon atoms as it is indicated. The high field shifted signal belongs to the C-1 atom, in the neighbourhood of the silicon (12.832 ppm). The low field shifted signals are characteristic for the carbon atoms C-2 (23.07 ppm, as a shoulder) and C-3 (27.859 ppm) of the mercapto grups.



Figure 5. ¹³C-CP/MAS NMR spectrum of alumina-SH

The comparative *mass spectrometry* study of unmodified and chemically modified alumina N puts in evidence the presence of the adequate chains on the modified alumina surface.

The mass spectrum (FIGURE 6) of *n*-octadecyl alumina N is dominated by the series of ions C_nH_{2n+1} (29, 43, 57, 71, ..., M-15) characteristic for the long normal hydrocarbon chain [14,15]. These ions are accompanied by those of C_nH_{2n-1} (27, 41, 55, 69,...) series. The formation of both ion series can be explained based on the fragmentation rules of hydrocarbons with normal chain. The periodicity of the main ions with a mass difference of $\trianglement multiplement multinter multiplement multiplement multinter multiplem$



Figure 6. Mass spectrum (EI-MS) in the maximum intensity point for alumina-C18





Figure 7. Intensity of characteristic ions *versus* temperature for alumina-C18: **a**) I total, **b**) $m/z = 44 [CO_2]^+$, **c**) $m/z = 36 [HCI]^+$ ion, **d**) m/z = 57 (organic part) ion

Over 100°C the characteristic ions for HCl (*m*/*z* 36 and 38) are visible, that confirms the fact that some initial radicals –Cl remain on the modified alumina surface. FIGURE 8 and FIGURE 9 show the mass spectrum in the maximum intensity point and the intensity of characteristic ions *versus* temperature, for 3-mercaptopropyl alumina N, respectively.



FIGURE 8. Mass spectrum in the maximum intensity point for alumina-SH

In the mass spectrum the main ions observed with: m/z 34, 47 and 76 correspond to SH₂, HS-CH₂ and HS-CH₂-CH₂-CH₃ structures and start around of 220°C. The characteristic ions appear around 250°C.

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FIGURE 9. Intensity of characteristic ions *versus* temperature for alumina-SH: **a**) I total, **b**) m/z = 34 $[SH_2]^+$ ion, **c**) m/z = 32 $[CH_3-OH]^+$ ion, **d**) m/z = 44 $[CO_2]^+$ ion, **e**) m/z = 76 $[C_3H_8S]^+$ ion

Over 180°C are visible the characteristic ions for CH_3 -OH (*m/z* 32) too, confirming the remaining of some initial radicals $-O-CH_3$. Thus the 2,3 functionality of trifunctional modifier is explained.

The comparative study of mass spectra of unmodified aluminas *versus* chemically modified alumina, puts in evidence the modifications of alumina surface with the corresponding *n*-octadecyl and 3-mercaptopropyl chains.

The **thermoanalytical (TG, DTG, DTA)** method gives quantitative information about the water removed by the desorption and dehydration processes (endothermal effect) and the elimination process of organic part (exothermal effect) of the studied stationary phases and data referring to the temperature ranges where the thermal effects take place.

Thermoanalysis results for aluminas (unmodified and chemically modified) are presented in TABLE 2 and FIGURE 10.

TABLE 2

							inas	
	Stationary	Endothermal	٨m	Exothermal effects				лм
	phase	effect t℃	%	Effect I, t°C	⊿m _l %	Effect II, t℃	⊿m _{ll} %	%
	alumina	80 (35–90)	1.0	165, 350 (90–700)	11.5	_	-	12.5
	alumina -C18	95 (35–160)	1.8	360 (160–480)	24.2	560 (480–700)	5.5	31.5
	alumina -SH	80 (100–240)	8.5	340 (240–410)	11.0	520 (410–800)	12.0	31.5

Thermoanalysis results for unmodified and chemically modified aluminas

The TG data indicate a continuous mass loss which depends on the nature and length of the chain bonded to alumina surface. The mass loss is less for unmodified alumina than for the modified material.

The DTA data show two types of thermal effects. The endothermal effect was attributed to the desorption and dehydration processes of superficial water. For the modified alumina N, the observed exothermal effects are assigned to the

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removal of species from the surface and to the elimination process of the organic part (see TABLE 2).

The different thermal behaviour of unmodified and modified alumina revealed the differences among these compounds as a result of the organosilanization reaction.



Figure 10. Thermoanalytical (TG, DTG, DTA) diagrams for: a) alumina, b) alumina-C18; c) alumina-SH

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The *chromatographic testing* of unmodified and chemically modified alumina N has been studied for the separation of some dyes and benzo[a]pyrene derivatives.

The parameters that define the TLC behaviour of the separated compounds on the unmodified and chemically modified aluminas [selectivity (β), efficiency (*E*) and resolution (R_s)] were calculated according to Equations 1-3 [18], for two neighbouring spots of studied compounds.

$$\beta = \frac{R_{f2} - R_{f1} \times R_{f2}}{R_{f1} - R_{f1} \times R_{f2}}$$
(1)

$$E = \frac{(\beta - 1)^2}{4\beta} \times (1 - R_{f_2}) \times (1 - R_{f_1}) \frac{1}{\sqrt{H_2 H_1}}$$
(2)

$$R_{\rm S} = \frac{\Delta R_{\rm f} \sqrt{Z_{\rm f} - Z_0}}{2(\sqrt{R_{\rm fl}H_1} + \sqrt{R_{\rm f2}H_2})}$$
(3)

where:

$$H = \frac{Z_{\rm f} - Z_0}{N}$$
, is the theoretical plate height and $N = 16 \left(R_{\rm f} \frac{Z_{\rm f} - Z_0}{\delta_{\rm x}} \right)^2$

the theoretical plate number. R_{f1} , R_{f2} , represent the retention factors of a neighbouring pair of substances, Z_{f} - Z_0 , the distance between origin and the mobile phase front and δ_x , the spot diameter.

The experimental average values for δ_x are 0.5 cm for unmodified alumina and 0.4 cm for chemically modified aluminas.

In FIGURE 11 are presented the chromatographic results obtained at the separation of some dyes on thin layers of unmodified and chemically modified alumina N using the dichloromethane : diethyl ether : acetone, (20 / 70 / 10, v/v) mixture as mobile phase.





The chromatographic results obtained at the separation of benzo[a]pyrene derivatives on unmodified and chemically modified aluminas are presented in FIGURE 12.

The results show that the values of the chromatographic parameters depend on the retention factors of two neighbouring compounds, the migration distance of mobile phase, the diameter of spots and the type of mobile and stationary phases, respectively.

Analysing the results presented in FIGURE 12 we can conclude:

- for R_f values from the 0.2-0.8 range, the selectivity has a small variation;

- the separation degree of two neighbouring compounds is given by the value of resolution;

- the efficiency of separation of two neighbouring compounds depends on the selectivity, $R_{\rm f}$ values and spot diameter.

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Mobile phase: benzene:acetone (70 / 30, v/v);

Benzo[a]pyrenes: Benzo[a]pyrene-*trans*-7,8-diol-9,10-epoxy (*syn*) (B1), Benzo[a]pyrene*trans*-9,10-dihydrodiol (B2), Benzo[a]pyrene-*trans*-7,8-dihydrodiol (B3), Benzo[a]pyrene-8hydroxy (B4), Benzo[a]pyrene-7-hydroxy (B5

The best values of the chromatographic parameters at the separation of some dyes and benzo[a]pyrenes derivatives have been obtained on *n*-octadecyl alumina N thin layers.

The chromatographic results demonstrated new retention properties for the chemically modified aluminas and the possibility to use them at different chromatographic applications.

CONCLUSIONS

Chemically modified alumina N samples have been obtained by the silylation reaction of basic alumina N with trifunctional modifiers, namely *n*-octadecyltrichlorosilane and 3-mercaptopropyltrimethoxysilane.

Elemental analysis and FTIR spectra of modified alumina show the presence of organic modifier on the alumina surface. An important change of specific surface area is obtained.

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The comparative mass spectrometry and ¹³C CP/MAS NMR spectroscopy study of unmodified and chemically modified alumina put in evidence the presence of the adequate chains on the modified alumina surface.

Thermoanalytical diagrams permitt to find out the type of thermal effects and the temperature ranges where they take place, as well as the corresponding mass loss values.

The chromatographic behaviour of chemically modified alumina proves their new retention properties at the separation of some dyes and benzo[a]pyrene derivatives.

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STUDIA UNIVERSITATIS BABES - BOLYAI, CHEMIA, LI, 2, 2006

Dedicated to professor Gh. Marcu at his 80th anniversary

SYNTHESIS AND VIBRATIONAL STUDIES ON NEW COMPLEXES OF MONODEPROTONATED (4H-5-MERCAPTO-1,3,4-THIADIAZOL-2-YL)THIOACETIC ACID

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ABSTRACT. Partial neutralisation of (4*H*-5-mercapto-1,3,4-thiadiazol-2-yl)thioacetic acid with 1 eq. of sodium bicarbonate produced the sodium carboxylate salt. This salt was further used in the preparation of the title compounds as four d metal complexes, Co, Ni, Cu and Zn. Preliminary vibrational studies on this new series are discussed in terms of FT-IR and FT-Raman spectroscopy. The spectral data are consistent with the coordination of the deprotonated carboxylic groups to the metal centers. In all cases, the occurrence of the thione tautomeric form of the heterocycle was found.

Key-words: (4*H*-5-mercapto-1,3,4-thiadiazol-2-yl)thioacetic acid, metal complexes, FT-IR and Raman spectroscopy.

INTRODUCTION

We have an on-going interest in the chemistry of mercapto-aza type heterocyclic derivatives mainly from the aspect of their coordination and supramolecular chemistry, and also because of their various applications in analytical chemistry, materials science, etc.¹⁻¹¹ Previous structural studies performed on well known analytical reagents revealed their extraordinary potential in this field. For example, 1,3,5-triazine-2,4,6-trithione, C₃N₃S₃H₃ (also known as trimercaptotriazine or trithiocyanuric acid) and 4-phenyl-2-mercapto-1,3,4-thiadiazole-5-thione (also known as *Bismuthiol II*) have been intensively used in removing heavy metals from waste waters and also as industrial precursors.¹⁻⁶ Recently, the co-crystallization of trithiocyanuric acid with melamine, tricyanuric acid, 4,4'-bipyridine, etc. generated supramolecular structures with nanometric cavities and channels. Many of these compounds have proved excellent zeolitic properties.⁷⁻⁹ Furthermore, crystallographic investigations on (organo)metallic complexes of such ligands revealed interesting 1D, 2D and 3D supramolecular architectures build up *via* intermolecular covalent and/or secondary interactions.^{10,11}

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It was concluded that the flat, rigid molecular geometry of the ligands, along with their increased number of heteroatoms E and EH groups (E: O, N, S, etc.), encourage the self-assembly of these molecules into layered channel-type crystal structures. Moreover, the involvement of the NCS groups into the building pattern is crucial.^{10,11} For example, the presence of exo- and endocyclic sulfur atoms may give rise to S···S interactions, which have been found of great importance in the supramolecular construction of molecular electronics.¹²

We have recently initiated the vibrational investigation of a new range of π -excessive heterocyclic systems based on dimercapto-thiadiazole, $C_2H_2N_2S_3$ (also known as *Bismuthiol I*).^{13,14}

We decided to use one of these compounds, the (4*H*-5-mercapto-1,3,4-thiadiazol-2-yl)thioacetic acid, $C_2HN_2S_3$ -CH₂COOH **1** (Scheme 1), in the syntheses of five new metallic derivatives: M($C_2HN_2S_3$ -CH₂COO)_n, n = 1, M = Na, **2**; n = 2, M = Co, **3**; Ni, **4**; Cu, **5**; Zn, **6**.





Hence, the aim of this work is the synthesis of the above series of derivatives together with their preliminary IR and Raman characterization. It could infer us about i) the occurrence of the thione *vs.* thiol tautomeric form of the heterocyclic moiety in solid state and ii) to suggest the most probable coordination sites in the series 2-6.

EXPERIMENTAL

FT-IR and FT-Raman spectra on solid samples were recorded using a Bruker FT-IR Equinox 55 Spectrometer equipped with an integrated FRA 106 S Raman module. The excitation of the Raman spectra was performed using the 1064 nm line from a Nd:YAG laser with an output power of 250 mW. An InGaAs detector operating at room temperature was used. The spectral resolution was 2 cm⁻¹.

The starting materials were purchased from commercial sources as analytical pure reagents and were used with no further purification. Compound **1** was prepared following a literature protocol.¹³

The pure monosodium salt of 1, $C_2HN_2S_3$ -CH₂COONa 2 was obtained by reacting stoechiometric amounts of acid 1 and sodium bicarbonate in aqueous solution, at room temperature and subsequent recrystallization of crude 2 from hot distilled water.

The transition metal complexes of **1** as monodeprotonated forms **3-6**, $[M(C_2HN_2S_3-CH_2COO)_2]$, M = Co, **3**; Ni, **4**; Cu, **5** and Zn, **6** were synthetised by reacting stoechiometric amounts of **2** with the corresponding metal salt $Co(NO_3)_2\cdot 6H_2O$, $NiCl_2\cdot 6H_2O$, $CuSO_4\cdot 5H_2O$ and $ZnCl_2$, respectively, in aqueous solution, at room temperature. In all cases, the complexes precipitated at once. After stirring the reaction mixtures for 1 - 2 hours, the crude products were filtered and purified by recrystallization from warm distilled water (**3** and **5**) or by washing

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with warm water (5 and 6). Preparation details and brief characterization of compounds 2 - 6 are given in Table 1.

Table 1

	Preparation details and characterization of compounds 2 – 6.						
No.	Compound	Yield (%)	mp (°C)	Aspect			
2	C ₂ HN ₂ S ₃ -CH ₂ COONa	86	268 - 270	colorless, crystalline			
3	$[Co(C_2HN_2S_3-CH_2COO)_2]$	52	151 - 153	pink , crystalline			
4	$[Ni(C_2HN_2S_3-CH_2COO)_2]$	57	183 - 185	pale green, crystalline			
5	$[Cu(C_2HN_2S_3-CH_2COO)_2]$	58	199 - 201	green, powder			
6	$[Zn(C_2HN_2S_3\text{-}CH_2COO)_2]$	52	218 - 220	white, powder			

RESULTS AND DISCUSSION

Partial neutralisation of (4*H*-5-mercapto-1,3,4-thiadiazol-2-yl)thioacetic acid, $C_2HN_2S_3$ -CH₂COOH **1** with sodium bicarbonate produced the monosodium salt, $C_2HN_2S_3$ -CH₂COONa **2**. Compound **2** was further used in the preparation of four d metal complexes, [M($C_2HN_2S_3$ -CH₂COO)₂], M = Co, **3**; Ni, **4**; Cu, **5** and Zn, **6** (Scheme 2). All the reactions worked in aqueous media, at room temperature. Compounds **2** – **4** are soluble in water and were isolated as crystalline solids. Compounds **5** and **6** are not soluble in common solvents and were isolated as powders. All metal derivatives are air and light stable in solid state.

$$2 C_2 HN_2 S_3 - CH_2 COOH \xrightarrow{+ 2 NaHCO_3} 2 C_2 HN_2 S_3 - CH_2 COONa \xrightarrow{+ M^{2+}} [M(C_2 HN_2 S_3 - CH_2 COO)_2]$$

$$\xrightarrow{- 2 H_2 O} 2 Na^+$$

$$M = Co, 3; Ni, 4; Cu, 5; Zn, 6$$

Scheme 2

The FT-IR and Raman spectra of 2 - 6 were recorded in the 4000 – 400 and 3500 – 200 cm⁻¹ spectral ranges, respectively. For the copper(II) complex **5**, the Raman spectrum could not be recorded. The most relevant spectral data are listed in Table 2. The following discussion is based on the comparison between our spectral data recorded for 2 - 6 and the known literature data for **1** and *Bismuthiol L*^{13,14}

The $4000 - 2000 \text{ cm}^{-1}$ *spectral range* was relevant for the characterization of NH, OH, SH and CH groups in IR and/or Raman spectra.

M.M. VENTER, V. CHIŞ, S. CÎNTĂ PÎNZARU', V.N. BERCEAN, M. ILICI, I. HAIDUC

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However, the presence of the carboxylic OH group in the starting acid **1** gave rise to a large and complex IR band (approx. $3100 - 2500 \text{ cm}^{-1}$) which overlapped the other expected fundamentals and makes their assignation less accurate. In the IR spectra of **2** - **6**, the title band disappeared as a result of carboxylic group deprotonation. On the other hand, the same spectra show broad bands at 3487 - 3385 cm⁻¹ which may be assigned to the v(H₂O) mode, as well as weak to medium bands at 3106 – 3090 cm⁻¹ which are tentatively assigned to the v(NH) fundamental.

The Raman spectra of **1** – **6** were more suitable for the interpretation of the tautomeric form of the heterocycle and the characterization of the alkyl fragment. Thus, all spectra lack the characteristic v(SH) bands, usually assigned in the 2600 – 2400 cm⁻¹. In addition, the $v_{as}(CH_2)$ and $v_s(CH_2)$ stretching vibrations may be clearly assigned at 3022 – 2953 and 2944 – 2905 cm⁻¹, respectively.

It can be concluded that the deprotonation of the acid **1** took place at the COOH group and the ligand participates in coordination as carboxylato anion. In addition, we tentatively suggest that the remaining proton was located at an azaatom, which gave option to the thione tautomer form of the heterocyclic unit.

The 2000 – 1000 cm⁻¹ spectral range was relevant for both heterocyclic and alkyl-carboxylic units of the coordinated ligand. The most important as characteristic bands for the heterocyclic fragment fall in the 1541 – 1433, 1120 – 1103 and 1059 – 1045 cm⁻¹ regions and are assigned to the v(C=N), v(N-N) and v_{as}(S-C=S) modes, respectively. The first stretching mode consists of two components, which are assigned in the spectra of **2** - **6** at 1541 – 1515 and 1457 – 1433 cm⁻¹. These components are comparable with the two corresponding fundamentals found for **1** (1494 and 1448 cm⁻¹) and related mercapto-thiadiazoles (e.g. 1510 –1506 and 1452 -1450 cm⁻¹ for *Bismuthiol I*). Similarly, the assignment of the other two stretching modes is consistent with the literature data for **1** and *Bismuthiol I*.

As it concerns the alkylcarboxylic substituent, the title spectral range offers consistent information about both CH₂ and COOH groups but the most outstanding behavior was revealed by the COOH / COO⁻ groups. According to literature, ¹³⁻¹⁵ the COOH group is defined by two major modes: v(C=O) and v(C-OH); the later may be represented by two bands. As the deprotonation occurs, the resulting COO⁻ groups is expected to adopt a C_{2v} symmetry and to reveal the v_{as}(COO) and v_s(COO) fundamentals. In the case of compounds **1** – **6**, the first mode of both groups was easily assigned. Thus, for compound **1** the band corresponding to v(C=O) falls at 1693 cm⁻¹ in both IR and Raman spectra while in the spectra of the metal derivatives the same mode is shifted dramatically to lower wave numbers (1641 – 1558 cm⁻¹) and can be related to the v_{as}(COO) fundamental. The assignment of the v(C-OH) / v_s(COO) modes in the expected region (1400 – 1200 cm⁻¹) proved very difficult due to the overlap with a significant number of bands. Despite this ambiguity, we propose a tentative assignment of these fundamentals at 1303 in **1** and 1333 – 1293 cm⁻¹ in the series **2** – **6**.

The 1000 – 200 cm⁻¹ spectral range shows no unusual features. The plethora of bands located in this region was assigned to group bending and other skeleton vibrational modes (i.e. $v_{as}(CSC)_{endo}$ 738 – 717 and $v_s(CSC)_{endo}$ 678 – 657 cm⁻¹).

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CONCLUSIONS

In the explored series of metal complexes, the ligand (4*H*-5-mercapto-1,3,4-thiadiazol-2-yl)thioacetate is present as carboxylato anion, saving the thione tautomer form. The coordination of the ligand to the metal centers M^{2+} (M = Co, Ni, Cu and Zn) is suspected to occur through the carboxylato group. We avoid, however, to evaluate its coordination behavior by comparing the $\Delta v = v_{as}(COO) - v_s(COO)$ values until more spectral and structural investigations will be completed on the title metal complexes.

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Dedicated to professor Gh. Marcu at his 80th anniversary

ACCELERATOR MASS SPECTROMETRY RADIOCARBON DATING OF AN OLD TROPICAL TREE: PRELIMINARY REPORT. 1. RADIOCARBON DATES.

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ABSTRACT. A number of 11 samples collected from the remains of a very large African baobab (*Adansonia digitata*) were processed and investigated by AMS radiocarbon dating. Radiocarbon dates of several samples, collected from different areas of the collapsed trunk, were found to be greater than 1000 yr BP. Several additional samples are under investigation.

KEY WORDS: radiocarbon dating, AMS, tropical trees, age determination, dendrochronology, dendroclimatology.

INTRODUCTION

In present, a number of 16 "millenarian" tree species with accurate dating, i.e. species with individuals that had lived over 1000 years (yr), are known [1, 2]. All these tree species are gymnosperms/conifers. The champion species is the bristlecone pine (*Pinus longaeva*), with an age of 4844 yr for the cross-dated WPN-114 or Currey (Prometheus) tree [3].

Somewhat surprisingly, no angiosperm tree with a confirmed age of over 1000 yr has been identified, so far. The oldest angiosperm tree reported in the literature is an African baobab (*Adansonia digitata*), felled in 1960 at Lake Cariba (in former Rhodesia, today Zimbabwe). This specimen, with a girth of 14.4 m (at a height of ca. 0.6 m above ground level), was far from being one of the largest trees of the species. It was dated

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by Swart via radiocarbon dating (using the beta decay counting method), who reported a radiocarbon age of 1010 ± 100 yr BP for a sample collected from the core of the stump [4]. After calibration, this value corresponds to a calendar age of 925 ± 105 yr (in 1960, when the tree died).

One should mention a singular study, which claimed incredible age values of over 1000 years, determined by radiocarbon dating, for two tropical trees of the Amazonian rainforest (i.e. 1370 yr for *Cariniana micrantha* and 1170 yr for *Dipteryx odorata*) [5]. These values contradict dramatically other results concerning the age limit of similar tree species from the Amazonian rainforest and have been questioned by several researchers [6-8]. It was mentioned that the respective results were outliers to all other findings in the dating of tropical trees. Criticism was raised due to the fact that results were based on single samples of each tree and on single radiocarbon measurements, without any corroborating data or replicated analyses [8].

According to a number of researchers, the species of the genus Adansonia are the main angiosperm candidates to become "millenarian". The genus Adansonia, which belongs to the Bombacacee family, comprises eight species of deciduous trees: six are endemic to Madagascar, one to Australia and one is widespread in continental Africa [9].

The African baobab (*Adansonia digitata* L.), which can be considered a tropical tree, is the largest and best-known of the eight *Adansonia* species. It is widespread south of the Sahara, especially in savanna regions. The familiar picture of the African baobab is of an almost grotesque tree, with a girth out of proportion to its height. Therefore, the African baobab is often named the upside-down tree. Although the baobab is a relatively short tree, with an usual height around 20 m, some individuals can reach an exceptionally large girth of over 25 m [10-12]. The African baobab is a very special tree. Baobabs are more similar to succulents than they are to woody trees. Thus, as they die, they might best be described as rotting, rather than slowly decaying. The trunk and the branches are constructed in concentric layers of succulent tissue, reminiscent of the layers of an onion. The wood is soft, spongy, fibrous and light; between each layer of xylem cells there is a layer of parenchyma cells that store water [10]. Thus, the African baobab can be considered a stem-succulent tree species. In the rainy season, baobabs store enormous quantities of water in their bulbous trunks, which they use to survive during the dry season.

The tremendous size of certain individuals has led many observers to state that the baobab lives to a long age. However, the age of the oldest baobab trees, i.e. the age limit of the African baobab, remains a controversial topic, which has generated two competing hypotheses:

(i)*The long lived baobab hypothesis* (baobabs as millenarian trees) is based on local traditions, the statements of pioneer explorers and the study of several large and old specimens. Early African explorers, such as Adanson and Livingstone, ventured to extrapolate the very slow growth rate of old baobabs over their entire life cycle, without any corrections, thus claiming incredible age values of several thousands years (up to 5150 yr) for the largest individuals [13, 14]. Modern measurements of several huge individuals from Botswana and Mozambique, performed by Guy, when compared to historic records of the same trees, evinced a very small increase or even a decrease in girth during a time span of ca. 110 yr, namely from ca. 1850–60 to1966 [15]. The studies of Guy and Swart evinced very ACCELERATOR MASS SPECTROMETRY RADIOCARBON DATING OF AN OLD TROPICAL TREE: ...

small growth rates (i. e. annual increase in radius) for old baobabs of Southern Africa (South Africa, Zimbabwe, Botswana, Mozambique) [15, 16, 4]. The singular result of Swart also suggests that much larger individuals than the radiocarbon dated tree could be well over 1000 yr old. [10, 12, 16].

(ii) *The short lived baobab hypothesis* (baobabs as centenary trees), promoted by many contemporary researchers, is based on several relatively recent studies that reveal very fast growth rates, especially for young baobab trees of Central Africa (Zambia, Sudan, Mali, Kenya, Tanzania) and of South Africa, as well [18-23]. These researchers, who discount or question the dating result of Swart, claim that the age limit of the African baobab is around 500–800 yr, perhaps reaching values close to 1000 yr, in the case of the largest trees [22-25]. Several additional remarks should be made. Even if an enormous girth would suggest a very old tree, the largest baobabs are not necessarily the oldest ones. As in the case of other tree species, the size of the baobab depends not only on its age, but also on the genetic variability and on its location, especially its access to moisture [26]. Consequently, there is a big problem to estimate the age of baobabs via girth measurements, due to the impact of the hydrostatic conditions within the tree and to the large variation in growth rates between individuals. In addition, most baobab researchers consider that the growth rate decreases severely with age.

Given that the trunk of the baobab is a water-storing organ, it swells and shrinks periodically, in direct relation to the water content. Seasonal, annual and long period girth variations have been reported; they may sometimes mask the radial increase du to cambial activity.

The baobab stops growing during the dry season and produces a sort of rings. These faint rings are believed by modern researchers to be annual rings, even if this has not been yet demonstrated conclusively [24]. The majority of researchers agree, however, that in areas with a distinct dry season, the baobab exhibits annual rings. Ring counting of young trees of known age have been shown to be within 2 % of the real age [10]. However, ring counting cannot be used to aging large baobabs, because of the presence of large hollows in the trunk.

Further, large baobabs have sometimes multiple stems, more or less fused in a single trunk. This could either be ascribed to multiple sprouting from the same rootstock or to the simultaneous germination of several seeds [24]. A baobab tree with two or more fused stems at an earlier stage of its life can be younger than its girth would suggest [12].

It should also be noticed that, when they die, baobabs collapse into a huge fibrous mass. The wood fibers dry out rapidly and the heat of the desert during the dry season may ignite them. It is also possible that the dehydrated, dry and friable wood fibers may be washed away by heavy rains over the rainy season or that they may crumble and be blown away by winds. Hence, a huge collapsed tree may disappear in a relatively short time, with no trace left behind. This fact may be at the origin of the popular myth which states that, after they die, baobabs catch fire spontaneously [24].

Hence, the sole technique for determining the age of old baobabs is radiocarbon dating of wood samples collected from a recently fallen tree, namely by measuring their ¹⁴C content relative to the content of stable carbon.

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In 1949, Libby pioneered the first radiocarbon measurements by monitoring the beta radioactive decay of individual ¹⁴C atoms with modified Geiger counters. He used samples of several grams of carbon-black powder. In the 1950's period, when atmospheric nuclear bomb tests were performed, this method was relatively insensitive and subject to statistical errors due to the absorption of nuclear contaminant [27].

More accurate methods in the decay counting technology had been developed by the conversion of sample solid carbon to CO₂ for measurement in gas-proportional counters and liquid-scintillation counters. Both methods rely on observing the decay of the radioactive ¹⁴C atoms. When a ¹⁴C atom decays, it emits a beta particle, which can be counted in a gas by the electrical pulse it generates. In a liquid-scintillation counter, the beta particle excites the emission of light from the molecules of an organic solvent, which acts as a "scintillant". Because only ca. 13.5 decays per minute occur in 1 g of modern carbon, it is necessary to use fairy large samples, up to several grams of carbon. Today, the majority of radiocarbon laboratories utilize these two dating methods [27].

It was recognized that direct counting of ¹⁴C atoms in the sample would greatly enhance the sensitivity and accuracy of the dating. Some unsuccessful attempts were made by using conventional mass spectrometry. In 1977, a new development added a particle accelerator into a mass spectrometer to produce an accelerator mass spectrometer [28]. The technique uses a very high performance particle accelerator to accelerate sample atoms as ions to high energies. After a first attempt which utilized positive ion acceleration [29], the superior approach with negative ion acceleration was developed [30-31]. This so-called Accelerator Mass Spectrometry (AMS) method allows the determination with high accuracy of tiny amounts of isotopes with a small relative abundance. The dating with radiocarbon by AMS is of very high sensitivity, up to 1:10¹⁵ atoms for carbon, which is 10³ to 10⁵ higher than the older decay counting methods. Thus, it allows for the analysis of very small samples, well below 1 mg, in a much shorter time. The higher costs are justified by the value of results. Over the past years, new concepts and technologies have been brought into the field of AMS radiocarbon dating and new types of accelerators and ion sources have been developed [32].

Grootboom, an exceptionally large African baobab (Photo 1), collapsed unexpectedly in north-eastern Namibia, at the end of 2004. The huge dimensions of the tree, associated with the unusual fact that it did not have very large hollow parts, offered an unique opportunity to clarify certain aspects concerning the controversial age of large baobabs.

An international joint research project was initiated, with the following aims: 1) to determine accurately the true age of the tree; 2) to establish whether Grootboom's trunk was a single unit or was made up of several fused stems; 3) to learn about the dynamics of Grootboom's growth rate during its life cycle.

Several samples were collected from the remains of the collapsed tree and were processed and analyzed by AMS radiocarbon dating.

EXPERIMENTAL SECTION

Collection of samples. Grootboom collapsed stepwise and very chaotically into six stems. Several sets of wood samples have been collected from two fallen stems (samples Nos. 5–10), as well as from the remaining stumps of two other stems (Nos. 1–4), that broke at a variable height from the ground and collapsed. Two stems were not available for sampling, one being already severely decayed, while the other 74

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was covered and partially crushed by a larger stem. Another sample (No. 11) was collected from towards the upper part of the tallest unbroken fallen branch.

Several additional samples collected from the remaining stumps, from the fallen stems and from the thickest exposed root, as well, are being currently dated.



Photo 1. The image (facing north) shows the heavy trunk of Grootboom and its rich crown full of leaves, at the beginning of the rainy season 2001–2002.

The sampling of Grootboom was recorded on videotape, a number of photographs were taken and several measurements of the remains of the collapsed tree were undertaken. This allowed the subsequent establishment of the position of each sample in the trunk of the tree prior to its collapse.

Sample preparation. The preparation of the samples for AMS radiocarbon dating involved the standard steps: i) pretreatment of the wood samples to isolate cellulose from the bulk tissue; ii) combustion of the separated cellulose to CO_2 ; iii) reduction of CO_2 to graphite.

Pretreatment of samples. Wood samples (with a mass of 0.7–2.0 g) were subsampled by cutting with a clean razor blade. The wood shavings were weighed and placed into a pre-combusted glass centrifuge tube to undergo numerous wet chemistry pretreatment steps. The woody tissues of trees, involved in water transport, contain several hydrolizable and mobile organic substances. Pretreatment intends to remove non-structural mobile carbon and to isolate only structural non-mobile carbon components, mainly cellulose, which had been synthesized at the original formation of the respective woody tissue.

The pretreatment method used was, with the exception of one sample, the so-called Acid-Base-Acid (ABA), which is widely employed in the case of wood samples prepared for AMS analysis. This classic method consists of a sequential washes of weak acids and bases and yields a residue consisting primarily of cellulose [33].

First, wood samples were treated with HCl solution (10% v/v), and then placed in a water bath at 60° C, for 1 h, to remove any inorganic carbon. The acid solution was discarded by means of a pipette and the sample rinsed with organic-

free water to neutral pH. NaOH solution (2% w/v) was added to the centrifuge tube, which was placed in the water bath for 1 h. The samples were treated with multiple base rinses until the solution was clear and free of humic or fulvic acid discoloration. The base solution was removed with a pipette and the sample was rinsed to neutral pH with organic free water. HCl solution (10% v/v) was added to drive off any CO₂ gas that may have adsorbed onto the sample during processing. After being heated in the water bath for 1 h, the final acid solution was removed by pipette and the sample was rinsed again to neutral pH. Subsequently, the remaining residues of each sample were filtered on pre-combusted quartz fiber filters, which were dried overnight in a drying oven at 60°C. All samples were stored in a dessicator, to await combustion.

One sample (No. 7) had the wood fibers quasi-totally destroyed, due to severe decay caused by microorganisms. Because in such cases the ABA method can produce large errors, the α -Cellulose (Jayme-Wise) method was used for pretreatment of this sample. The method involves an organic solvent extraction, followed by a bleach delignification process, which leaves only α -cellulose as residue.

 α -Cellulose was extracted according to the method described by Jayme and Wise [34] and modified by Loader et al. [35]. Briefly, dried wood shaving was extracted with 9:1 (v/v) dichloromethane and methanol, using an accelerated solvent extractor (ASE 200, Dionex Corp., Sunnyvale, USA) at 100 °C and 1000 psi. The resulting sample was dried in an oven at 50 °C. Then, the sample was introduced in a glass centrifuge tube and a bleaching solution of sodium chlorite (0.19 mol dm³) and HCI (0.07 mol dm⁻³) was added. The centrifuge tube was placed in an ultrasonic bath at about 75°C for 2 h, after which the acid solution was removed with a pipette. Three further additions of the bleaching solution were made, one after each hour. The resulting sample was thoroughly rinsed with pure water.

Next, a solution of NaOH (10% w/v) was added to the tube. The tube was placed in the ultrasonic bath for 45 min at 75 °C. The sample was washed with pure water and treated with a solution of NaOH (17% w/v) in the ultrasonic bath at room temperature. The sample was washed with water and then with a diluted solution of H₃PO₄ (3% v/v). Finally, a large volume of cold pure water was used to wash the sample to neutral. The remaining α -cellulose residue was dried at 50 °C.

Combustion to CO_2 . The combustion method used for the resulted cellulose samples was the Closed Tube Combustion (CTC) method [36]. Samples were weighed and placed into pre-combusted quartz combustion tubes with 2 g copper oxide wires (to provide oxygen for the combustion) and 100 mg of silver powder (to scavenge sulfur and chlorine gases). The combustion tubes were then placed in a vacuum line where they were evacuated, leak-checked and flame-sealed. They were combusted in a muffle furnace at 850°C, for 5 h. The generated CO_2 was cryogenically purified, quantified and transferred to a graphite reactor.

Reduction to graphite. In the graphite reactor, the generated CO_2 was converted to graphite by Fe catalysed reaction in a H_2 atmosphere. The graphite samples were submitted to AMS analysis.

AMS radiocarbon analysis. The radiocarbon (¹⁴C) content of the graphite samples was determined at the National Ocean Sciences Accelerator Mass Spectrometry (NOSAMS) Facility of the Woods Hole Oceanographic Institution.

Graphite derived from each sample was compressed into a small cavity in an aluminum "target", which acted as a cathode in the ion source. The surface of the graphite was sputtered with heated cesium and the ions produced were extracted and accelerated in the 3 MV Tandetron (TM) AMS system. After acceleration and ACCELERATOR MASS SPECTROMETRY RADIOCARBON DATING OF AN OLD TROPICAL TREE: ...

removal of electrons, the emerging positive ions were magnetically separated by mass and the ¹²C and ¹³C ions were measured in Faraday Cups, where a ratio of their currents was recorded. Simultaneously, the ¹⁴C ions were recorded in a gas ionization counter, so that instantaneous ratios of ¹⁴C to ¹³C and ¹²C were recorded. These are the raw signals that were ultimately converted to a radiocarbon age [37].

The AMS analysis of the samples was performed relative to an array of primary and secondary standard samples. Two primary standards were used: NIST Certified Oxalic Acid I (NIST-SRM-4990) and the derived Oxalic Acid II (NIST-SRM-4990C). The ¹⁴C activity ratio of Oxalic Acid II (Fm = 1.3605; δ^{13} C = -17.3 ⁰/₀₀) to Oxalic Acid I (Fm = 1.0520; δ^{13} C = -19.0 ⁰/₀₀) is taken to be 1.293. Three secondary standards were also used, namely FIRI B (Old Wood, Consensus Fm = 0.0033, NOSAMS Fm = 0.5668) and FIRI H (Dendro-dated Wood, Consensus Fm = 0.7574, NOSAMS Fm = 0.7450). System background measurements were performed with petroleum-derived Johnson-Matthey 99.9999% graphite powder (no chemical preparation, Fm < 0.002) and IAEA C-1 hydrolyzed Carrara marble (Fm < 0.002). The measurement of the secondary standards was used to assess the accuracy of the sample analysis. The overall system error is a function of the average deviation of the secondary standard results from their respective consensus values [38-40].

Radiocarbon date/age and auxiliary quantities/values. The quantities/values calculated from the experimental results of the AMS analysis are the following:

Fraction modern (carbon). Fraction modern (carbon) is a measurement of the deviation of the ¹⁴C/¹²C ratio of a sample from "modern." Modern is defined as 95% of the radiocarbon concentration (in AD 1950) of a NIST

Oxalic Acid I normalized to $\overline{\delta}^{13}C_{VPDB} = -19^{-0}/_{00}$ [E7]. AMS results are calculated using the internationally accepted modern ${}^{14}C/{}^{12}C$ ratio of 1.176 ± 0.010 x 10⁻¹² [38]; all results are normalized to -25 ${}^{0}/_{00}$ using the $\overline{\delta}^{13}C_{VPDB}$ of the sample. The value used for this correction is specified in the report of final results. Fraction modern (Fm) is basically computed from the expression: Fm = (S – B) / (M – B), where B, S and M represent the ${}^{14}C/{}^{12}C$ ratios of the blank, the sample and the modern reference, respectively. $\overline{\delta}^{13}C$ *Correction*. In addition to loss through decay of radiocarbon, ${}^{14}C$ is affected by natural isotopic fractionation. Fractionation is the term used to describe the differential uptake of one isotope with respect to another. While the three carbon isotopes are chemically indistinguishable, lighter ${}^{12}C$ atoms are preferentially taken up before the ${}^{13}C$ atoms in biological pathways. Similarly, ${}^{13}C$ atoms are taken up before ${}^{14}C$. The assumption is that the fractionation of ${}^{14}C$ relative to ${}^{12}C$ is twice that of ${}^{13}C$, reflecting the difference in mass. Fractionation must be corrected for in order to make use of radiocarbon measurements as a chronometrical tool for all parts of the biosphere. In order to remove the effects of isotopic fractionation, the Fraction Modern is then corrected to the value it would have if its original $\overline{\delta}^{13}C$ were $-25^{-0}/_{00}$ (the $\overline{\delta}^{13}C$ value to which all radiocarbon measurements are normalized).

 δ^{13} C were $-25^{0}/_{00}$ (the δ^{13} C value to which all radiocarbon measurements are normalized). The Fraction modern corrected for δ^{13} C, noticed by $\text{Fm}_{\delta}^{13}_{\text{C}}$, is calculated by: $\text{Fm}_{\delta}^{13}_{\text{C}} = \text{Fm} \left[(1 - 25 \cdot 10^{-3}) / (1 + 10^{-3} \delta^{13} \text{C}) \right]^2$. *Errors*. Atoms of ¹⁴C contained in a sample are directly counted using the

Errors. Atoms of ¹⁴C contained in a sample are directly counted using the AMS method of radiocarbon analysis. Accordingly, an internal statistical error is calculated using the total number of ¹⁴C counts measured for each target, $\pm(n)^{1/2}$. An external error is calculated from the reproducibility of multiple exposures for a given target. The final error is the largest of the internal or external errors.
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Aside from the normal statistical errors intrinsic to the counting of ¹⁴C events, there are additional statistical errors associated with the corrections applied to the Fm value. The error associated with δ^{13} C is described by : δ^{13} C error = $4 \cdot 10^{-8} / (1 + 10^{-3} \delta^{13}C)^2$. This component of the Fm error is added in guadrature as follows: Fm $_{\delta}^{13}$ C error = Fm $_{\delta}^{13}$ C [(Fm error² / Fm²) + δ^{13} C error²]^{1/2}. *Radiocarbon date/age* (¹⁴C age). Conventional radiocarbon age, the main

result of radiocarbon measurements, is calculated according to the formula:

⁺C age =
$$-\tau \ln (Fm_{\delta}^{13}C)$$
,

where $\tau = T / \ln 2 = 8033$ yr is the Libby mean life and T = 5568 yr is the Libby half-life. Radiocarbon ages are calculated using the Libby half-life of ¹⁴C.

The ¹⁴C age is expressed in vr BP (radiocarbon years before present, i.e. before AD 1950).

The ¹⁴C age error is given by: ¹⁴C age error = τ (Fm₅¹³_C error).

According to the convention outlined by Stuiver and Pollach [41] and Stuiver [42], radiocarbon dates are reported as follows: ¹⁴C age values < 1000 yr BP are rounded to the nearest 5 yr, 14 C age values > 1000 yr BP are rounded to the nearest 10 yr and the 14 C age errors < 100 yr are rounded to the nearest 5 yr etc. However, not all laboratories subscribe to this convention.

In our report, the radiocarbon age of the samples and the corresponding error values were rounded to the nearest 5 yr

RESULTS

General research on Grootboom. The area of Northeastern Namibia. located between the village of Tsumkwe and the border with Botswana, which belongs to the Tsumkwe Constituency, traditionally known as Bushmanland, has several large African baobab trees. Three of these baobabs, known under the names of Dorslandboom, Grootboom and Holboom, are of outstanding size.

Die Grootboom (The Big Tree in Afrikaans) site is situated at 15 km (as the crow flies) east of Tsumkwe, in close proximity of the Makuri and Djokhoe campsites. The position of *Grootboom* is defined by the following parameters: GPS coordinates 19.649317 S-lat, 020.657583 E-long (19°38'57.5" lat S, 20°39'23.7" long E); altitude 1149 m. Geographically, the region belongs to the Northern Kalahari Woodland basin and the corresponding vegetation type is known as woodland savanna. The semi-arid climate has a distinct dry season (April-October) and a rainy season (November-March) with ca. 60-100 rainy days. Average annual rainfall is 451 mm (measured from 1965 to 2005).

Grootboom was the largest tree of a so-called "Group of Seven", composed initially of seven baobabs spread over an area of ca. 2 hectares, under the protection of the Nyae Nyae Conservancy.

The first records of Grootboom are found in the writings of the "Dorslandtrekkers" (Thurstland trekkers), that moved from South Africa in 1875 through Namibia onwards to Angola. They were Afrikaan Boers oppressed by the English in South Africa in those years. In their writings, the Dorslandtrekkers state that they moved through the "Tebra veld" (Bushmanland), where they encountered large numbers of wildlife and big "kremetartbooms" (baobab trees). Around 1890, they visited a huge baobab, known as Homasi by the San people (Bushmen), after the name of the respective place in the Juhoansi tongue (San dialect). The Dorslandtrekkers,

which renamed the tree *Grootboom*, were impressed not only by its extraordinary dimensions, but also by its excellent condition.

The heavy trunk of *Grootboom* was accurately measured by the German researcher Rudolf Wittmann [43]. The measurements done in November 2001, at the beginning of the rainy season, when the tree was in leaf, indicated a cbh (circumference/ girth at breast height, i.e. at 1.30 m above ground) of 30.6 m. This value (that might have been even larger by the end of the rainy season) corresponds in a very illusory circular approximation to a dbh (diameter at breast height) of 9.74 m. The basal circumference of *Grootboom* was of ca. 36 m and the largest base diameter (west to east) was found to be 12.1 m.

The bark was very thin, measuring only 1-2 cm and its color was elephant gray with reddish spots. The trunk of Grootboom divided at a height of 2-6 m in seven branches, out of which the tallest four were pointed straight upwards. The height of these branches, estimated from the largest unbroken fallen branch, from other broken branches and from several photographs of the tree were (from west to east and than to northeast, facing north) the followings: 14.0, 26.5, 30.5, 32.0, 27.0, 20.5 and 17.0 m (estimated error: ± 0.5 m). The maximum height of 32.0 m is also exceptional for an African baobab. The estimated mean crown spread was of ca. 32 m.

Taking into account the 3 standard dimensions (height, circumference and crown spread), Grootboom can be considered the biggest known African baobab.

As for many other large baobabs, the shape of the trunk was very irregular. Several deep incisions and some obvious fissures determined certain researchers to consider that the huge trunk of *Grootboom* was possibly composed of 3-5 fused stems. Other researchers considered that the incisions and fissures were generated by the twisting movement of branches in time and that the trunk of *Grootboom* was in fact a single entity.

At the end of the 2003/2004 rainy season, Grootboom was full of leaves and looked very healthy. In late June 2004, however, it started dying suddenly. Several branches fell first, approximately two months before the trunk began to collapse. The huge trunk collapsed successively into no less than six stems. The last stem fell around New Year's eve of 2005. Grootboom was probably killed by the mysterious baobab disease, which has affected many baobabs in Southern Africa since 1960.

The fallen stems are labeled (from west to east and than to northeast, facing north) as: A, B, C, D, E, F, in the reverse order of collapse (Photos 2–4). The largest stem (D) fell with roots exposed, three stems (C, E and F) collapsed totally with no roots exposed, while two stems (A, B) broke at irregular heights from the ground (between 1–2 m), also leaving behind stumps.

The original positions of the collected samples in the trunk prior to its collapse are shown in Photo 5.

Growth rings. Growth rings could be identified on only one single sample out of the collected. This is quite surprising, as one considers that in addition to the dated samples, some other large samples (with sizes up to $30 \times 30 \times 10$ cm) have been collected

Sample No. 3 exhibited two obvious growth rings and the external contour of a third ring, as well. The width values of the three consecutive rings, each composed of several wood layers, were the following: 4.8, 4.6 and 2.5×10^{-3} m (mm).

The alternation of two larger rings and one thinner ring is in agreement with the sequence of rainy seasons rich or scarce in precipitation, which have been recently ADRIAN PATRUT, KARL F. VON REDEN, DANIEL A. LOWY, PAULINE LINDEQUE, ET AL

recorded in the area. However, one cannot unmistakably state that the observed growth rings are true annual rings.



Photo 2. Location of stems on the southern flank.



Photo 3. Location of stems on the eastern flank.



Photo 4. Location of stems on the northern and northeastern flanks.



Photo 5. Original position of the samples prior to Grootboom's collapse.

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AMS dating results. The NOSAMS dating results are displayed in Table 1. The radiocarbon dates/ages of 3 samples (Nos. 1, 4 and 8), collected from three different stems, was found to be greater than 1000 yr BP, i. e. 1255 ± 35 , 1045 ± 20 and 1090 ± 55 yr BP, evidence for Grootboom's old age.

The calibration of radiocarbon dates to calendar ages is presented in part two of the report.

Credits:

Photos 1–5 were taken by Rudolf Wittmann on November 14, 2001 and are copyrighted: © Rudolf Wittmann (www.baumsachverstaendiger.de).

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Dedicated to professor Gh. Marcu at his 80th anniversary

ACCELERATOR MASS SPECTROMETRY RADIOCARBON DATING OF AN OLD TROPICAL TREE: PRELIMINARY REPORT 2. CALIBRATED RESULTS

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ABSTRACT. Calibration of radiocarbon dates of 11 samples collected from a recently collapsed African baobab is discussed. The 1- σ calibrated calendar ages of several old samples are over 1000 yr.

The obtained results also suggest that the investigated baobab was morphologically a multiple tree, with several fused stems, while genetically it was a single individual.

KEY WORDS: radiocarbon dating, AMS, tropical trees, age determination, dendrochronology, dendroclimatology.

Part one presented the AMS radiocarbon dating results of 11 samples collected from Grootboom, the world's largest African baobab, which had collapsed recently in Namibia [1].

Part two is dedicated to calibration of radiocarbon dates to calendar ages and to the discussion and interpretation of the resulting age values.

EXPERIMENTAL SECTION

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Calibration. The calculation of the radiocarbon age (¹⁴C age) of a sample assumes that the rate of production of ¹⁴C and consequently the specific activity/ concentration of the ¹⁴C in the atmospheric CO₂ is constant. This assumption is now known to be incorrect, meaning that radiocarbon years are not equivalent to calendar years. Long-term variations in the rate of production correspond to fluctuations in the strength of the Earth's magnetic field. Short-term variations, i.e. "wiggles," are known as the de Vries effect and may be related to variations in sunspot activity.

Radiocarbon ages are always reported as yr BP (before present, where 0 BP = AD 1950), assuming that the atmospheric ¹⁴C concentration has always been the same as it was in AD 1950 and that the half-life of ¹⁴C is T = 5568 yr (Libby half-life). Consequently, a calibration dataset is necessary to convert uncalibrated conventional radiocarbon dates/ages into calibrated calendar ages. Calibration curves also include the correction factor for conversion to the Cambridge half-life (T = 5730 yr). Calibrated ages are reported as calBP, calBC or calAD, as they are expressed in calendar yr BP, calendar yr BC or calendar yr AD.

Practically, calibration programs calculate, via a calibration dataset, probability distributions of the calibrated calendar age for a certain radiocarbon age (with a certain standard deviation). Probabilities are ranked and summed to find the 68.3% (1- σ ; one sigma) and 95.4% (2- σ ; two sigma) confidence intervals and the relative areas under the probability curves for the two standard intervals. Each 1- σ or 2- σ probability distribution corresponds, by the selected areas, to one or several ranges of calendar years.

The radiocarbon ages of samples were calibrated with the OxCal version 3.10 for Windows software/program [2-6] and also with the CALIB version 5.0 software/program [7-9], using the atmospheric data from Reimer et al., i. e. the IntCal04 terrestrial calibration dataset [10]. For one sample (No. 11), the atmospheric data from McCormac et al., i.e. the SHCal04 calibration dataset [11] was used .

All calibrated calendar ages and the corresponding errors were rounded to the nearest 5 yr (excepting the calAD ages calculated with the CALIB program, which were rounded to the nearest year).

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RESULTS

Calibrated ages. The calibration of radiocarbon dates/ages (¹⁴C ages) to calendar ages are listed in Table 1 (as calibrated with the OxCal v3.10 program) and Table 2 (as calibrated with the CALIB. 5.0 program). The 1- σ (68.2%) and 2- σ (95.4%) probability distributions, which define calAD age ranges of the dated samples, were obtained (calAD ages are rounded to the nearest 5 yr by the OxCal program and to the nearest year by the CALIB program).



Fig. 1. The plotted OxCal calibration for sample No. 1.

From these values, we chose as the calAD age of each sample the corresponding 1- σ range (when only one range exists) or the corresponding 1- σ range with the highest probability (when there is more than one range). The selected calAD range of each sample is marked in bold.

Tables 1 and 2 also include the mean calBP age values and the corresponding errors of the samples (each rounded to the nearest 5 yr), calculated form the selected calAD range.

Shown are also calendar ages of the samples at the death of the tree. They were calculated from the corresponding mean calBP ages extrapolated (from AD 1950) to AD 2005. AD 2005 was chosen as the reference year for calculating sample ages, as the stems of Grootboom, where the samples originated from, collapsed between September 2004 and New Year 2005. Also rounding reasons pleaded for this year. In the case of sample No. 11, no calAD age interval was selected, for reasons to be explained later in this paper.

Results obtained with the two different programs, using both the same calibration datasets, are very similar. The 3 samples with high radiocarbon date were found to be millenarian, with ages of 1275 (No. 1), 1000 (No. 4) and 1050–1060 (No. 8) cal yr.

The plotted OxCal calibration results for the 3 millenarian samples are presented in Figs. 1–3.





Fig. 2. The plotted OxCal calibration for sample No. 4.



Fig. 3. The plotted OxCal calibration for sample No. 8.

DISCUSSION

Northern vs. Southern Hemisphere calibration. The radiocarbon calibration dataset is the result of committed work and significant joint efforts of numerous researchers over several decades. It is based on dating results of samples with known age collected almost exclusively form the Northern Hemisphere, especially dendro-dated wood and marine records.

Several studies have revealed, however, regional radiocarbon offsets. The most important is the so-called interhemispheric offset. Certain comparative studies evinced that trees from the Southern Hemisphere dated older than identically aged trees from the Northern Hemisphere. Such results suggest hemispheric differences in the distribution of ¹⁴C throughout the troposphere. These differences are attributed to

the larger expanse of ocean in the Southern Hemisphere and the atmosphereocean exchange [12-17].

The very first Southern Hemisphere calibration dataset was developed in 2002, based on several research on dated wood samples from New Zealand, Tasmania, Chile and South Africa, covering the period AD 1955–955. The SHCal02 calibration [18] is based on the IntCal 98 dataset for the Northern Hemisphere [19], corrected with a number of results from the Southern Hemisphere. The SHCal02 dataset indicates older ¹⁴C ages by ca. 8–80 yr in the Southern Hemisphere, with a mean offset of 41 ± 14 yr for the period cal AD 1850–950 and also shows a periodicity in the offset of about 130 yr.

The International Radiocarbon Conference in Wellington, New Zealand recommended the use of the SHCal02 dataset for samples with cal AD 1850–950 (0–1000 cal yr BP) ages from the Southern Hemisphere [11]. The Southern Hemisphere is defined as south of the thermal equator or south of the Intertropical Convergence Zone (ITCZ). The ITCZ is the region that circles the Earth and extends from about 5° N to 5° S, where the northeast and southeast trade winds converge in a low pressure zone.

There are, however, seasonal shifts in the ITCZ that may bring atmospheric CO₂ from the Northern Hemisphere to a site for part of the year and from the Southern Hemisphere for another part. This seasonal migration of the ITCZ, combined with multi-decadal and millennial migrations in the past, poses some uncertainties for calibration of dated samples from tropical and neo-tropical sites [11].

The SHCal02 dataset, corrected and completed with new research data, determined in 2004 the development of the SHCal04 calibration dataset, which covers a much longer period, between 0–11 cal kyr BP [11]. The SHCal04 Southern Hemisphere calibration is associated with the new developed IntCal04 Northern Hemisphere calibration for the period 0-26 cal kyr BP [10] (Fig. 4). SHCal04 shows a larger interhemispheric offset than the previous SHCal02 that varies only slightly from 55 to 58 yr.



Fig. 4. Plotted IntCal04 and SHCal04 calibration datasets for the period 0–2000 cal yr BP. The difference (for a certain calendar year BP value) between the SHCal04 and IntCal04 curves (expressed in radiocarbon years BP) corresponds to the offset.

Nevertheless, the calibration of the dated samples collected from Grootboom was performed by using the atmospheric data from Reimer et al., i. e. the very large and well documented IntCal04 terrestrial calibration dataset [10]. The reasons for this choice are the following:

i) The SHCal Southern Hemisphere calibration is based on a relatively scarce number of dating results of wood samples from mid-latitude areas, which suggest a mean interhemispheric offset of several decades. On the other hand, alternative studies show no or very small interhemispheric offset values [20-22]. A very recent extended research done on Tasmania huon pine (*Lagarostrobus franklinii*) decadal samples do not document a distinguishable offset from the Northern Hemisphere for the time period 1550–2100 cal yr BP [23].

ii) The Grootboom site is located at relatively small latitude, ca. 19° S, right in the intertropical area, for which there are some uncertainties concerning calibration. We outline that all results used for the Southern Hemisphere calibration originates from wood samples collected from greater latitudes. It should be also noticed that some research suggests intrahemispheric offsets between samples collected form different latitudes of the same hemisphere [24-28, 14-15].

Only the youngest sample (No. 11), with a 14 C age of 30 ± 40 yr BP, was calibrated using the atmospheric data from McCormac et al., i. e. the SHCal04 calibration dataset [11], because this recent period is much better documented and a small interhemispheric offset was demonstrated.

One should notice that the calibration of the samples using the SHCal04 dataset leads to lower calendar age values, with an offset of several decades.

Age of samples. Trees take up atmospheric CO_2 by photosynthesis. CO_2 is being used in part for the quasi-continuous building of the wood texture of the trunk and branches. As other trees, the African baobab grows due to cambial activity which, during the rainy season, adds every year new layers of woody tissue. Dead wood does no longer exchange CO_2 with the environment.

Beside structural non-mobile carbon components, mainly cellulose, which have been synthesized at the very moment of the original formation of the respective woody tissue, the wood (xylem) also contains non-structural mobile carbon components, which are formed earlier or later and are removed by pretreatment of samples

While the stable isotope (i.e. ¹²C and ¹³C) content of the wood remains constant over time, the amount of the unstable ¹⁴C isotope, also called radiocarbon, gradually decreases via radioactive beta minus decay. Thus, ¹⁴C emits continuously electrons and is converted to nitrogen, according to reaction:

 ${}^{14}C \rightarrow {}^{14}N + e^{-}$.

By determining the ¹⁴C content, relative to the content of stable carbon, of a pretreated wood sample, one can date that particular sample.

The age of a certain position in the trunk or branches is calculated starting with the moment of the original formation of the respective woody tissue. Theoretically, the true age of a certain position is equal to the age of a pretreated wood sample collected from the respective location, which was dated and calibrated.

However, in certain cases, especially of tree species with soft and spongy wood, the age of one particular sample may differ, more or less, from the true age of the corresponding position.

The age of several samples collected from the remains of Grootboom belong to this category, such as the dating results suggest. One can distinguish the following three cases:

i) Samples collected from the rim of hollow parts. Beside hollows generated by elephants or by fire, hollows are generally formed by twisting movements of large branches, due to their growth and/or excessive weight, which may rotate and break the soft wood of the baobab trunk. Thus, at the moment when a hollow forms, woody tissues are displaced from their original position. On the other hand, certain tree species, including the African baobab, show the tendency to partially repair over time the hollows with amounts of new wood, especially for hollows relatively close to the external surface of the trunk. In special cases, such as the Livingstone baobab of Chiramba (Mozambique) or Holboom, another huge Namibian baobab located at only 8 km from the Grootboom site, it was reported that rims of an internal hollow can be covered over time with new bark.

ii) Samples collected in the vicinity of contact or fusion areas between neighboring stems. For trees with multiple stems, the neighboring stems may eventually get in contact and may even fuse together. Production of woody tissue can continue in contact and in fusion areas, even when no free place would be available for growth. Thus, a wood sample collected from such a location may contain woody tissues of different ages, which are mixed together.

ii) Samples collected from collapsed stems. In the case of tree species with soft and spongy wood, the collapsed stems are damaged and get bent due to their own weight. Thus, woody tissues from different positions may mix together. Samples collected along the central axis, parallel to the ground, may also contain younger wood, which originates from positions located towards the upper part of the fallen stem.

Consequently, samples collected from the rim of hollows, in the vicinity of contact or fusion areas between stems or from collapsed and damaged stems may date younger than the true age of the position from where they are collected. On the other hand, such samples may even be non-homogenous, containing woody tissues of different ages in variable proportions.

In the light of these remarks, the only samples whose dated and calibrated age is identical or very close to the true age of their corresponding position in the trunk are those which were collected from a remaining stump of a fallen stem, from a location away from the rim of a hollow and also not very close to a contact or fusion area between two neighboring stems. These conditions are met only by one of the samples collected from Grootboom, namely No. 3.

Discussed below are calibrated ages of the dated samples (Tables 1 and 2). The samples collected from four different stems are numbered (from No. 1 to No. 10) from west to east, facing north, considering their original position in the trunk prior to collapse [1].

Sample No. 1, ca. 1275 yr. This is the oldest sample of the tree, collected at a height of ca. 1.20 m and at a distance of ca. 0.40 m from the center of the remaining stump of stem A, the last to collapse around New Year's eve 2005.

Sample No. 2, ca. 510 yr. The sample originates from the stump of stem B and it was collected at a height of ca. 1.90 m and a distance of only ca. 0.20 m from the contact and partial fusion line between stems A and B. Its age is suspected to be younger than the true age of the corresponding position. In addition, the wood of the sample is much more compact and dense relative to other samples, suggesting that a large number of woody tissues had been added to this location and had been pressed together over time.

Sample No. 3, ca. 515–520 yr. This is an extreme lateral sample, collected at a height of ca. 2.30 m and at only 0.10 m from the southern edge of the remaining stump of stem B.

Sample No. 4, ca. 1000 yr. It is another millenarian sample, collected from the stump of stem B, at a height of ca. 1.40 m and also from the rim of the largest internal hollow of Grootboom. The corresponding position was shifted towards the southern flank.

Sample No. 5, ca. 600 yr. This sample was collected from the collapsed stem C. In the standing stem, the corresponding position was located close to the rim of a large hollow, at a height of ca. 2.60 m above ground level.

Sample No. 6, ca. 560 yr. It was collected, like sample No. 5, from the fallen stem C, from a position located at the rim of the same very large hollow and a height of ca. 2.40 m above ground level.

Sample No. 7, ca. 815–820 yr. This sample was collected from the rim of a small hollow, located just in the core/center of the fallen stem D, the largest of the tree. It had a rather sandy aspect, being severely decayed due to microorganisms which destroyed the fibrous structure of the wood. Its age is suspected to be considerably younger than the true age of the corresponding position. All samples originating from stem D were located near ground level in the standing trunk.

Sample No. 8, ca. 1050–1060 yr. It is the second oldest sample of the tree, which was collected from the rim of a hollow of the collapsed stem D, from a position located initially at 1.00 m towards east, relative to sample No. 7.

Sample No. 9, ca. 590 yr. The sample was collected from the fallen stem D, from a position situated at 1.20 m towards east, as compared to sample No. 8.

Sample No. 10, ca. 575 yr. This is another extreme lateral sample, collected from the fallen large stem D. Prior to collapse, it was located at only 0.25 m from the eastern extremity of the stem and of the whole trunk.

Sample No. 11, no selected age. This sample was collected at 1.60 m from the top of the unbroken fallen high branch, with a measured length of 26.31 m, following its contour.

The calibration and interpretation of radiocarbon dating for samples formed after AD 1640 is problematic and is characterized by low accuracy. This is mainly due to industrial fossil fuel burning, which involved large emissions of million-year old practically ¹⁴C-free carbon in atmosphere. The high variations of ¹⁴C concentration in the atmosphere over the industrial period, known as the Suess effect, affected to a larger extent the Northern Hemisphere, which was more advanced [29]. The incorporation in living tissues, including those which became shortly thereafter dead wood, of CO₂ from the period of the Suess effect produces generally an overestimation of the ¹⁴C age of the corresponding samples. This determines that a calibration of a sample from this period results in up to five cal age ranges for one ¹⁴C age [30].

The ¹⁴C age of sample No. 11, i. e. 30 ± 40 yr BP, corresponds to three 1- σ ranges (calibrated with OxCal and CALIB), respectively to three 2- σ ranges (calibrated with OxCal) and even to six 2- σ ranges (calibrated with CALIB) (Tables 2-3). The 1- σ range and also the 2- σ range with the highest probability correspond, with both calibration programs, to the period AD 1950–1960 (rounded values).

The 1950–1960 decade is also known for the beginning of the dramatic period of troposphere bomb tests. Due to the 404 atmospheric nuclear detonations between 1953–1963, a large amount of artificial ¹⁴C was injected into the stratosphere. Consequently, the ¹⁴C concentration in the troposphere almost doubled from 1950 to 1964 [31-33]. Again, the Northern Hemisphere, where the large majority of nuclear weapon tests took place, was affected earlier and to a more significant extent than the Southern Hemisphere. Incorporation in living tissues of CO₂ from the bomb tests period, which contains modern carbon very rich in ¹⁴C, determines generally an underestimation of the ¹⁴C age of the respective sample. The calibration of a sample from this period also results in three or more cal age ranges for one ¹⁴C age.

Given that both the Suess effect and the bomb tests affected to different extents the two hemispheres, these differences being relatively well documented, determined us to use the SHCal04 dataset to calibrate the ¹⁴C age of sample No. 11.

Selection of a reliable/accurate cal age range is, however, almost impossible. The range AD 1950–1960, which has the highest probability, is practically excluded, even if the ¹⁴C age is probably underestimated. This is due to a large number of testimonies that state that Grootboom practically had not changed at all from its discovery in ca. 1890 until its sudden death. What is certain is that sample No. 11 was formed after AD 1640, thus having an age of less than 365 cal yr. In our opinion, the age of sample No. 11 is probably between 200–300 yr.

Structure of Grootboom. The enormous trunk of Grootboom collapsed very chaotically in all directions, so that six fallen stems have been eventually identified. After the fall of the last stem to the ground, the remains of Grootboom looked as if the tree would had been bombed or blasted with dynamite.

The six stems were marked (from west to east and than to northeast, facing north) by: A, B, C, D, E and F, in the reverse order of collapse [1].

The dated samples were collected from four stems, namely A, B, C and D. The fluctuation of the age of samples and the presence of 3 millenarian samples in three different stems strongly indicate that the trunk of Grootboom was made up of several (the more or the less) fused stems.

Considering the dating results and other information, we will try to answer the following question: How many and which of the six fallen stems had been independent, i. e. belonged to the original structure of Grootboom?

Discussed and analyzed below are the six stems, which were all of oval shape. The estimated values of the largest dbh (diameter at breast height) and of the dbh in perpendicular direction are presented in brackets.

Stem A (ca. 2.5 m x 1.2 m). This stem, which collapsed the last, was located at the western extremity of the trunk (facing north). It was located next to trunk B towards east, the two stems being partially fused up to a height of ca. 1 m above ground level. Its small dimensions are somewhat surprisingly, especially because it had enough free room to grow towards west. The oldest dated sample (No. 1) was collected exactly from stem A.

Stem B (ca. 3.1 m x 1.8 m). Stem B was located in between stems A and C and had no free room for additional growth. The third oldest sample (No. 4) was collected form this stem, more precisely from the rim of the largest hollow of the tree.

Stem C (ca. 1.6 m x 0.8 m). This small stem was fused up to a height of ca. 2 m with other three stems: B (towards west), D (towards east) and E (towards northeast). Its reduced dimensions and the relatively young age of the samples collected from it (Nos. 5 and 6) reveal that stem C must have been only a fusion product of the three stems, that continued to produce woody tissue after fusion.

Stem D (ca. 5.5 m x 3.2 m). Stem D, obviously the largest of the tree, was located at the eastern extremity of the trunk. It was the only stem which fell with roots exposed. Stem D was partially fused with stems C (towards west) and E (towards north). After breakdown, its base diameter parallel to the ground was measured and found to be 5.90 m. The second oldest sample (No. 8) was collected from this stem.

Stem E (ca. 3.0 m x 2.0 m). Stem E, which originates form the eastern flank of the trunk, was fused with stem D (towards south) and was in contact with stem F (towards north). It was unavailable for sampling, as it was partially hollow and also almost entirely covered and crushed by other fallen stems.

Stem F (ca. 3.5 m x 3.2 m). This stem, the first which collapsed, was almost isolated at the northeastern extremity of the trunk. It had only a connection area near ground level with stem E. The partially hollow stem F had already been severely decayed at the moment of the first sampling and no sample was collected from it.

These data suggests that five stems (A, B, D, E and F) may be considered independent and they are likely to belong to the original structure of Grootboom. The sixth stem (C) was much younger, being the result of fusion of three vicinal independent stems.

As already mentioned, there are two possibilities for the genesis of a baobab with multiple stems [34]. The first possibility is multiple sprouting from the same rootstock of a fallen parent tree. Hence, the stems are clones of the parent tree and they look very similar, because they are genetically identical. The second possibility is the simultaneous germination of several seeds. In this case, the stems are genetically different and they look somewhat different, for instance they may break into leaf at a different time and the tint of leaves and bark may also be somewhat different.

No such differences have been observed for Grootboom. This fact supports the hypothesis that the five independent stems sprouted simultaneously and fused into a single trunk at some later time. Consequently, one can state that morphologically Grotboom was a quintuple tree, while genetically it was a single individual.

The moment of complete fusion can be estimated by means of the age of samples collected from the fusion stem C (Nos. 5 and 6) and near the fusion area in between stems A and B (No. 2). Considering some corrections, we estimate that the five independent stems fused into a single trunk ca. 600–800 yr ago.

It should be also noticed that other very large African baobabs obviously have or are suspected of having multiple stems. We mention here the following trees: Chapman baobab at Gootsa pan (in Botswana), Dorslandboom in Bushmanland (in Namibia), Platland baobab near Duiwelskloof (in South Africa), Big tree at Victoria Falls and Big baobab at Devuli/Mokore ranch (in Zimbabwe), Big baobab near Joal (in Senegal).

Age of Grootboom. The 3 oldest dated samples with ages of at least 1000 yr, collected from three different stems, evince beyond doubts that Grootboom was a millenarian tree. Grootboom's true age can be estimated from the age of the oldest sample and its position in the respective stem. Sample No. 1, with a calendar age of 1,275 \pm 50 yr, was collected from the stump of the relatively small stem A, located at the western extremity of the trunk, at a height of 1.20 m above ground and a distance of 0.40 m from the calculated position of its core which was hollow. These results reveal that the age of Grootboom was of +1,275 yr, i.e. the tree was older than 1,275 yr (our estimate is 1,350–1,500 yr).

Growth rate. The classic concept of growth rate of the whole trunk is meaningless for trees with multiple stems. One can only evaluate growth rates of independent stems and their dynamics, taking into account that the growth of each stem is limited by other stems in at least one direction.

In the particular case of Grootboom, one should notice that all stems had a clearly oval shape. Thus, the west-east diameter of the first three independent stems (A, B and D), from which samples were collected, was almost twice larger than their north-south diameter. This appears to be surprising as these stems had much more available room for growing towards north and south than towards west and east. Consequently, the growth rate in the west-eastern direction was almost twice greater than the growth rate in north-southern direction. For the other two stems (E and F), the west-east growth rate was also greater.

Because in case of multiple sprouting all independent stems are of the same age, i. e. +1275 yr, for each stem one can calculate the mean growth rate in two perpendicular directions. Mean growth rate values at breast height (grbh) for the entire life cycle of Grootboom are shown in Table 3. Given that in these calculations, we used the minimum age value (1275 yr), the growth rate data presented can be considered as representing the upper range values. The presented values for the mean growth rate (expressed in $10^{-3} \text{ m·yr}^{-1}$) correspond to the mean annual increase in radius (expressed in 10^{-3} m or mm).

Table 3.

Stem	Age (cal yr)	Diameter at breast height (m)		Mean growth rate at breast height/ /annual increase in radius (10 ⁻³ m·yr ⁻¹)	
		dbh 1	dbh 2	grbh 1	grbh 2
			[INS]		[NS]
A	+1275	2.50	1.20	0.98	0.47
В	+1275	3.10	1.80	1.22	0.71
D	+1275	5.50	3.20	2.16	1.25
E	+1275	3.00	2.00	1.18	0.78
F	+1275	3.50	3.20	1.37	1.25

Mean growth rate values of the independent stems

The largest mean growth rate is the west-east value for stem D (2.16 x 10^{-3} m·yr⁻¹), while the smallest is the north-south value for stem A (0.47 x 10^{-3} m·yr⁻¹).

These mean values do not reflect, however, the growth dynamics of Grootboom's stems. According to published accounts, the growth rate of the African baobab decreases severely with the age [35, 36, 37, 38]. This statement can be also verified in the case of

Grootboom, by the ratio of the position of the extreme lateral samples to their age.Ages of extreme lateral samples (Nos. 3 and 10) show that over the past ~520 yr stem B grew by only 0.10 m towards south, while stem D grew by 0.25 m towards east in ~575 yr, even though there was enough room in both directions (no neighbouring stem). The calculated growth rate values are very small: 0.19 and 0.43 x 10^3 m yr⁻¹ (corresponding to a mean annual increase in radius by only 0.19 and 0.43 mm). Such values reveal that over the past 500–600 yr Grootbooom almost ceased growing.

When compared to historic records of the same trees, measurements of several huge individuals from Botswana and Mozambique showed a very small increase or even a decrease in girth during a time span of ~110 yr, from 1850–60 to 1966 [39]. These results were attributed to an obvious decrease of rainfall in Central Southern Africa. Our dating results of Grootboom suggest that the period of prolonged drought may have begun several centuries earlier, probably around AD 1400–1500.

Age limit of the African baobab. Preliminary dating results of the samples collected from Grootboom strongly support the long lived baobab hypothesis or at least that certain individuals of the species may become millenarians. The oldest sample indicates an age of +1275 yr for Grootboom, while our estimates augment the age up to 1350–1500 yr. One should consider that shortly before its demise Grootboom looked healthy and seemed not to be a very old and decrepit tree, nearing its end. It is very likely that it was attacked and killed by the poorly studied baobab disease. Had this unfortunate event not happened, Grootboom might have probably lived for additional years. Therefore, we estimate that the age limit of the African baobab, which is probably identical to the age limit of angiosperms, is around 1,500 yr.

History of Grootboom. The acquired data suggests the following scenario for the history of Grootboom. Over 1275 yr ago (prior to AD 730), maybe 1350–1500 yr ago (around AD 500–650), a very large African baobab collapsed somewhere in Central Southern Africa, in a semi-arid area habited only by San people or Bushmen. Five sprouts/shoots developed from the prostrate parent, which could be named Ur-Grootboom.

The five young baobabs, which were clones of the parent tree, grew gradually, until 600–800 yr ago (around AD 1200–1400), when they fused together into a single huge trunk, which came to be known much later as the Grootboom baobab. About 500–600 yr ago (around AD 1400–1500), the trunk of Grootboom was almost as large as at the time of its death. Presumably, climate changes in the area, especially shift towards conditions of prolonged drought, almost stopped its growth.

In AD 1890, Grootboom was discovered by the modern world, namely by the Dorslandtrekkers, a group of Boers who withdrew from South Africa to Angola. The tree, which looked very healthy, had not changed its physical appearance until AD 2004, when it unexpectedly died, being probably killed by the still mysterious baobab disease. Grootboom begun to collapse stepwise, until the last stem fell around New Year AD 2005. At the beginning of AD 2006, the last traces of Grootboom disappeared completely, leaving behind only the memories of what once was a mighty baobab tree.

CONCLUSIONS

Grootboom, the world's largest African baobab tree, dies and collapsed unexpectedly in Bushmanland, Namibia, in late 2004. Six fallen stems have been identified on the ground.

The international research project started on this occasion had the following aims: 1) to determine accurately the true age of the tree; 2) to establish whether Grootboom's trunk was a single unit or was composed of several fused stems; 3) to learn about the dynamics of Grootboom's growth rate during its life cycle.

11 samples collected from the remains of the collapsed tree were processed and analyzed by AMS radiocarbon dating. Several additional samples are under investigation.

The presented results and conclusions of the research can be summed up as follows:

1) The radiocarbon age (14 C age) values of 3 samples (Nos. 1, 4 and 8), collected from three different stems, was greater than 1000 yr BP, i. e. 1255, 1045 and 1090 yr BP. These are the first samples of an angiosperm tree, with accurate dating results, older than 1000 yr.

The radiocarbon age of the oldest sample (No. 1) was found to be 1255 \pm 35 yr BP, which corresponds to a 1- σ calendar age of 1275 yr, at the moment of the tree's demise.

Grootboom becomes the first millenarian angiosperm tree, with an age of +1275 yr. Our estimate of the age is around 1350–1500 yr, according to the position of the oldest sample in the corresponding stem prior to collapse.

2) The acquired results indicate that five stems may be considered independent, belonging to the original structure of Grootboom, while the sixth stem was the result of fusion of three vicinal independent stems.

The five (genetically identical) independent stems sprouted simultaneously from a parent tree and they fused into a single trunk at some later time. The age of samples collected from the fusion stem (Nos. 5 and 6) and near the fusion area of two neighboring stems (No. 2) suggest that the five stems fused into a single trunk around 600–800 yr ago.

One can state that morphologically Grootboom was a multiple tree, but genetically it was a single individual.

3) The age of the extreme lateral samples (Nos. 3 and 10) reveals that Grootboom grew by only ca. 0.10 m towards the southern edge and 0.25 m towards the eastern extremity over the past ca. 500–600 yr. The severe decrease of the growth rate could be ascribed to climate changes in Central Southern Africa, i.e. a shift towards conditions of prolonged drought.

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Dedicated to professor Gh. Marcu at his 80th anniversary

VOLTAMPEROMETRIC DATA ON THE BIOLOGICAL ACTIVE SYSTEMS CU(II), PD(II) AND NI(II) – SALICYLIC ALDEHIDE THIOSEMICARBAZONE

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ABSTRACT. The voltamperometric behavior of the systems Cu(II), Ni(II) and Pd(II) – thiosemicabazone of salicylic aldehyde (TSCSA) has been investigated on glassy carbon electrodes (cyclic voltametry) in DMF/LiCIO₄ 0,2 M medium, using computer aided electrochemical systems BAS 100W and AUTOLAB – ECOCHEMIE.

In the case of all complexes can be seen that the peak related with the sulphur oxidation was moved towards positive higher values, which confirm the involving of sulphur in the coordination. From the point of view of the donor – acceptor interaction between metallic ion and sulphur atom as donor, it can be affirmed that, in these conditions, the Cu(II) complex is more stable than the Ni(II) and Pd(II) complexes. This conclusion is in accordance with the especial affinity of the Cu(II) ion against sulphur.

Based on the shifts of reduction potentials of the metals, $\Delta \varepsilon$ = 215 mV for Cu complex, $\Delta \varepsilon$ = 162 mV for Ni complex and $\Delta \varepsilon$ = 180 mV for Pd complex, various degrees of interaction have been identified. They are qualitatively correlated with the stability of the complexes.

Key words: thiosemicarbazones, metal complexes, cyclic voltammetry.

Introduction

Thiosemicarbazones and related compounds, as well as their metal complexes have been the subject of great interest of many researchers, the first of all because they are active from the biological point of view, some having antitumoral activity. The statement is supported by a large number of papers, some of them in the last years [1,2]. Because in the human body there are redox processes it is important to have the information about the electrochemical behavior of the systems metal ions, Cu(II), Pd(II) and Ni(II) - thiosemicabazone of salicylic aldehide in solution. Apart from their diverse chemical and structural characteristics, the significance of these compounds, especially related to thiosemicarbazones and theirs metal complexes, the relation between structure and biological activity has

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been covered in papers by West et al.[3,4]. The complexes have been prepared using the interaction of Ni(II) and Cu(II) sulphates, respectively $[PdCl_4]^{2^{\circ}}$ with the thiosemicarbazone of salicylic aldehide (TSCSA) in aqueous solutions of ethylic alcohol, in echimolar amounts, resulting 1/1 type ligand thiosemicarbazone metal complexes [5].

In order to characterize the redox processes of these systems, which may occur on the ligands and complexes, as well as to compare the stability of the complexes and the coordinating ligand position involved in complex formation in solution, electrochemical investigations are usually performed, in suitable solvents. Although the thiosemicarbazone – based ligands were investigated by electrochemical methods [6,7], for the system (TSCSA) – Me(II) there are not the data in the literature.

The investigations have been made in organic media (DMF, THF, AC), in the presence of several supporting electrolytes, usually at glassy carbon electrode. The most important conclusion of these voltamperometric determinations was the complexity of the electrochemical answer, involving both the redox processes on the level of ligand moiety and metal ion [8].

Experimental

The tests were performed in a classical cell having three electrodes. All the samples were deaerated 20-30 minutes before each test. Because of low solubility of the complex in the protic medium, the non aqueous medium of DMF was chosen.

Cyclic voltammograms were recorded in a dried cell purged with argon. DMF used as electrolyte solvent was purified according to standard procedures [9]. LiClO₄ dried in an oil pump vacuum at 100° C was added as supporting electrolyte at a concentration of 0.2 M. Compounds under investigation were added at 1 mM concentration. The working electrode (GC, Hg and Pt) and platinum wire counterelectrode were used. An Ag/Ag⁺, AgCl electrode, in a separate compartment served as reference electrode. CVs were recorded both in negative and positive-going direction at the starting potentials, at different scan rate using a Autolab potentiostat (Ecochemie) equipped with PGSTAT 12 soft. All experiments were run at room temperature (22° C).

Results and discussion

The ligand (TSCSA), produced from the condensing salicylic aldehide with thioureea (TU), has basically three electroactive positions: hydroxyl oxygen, the sulphur from the remaining thiourea and the azometin grouping. From these, the last two unsaturated groups can be reduced and the OH can be easily oxidized.



The electronic interaction between datively bonded ligand (TSCSA) and divalent metal ions has been studied with cyclic voltammetry and polarography, both in reduction and oxidation potential range.

Typical CVs of complexation in Cu(II)-TSCSA system, in DMF, in negative – going direction are displayed in Fig. 1.



Figure 1. Comparison between the electrochemical behavior of (TSCSA) and Cu (II) complex in negative – going direction.

In DMF medium, on glassy carbon, the ligand presents two oxidation peaks, the first in the 700-850 mV range *vs* Ag/AgCl,KCl, which can be related to phenolic group [10, 11], Fig.2a.



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Figure 2. CVs for the oxidation of ligand and (TU) on GC electrode in DMF/ 0.2 M LiClO₄ for : $a - oxidation of 10^{-3} M (TSCSA)$

b – oxidation of 10^{-3} M (TU)

and the second in the high and positive range (>1 V vs Ag/AgCl) which is specific to π carbon-sulphur oxidation; this last peak's attribution was confirmed by the voltamperometric response of (TU), Fig 2b.

To spot the entail the complexation of metal ions, respectively the processes of reduction of the azometin group of ligand, it was followed in the same parameters the voltamperometric response of the components and of the system in an extended range of negative voltages, allowed by the use of non-aqueous electrolyte, (ε_0 = - 0.4 V/ Ag/AgCl, KCl and ε_f =-2.2V/).

The irreversible reduction of azometin group in (TSCAS) might occur at the negative electrode potential, to $C_{-C=N-}$ - 1.8 V/ Ag/AgCl, KCl, Fig.3.



Figure 3. Reduction of azometin group in (TSCAS).

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By comparing VCs of the free ligands (TSCSA) and TU, and for the complexed metalic ions can be seen differences of the peak voltages and for the peak currents because of complexation, Table 1.

According to other authors, confirmed for related compounds in solid state [8], usually thiosemicarbazones act such tridentate ligand (O, S, N):



Me: Ni(II), Cu(II), Pd(II)

but in some cases, M=Ru(II),Os(II) the phenolic oxygen is not included in the coordination [14] and the dimer structures have been demonstrated for TI complexes with related ligands.

Table 1.

The influence of complexation on the peak potentials in DMF [V/ Ag/AgCl, KCl].

Compound	Oxidation		Reduction	
Compound	Сон	€c=s	- Є _{-СН=N} -	- C _{Me(II)}
TSCAS	0.81	1.09	1.47;1.82	-
Cu (II) - TSCAS	0.86	1.43	1.40; 2.12	0.40; 1.20
Cu(II)	-	-	-	0.30
Ni (II) - TSCAS	0.91	1.18	1.53; 2.02	1.53
Ni(II)	-	-	-	1.28
Pd(II) - TSCAS	0.98	1.28	1.46; 1.98	1.55
Pd(II)	-	-	-	0.55;1.30

Comparing our data in DMF for ligand and complexes, Table 1, we can see that the redox properties of the linkages involving donor atoms were stabilized by complexation, Table 2. These facts are valuable both for OH and C=S, when the oxidation potential is displaced to more positive values. For CH=N, the second reduction step could be easier, at lower negative potentials. We consider that the displacement of peaks potentials is due to the complexation and the value of the shift is in relation with the stability of the complex.

Table 2.

Displacement [mV] of the peak potentials of donors and metal ions, by complexation in DMF/0.2M LiClO₄.

Me(II)	Donor group	Δε* [mV]	Δε** [mV]
Cu(II)	–OH	50	
	=S	340	215
	–N=CH–	- 300	
Ni(II)	–OH	100	
	=S	90	162
	–N=CH–	- 200	
Pd(II)	–OH	170	
	=S	190	180
	-N=CH-	- 160	

* Average of 3 determinations.

** For complexed metal ions comparing to metal ion coordinated only with perchlorate.

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The explanation of the shifts of oxidation potentials of donors after complexation is easier for the oxidable groups (OH and =S): the density of electrons after complexation becomes lower, so the oxidation becomes more difficult. For the azometin reducible group, the π character of double bond diminishes by complexation, the linkage becomes weaker and its reduction is possible to smaller negative potentials.

Taking the reduction potential of metal ions complexes coordinated with perchlorate anion, even in dissolved state, results in a negative shift, by complexation with (TSCSA), Table 2, corresponding to the stability of the complexes.

Conclusions

The following qualitative remarks were deducted on complexation interactions:

- Powerful engaging of the sulphur donor atom and azometin group in complexation with Cu(II) ions, more comparing to Ni(II) and Pd(II).

- Participation of the hydroxyl oxygen in complexation with Ni(II) and Pd(II) more than for Cu(II).

- Engaging of the $\boldsymbol{\pi}$ electrons of the azometin linkage in complexation in the following order:

Cu(II) > Ni(II) > Pd(II)

- According to shift of the reduction potentials of metals ions, the relative stability of the complexes is:

Cu(II) > Pd(II) > Ni(II)

Based on the information reported here, future experiments will be realized to establish the quantitative data on the stabilization of these metal ions through complexation.

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Dedicated to professor Gh. Marcu at his 80th anniversary

ADSORPTION AND ELECTROCHEMICAL DATA FOR p-NITROPHENOL REMOVAL FROM SYNTHETIC WASTEWATERS

MARIA JITARU*, BOGDANA KOUMANOVA**

ABSTRACT. There are three main results reported in this paper refer to: (*i*) the equilibrium adsorption experiments of recalcitrant toxic p-nitrophenol (p-NP) on natural zeolites from Balkan area; (*ii*)voltamperommetric data on the electrochemical behavior of p-nitrophenol, depending on the electrode nature and working conditions (pH, electrolyte composition, hydrodynamic parameters (*iii*) and electrochemical oxidation to remove the remaining p-nitrophenol on different electrodes after the adsorption.

The zeolite from Mirsid, Romania (zeolite-2) seems to have the best adsorption properties for p-nitro phenol, comparing to classical sorbent, active carbon powder. The synthetic waste waters containing 0.01p-NP have been treated by adsorption on (zeolite-2) and the decrease of the p-NF concentration up to 0.01 mM has been obtained. The remaining p-NP was treated by electrochemical way (electro-oxidation). Applying the combined adsorptive-electrochemical oxidation procedure we have been obtained the decrease of COT up to 98%, on SnO₂/IrO₂/Ti (Modified Oxides Electrode- MOE). These results could be explained by the catalytic effect of the zeolite for the chemical phenol oxidation.

Key words: nitrophenols, adsorption, zeolites, electrochemical oxidation

Introduction

Due to the recognized toxicity of nitrophenols, the International Programme on Chemical Safety (IPCS), established in 1980, as a joint venture of the United Nations Environment Programme (UNEP) and the World Health Organization (WHO) elaborate the authoritative document, Concise International Chemical Assessment Document (CICAD 20), on the risk assessment of nitrophenols.

During the past decade, extensive research has been conducted to develop innovative, effective, inexpensive and promising adsorbent materials that are regenerated easily for dealing with the problem of the treatment of contaminated wastewater. Organically modified benthonites and zeolites, produced

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by replacing exchangeable inorganic cations with quaternary alkyl ammonium cations, have been reported as strong adsorbents for non-ionic organic pollutants.

The monosubstituted 4-nitrophenol is found in wastewaters discharged from various industrial activities such as pulp and paper industries, textile mills, steel plants, oil refineries, etc. [1]. This compound is also associated with agricultural activities as an intermediate for the production of pesticides, herbicides and insecticides [2].

Attempts have been made to remove mononitrophenols from wastewater by a number of methods: oxidation with strong oxidizing agents as H_2O_2 [3], biodegradation [4], biosorption [5], photocatalytic degradation [6], etc. With respect to adsorption of nitrophenols, activated carbon is one of the most commonly used adsorbents due to its high surface area and well developed pore structure [7-11]. As activated carbon is relatively costly, attempts have been directed to the utilization of low-cost and abundant natural materials as alternative adsorbents for nitrophenols removal from aqueous phase.

Electrochemical processes offer useful possibilities for *in situ* and local treatment of industrial waste-waters at anodes for destructive removal or modification of noxious solutes (especially organics) [9]. For this latter type of impurity solutes, the overall process most desirable is that of so-called "mineralization", *i.e.*, complete anodic oxidation to CO₂ and H₂O, or additionally to N₂ or NO₃⁻⁷, or to SO₄²⁻ in the case of N or S containing organics, respectively. The use of electrode processes has the advantage that no added oxidizing or reducing agent needs to be provided since electrons (at cathodes) or their vacancy states (at anodes) are direct reagents, so that no other undesired exogenous products will arise. The oxidation of phenols on different types of modified oxides anodes, including the DSA commercial anodes has been studied, by cyclic voltammetry, polarization measurements, and electrochemical impedance spectroscopy and potentiostatic transients in different aqueous solutions [11-14]. The formation of a phenoxy radical in a diffusion-controlled irreversible process is the initial step. In the concentrated phenols solution, the polymerization of phenoxy radicals leads to the formation of porous polyoxyphenylene film, strongly adherent to the electrode surface; this film inhibits partially the further oxidation [3].

Continuing our previous research focus on the electrochemical investigation of the phenols and other organic pollutants oxidation [12, 13], this paper presents the results obtained by combining two procedures in order to eliminate the nitrophenols: adsorption of nitrophenols from the quasi concentrated solution (10⁻²M) and the electrochemical oxidation of the remaining nitrophenols.

Experimental

Nitrophenols are sparingly soluble in water and therefore, both the adsorption and electrochemical properties have been investigated from their hydro alcoholic solutions (1/1 water/n-propyle alcohol mixture). For the electrochemical measurements the 0.1 M Britton-Robinson buffer electrolytes was used. The voltamperommetric and spectrophotometric control of nitrophenols has been achieved using a common three electrodes cell (WE: glassy carbon and; RE: Ag/Ag⁺/AgCl; AE: Pt wire).

The reactor setup used for adsorption measurement contains magnetic stirred and thermostated saturation vessel equipped with a reflux condenser. In a typical adsorption experiments, 0.5 mg of zeolites were been added to a glass reactor followed ADSORPTION AND ELECTROCHEMICAL DATA FOR p-NITROPHENOL REMOVAL FROM SYNTHETIC ...

by the addition of known quantity of nitrophenols in 1/1 water/i-propanol mixture. Initial sample was taken before starting the experiment. Each experiment was carried out at the constant temperature. The experiment was continued until analysis of two successive samples was constant, which confirmed the adsorption equilibrium stage.

The concentration of the remaining nitrophenols, after their adsorption on zeolites has been reduced by electrochemical way (oxidation or/and reduction), using a modified PRIAM reactor [12]. The intermittent samples were taken out at certain time intervals during both adsorption and electrochemical experiments and analyzed by UV-VIS (Pye UNICAM Helios β) and cyclic voltammetry (ECOCHEMIE-BAS100W).

The yellow benthonite used for the investigations was taken from deposits in the southern part of Bulgaria. The zeolites were taken from deposits in different regions in Romania: zeolite 1- from the region Marsid, zeolite 2- from Macicas and zeolite 3- from Sacaramb.

Results and discussion

Adsorptive properties of the investigated sorbents

The chemical composition and physical properties of the zeolite 1, zeolite 2 and yellow benthonite are presented in Table 1. Specific surface area and pore volume of the natural materials were determined using Sorptomatic 1990, Fisons instruments.

The adsorption isotherms, Figure 1 (plots of q - the quantity of adsorbed solute per unit weight of adsorbent, versus C_e - equilibrium concentration):

$q = V (C_0 - C_e)/w$

suggested that the isotherms of p-nitrophenols on zeolites are similar to Langmuir isotherms.

Table 1

Content, %	Zeolite 1	Zeolite 2	Yellow bentonite
SiO ₂	65.43	63.905	59-75
TiO ₂	0.29	0.385	0.1-0.8
Al ₂ O ₃	13.94	14.455	12-16
Fe ₂ O ₃	1.31	1.74	1-6
CaO	3.98	5.335	1.8-5
MgO	0.41	0.30	0.9-3
Na ₂ O	0.28	1.02	0.5-1
K ₂ O	2.06	0.86	0.5-1
Loss at ignition	12.3	12.00	-
Specific surface area m ² g ⁻¹	44.56	27.33	57.99
Pore volume at p/p ⁰ 0.99, cm ³ g ⁻¹	0.133	0.132	0.169
Monolayer volume, cm ³ g ⁻¹	8.8704	8.5030	13.3204

Characteristics of zeolites

The Langmuir adsorption isotherm has been successfully applied to many adsorption processes. A basic assumption of the Langmuir theory is that adsorption takes place at specific homogenous sites within the absorbent. It is then assumed that once an organic molecule occupies a site, no further adsorption can taken place at that site. Theoretically, therefore, a situation value is reached beyond which

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no further sorption can take place. The values of the calculated Langmuir constants (K_L, a_L) and the correlation coefficient (R^2) are listed in Table 2.

The adsorption of p-nitrophenol on zeolites was compared with the data on activated carbon indicating that the adsorption is quite similar, in both cases could be attributed to the physisorption.

The adsorbed quantity of monolayer, q_m was found to vary with temperature. The adsorption equilibrium constant, K, decreased with increase in temperature, which was also consistent with the Langmuir theory.

Form the comparative analysis of the experimentally obtained adsorption capacity of the studied natural materials it was established that zeolite 2 and yellow bentonite exhibited the highest affinity to 4-NP. The values of the equilibrium and monolayer adsorption capacity of the sorbents to 4-NP follow the order:



Yellow bentonite = zeolite 2 > zeolite 1

Figure 1. The adsorption isotherms comparing with Langmuir model
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				-		
Sorbent		Langmuir			Freundlich	
Solbent	KL	a∟	R^2	K _F	n _f	R^2
Zeolite 1	12.5313	3.4097	0.9911	2.3911	0.1827	0.6694
Zeolite 2	3.4435	0.5557	0.7642	1.3270	0.7259	0.8076
Yellow bentonite	4.2212	0.6809	0.9875	2.7077	0.3385	0.8787

Value of the calculated constants in the Langmuir and Freundlich models.

Adsorption experiments of p-nitrophenol on Zeolite-2

During the adsorption of p-NP on Zeolite-2 the change of color has been observed, as can see in UV-spectrum, Figure 2 (λ =400nm);



Figure 2. Evidence for p-NF oxidation after the adsorption on Zeolite-2, after 20 min of adsorption (see Fig.3).



Figure 3. Effect of the initial concentration on adsorption of p-NP on benthonite. P=300 rpm; W=2 g

Table 2

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The surface of sorbent has been totally covered after 10-20 min, depending on the initial concentration.

This behavior (Figure 2) could be explained by the catalytic activity of the Zeolite-2 for p-NP oxidation. The most important results of p-NP adsorption are: decrease of p-NP concentration with (5-18) %, depending on the zeolite nature and conditions and partial oxidation of p-NP to the quinone structure, easier to oxidize by electrochemical way.

Decrease of nitrophenols concentration by electrochemical oxidation According to our previous results the decrease of p-NF concentration could be realized both by reduction and oxidation [12].

The oxidation potential of 4-nitro phenol slowly decrease at the same time with increasing of phenols concentration, due the increasing of the coverage degree, Fig. 3



Figure 3. Dependence of oxidation potential of p-NP on concentration on GC.

Because each type of electrode shows a different behavior depending on its superficial characteristics and properties, by electrochemical oxidation on (MOE) and GC (glassy carbon electrode) the total mineralization takes place with different yields.

To make the comparison between these types of electrodes (GC and MOE), the oxidation of aqueous synthetic phenolic water was chosen.

As can be seen in figure 4, on (MOE) [16] an almost complete mineralization of the p-NP achieves and the mineralization rate is higher than those obtained on GC. The electrochemical total oxidation of the remaining p-NP, on (MOE) from about 1 mM to $10-50 \mu$ M.

On the other hand the analysis of the answer after multiples scans shows that polymerization is an important secondary reaction pathway on GC, Figure 5.

Preliminary data by the GPC liquid chromatography have demonstrated that the polymeric material developed in the process was a mixture of the various polymers with low molecular weight (ranging from 200 to 500 mg/mmol). Conversely, the oxidation of the phenol waste using (MOE) electrode deals to the sequential formation of aromatic compounds, carboxylic acids and carbon dioxide, Table 3.









Figure 5. Polymers formation on GC.

Table 3

Preliminary data concerning the intermediates and final products of electro-oxidation (mg/l)

τ, ο, ,	
GC	MOE
1.6	1.1
2.8	2.5
38.5	0.0
30.0	90.3
26.6	6.1
	GC 1.6 2.8 38.5 30.0 26.6

Results obtained in this work could be explained in the terms of the oxidations of phenol to phenoxy radicals (first step of the process). The oxidation performed in (MOE) is stronger dealing whit the quick formation of benzoquinone and to the split of the aromatic ring. On the contrary the softer oxidation performed in GC electrode.

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Conclusion

The combined procedure proposed in this paper is the adsorption of p-NP on zeolite in order to reduce the pollutant concentration up to 1 mM, followed by the electrochemical oxidation of remaining p-NP, up to 10–50 μ M. The electrochemical mineralization of the remaining p-NP on (MOE) could be realized at accessible temperatures and pH~2, when the competitive oxygen formation diminishes. The preliminary data on the nature and distribution of the oxidation intermediates and products has to be confirmed by future experiments.

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Dedicated to professor Gh. Marcu at his 80th anniversary

ON THE DEPOLLUTION OF SOME RADIOACTIVE EFFLUENTS

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ABSTRACT. In this review, parts of our recent analyses in the depollution of the radioactive wastewaters are reported. The studies were focused in two main directions: (1) laboratory analyses on biodepollution of low radioactive waste waters, both in the absence and in the presence of the ionic competition and (2) chemical treatment of the radioactive waste waters using microporous titanosilicates and calix[4]arene/ p-(*tert*-butyl)calix[4]arene. In some cases, different aspects of the chemical reactivity, the kinetic of the reactions and the thermodynamic properties are presented.

1. Introduction

The problems of environmental radioactivity pollution are of crucial concern to state and local authorities responsible for environmental protection and control of nuclear wastes and weapons. In principle, there are two sources of environmental radioactivity, namely natural and manmade [1].

The pathways of radionuclides through the environmental waters are extremely complex. Transportation by water contaminates soils and water sources (rivers, lakes and the sea), many of which are situated far away from the release point. This is why in the Laboratory of Radiochemistry of "Al.I. Cuza" University of laşi a reinforced research in this matter started about ten years ago.

2. Recent analyses of bioaccumulation using plant species and microorganisms

Application of bacteria, fungi, algae or even plants to clean up surface and ground water contaminated by radionuclides has been increasingly applied. Hence a large number of studies on bioaccumulation or biosorption of ${}^{51}Cr^{3+}$, ${}^{60}Co^{2+}$, ${}^{90}Sr^{2+}$, ${}^{137}Cs^+$, ${}^{204}TI^+$, Th(IV), U(VI) from liquid wastes on different biocollectors has been reported.

2.1. Bioaccumulation of uranium using different strains of Saccharomyces cerevisiae

Under conditions of a natural aqueous systems, the insoluble uranium rapidly corrodes forming yellow uranyl compounds, where the linear $[O=U=O]^{2^+}$ entity forms the characteristic structural elements. Most notably, the solution chemistry of

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U(VI) is relatively complex, with numerous mono- and polynuclear uranyl-hydroxide and uranyl-carbonate complexes being formed.

The possibility of bioaccumulation of uranium species in beer yeast was investigated [2]. The behaviour of the *S. cerevisiae* – $UO_2^{2^+}$ system was studied versus contact time, pH and anion nature without ionic competition. Analysis of the obtained data revealed the following optimal working conditions: 1 h contact time, pH = 6.5 and 0.1 M UO₂(CH₃COO)₂ solution as uranyl source; as result, the maximum degree of bioaccumulation attends a value nearly 8.75 mmol $UO_2^{2^+}$ /g yeast. Both scanning electron microscopy and amino acid determinations lead to the conclusion that the uranyl nitrate solution may devastate the yeast cells provoking membrane damage and the release of the cell constituents (including the bioaccumulated uranium species).

However, in natural contaminated waters, the uranium series decay products and the non-radioactive cations influence uranium bioaccumulation process. Consequently, five different strains of *S. cerevisiae* were tested to analyze the bioaccumulation of U(VI) from waste water containing competitive ions [3]. Samples of water passing out from a previous uranium mill were used. The accumulation capacities of the tested strains were different. The kinetics of bioaccumulation, the leaching degree, the influence of cell density and their origin were analyzed. Under the applied working conditions, more then a half of the total activity could be accumulated after 1 h contact time of 1 ml *S. cerevisiae* suspension and 5 ml of water. The heavy metals effectively competed the uranium accumulation.

2.2. Decontamination of radioactive liquid wastes by hydrophytes vegetal organisms

The bioaccumulation of some radioactive ions from contaminated waste solutions, on hydrophytic vegetal organisms. In order to follow the distribution of radioactive ions ⁵¹Cr³⁺ [4], ⁶⁰Co²⁺, ¹³⁷Cs⁺ [4,5] and ⁶⁵Zn²⁺ [6] in various cell components extracted from *Spirulina platensis, Porphiridium cruentum, Scenedesmus quadricauda, Lemna minor, Elodea canadensis, Pistia stratiotes, Riccia fluitans* and *Azolla caroliniana* the plants were "cultivated" in radioactive solutions.

The formed complexes were extracted with acetone or acetic acid and were chromatographically separated. The results show an intense activity of the polysaccharide and lipoid fractions in the bioaccumulation process. The unusually high removal activity of lipids was probably due to the partial hydrolysis into the fatty acids and triglycerides, with a greater chelating action of the cations. An example of thin layer radiochromatography experiment is presented in Fig. 1.

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Fig. 1. The amount of ¹³⁷Cs⁺ and ⁶⁰Co²⁺ – radioactive ions localized in different biochemical components of the living fern Azolla caroliniana Willd.

3. Chemical treatment of the low radioactive wastewaters 3.1. Sorption of $^{60}Co^{2+}$, $^{115}Cd^{2+}$, $^{137}Cs^+$ and $^{204}Hg^{2+}$ on ETS-4 and ETS-10 microporous titanosilicates

ETS-4 (synthesized from gel with following molar composition: $2.0Na_2O$: $0.3TiO_2$: 0.6KF: 2.56HCl: $1.49SiO_2$: $39.5H_2O$) and ETS-10 (synthesized from gel with following molar composition: $1.0Na_2O$: $1.49SiO_2$: $0.2TiO_2$: 0.6KF: 1.28HCl: $39.5H_2O$) were subjected to sorption of radioactive cations ${}^{60}Co^{2+}$, ${}^{115}Cd^{2+}$, ${}^{137}Cs^+$ and ${}^{204}Hg^{2+}$ from aqueous solution, in the absence of ionic competition [7,8]. The uptake of these radiocations was compared by means of the distribution coefficient (K_d) versus contact time, determining their cationic exchange capacity at equilibrium.

Table 1.

ETG-4 and ETG-10 litanosilicates.				
Radioactive ion	Temperature, K	K _d , ml/g	R, meq/g	
	277	436	3.03	
⁶⁰ Co ²⁺	293	733	4.23	
	313	760	4.31	
	277	631	3.87	
115 Cd ²⁺	293	791	4.43	
	313	811	4.48	
	277	1258	3.83	
¹³⁷ Cs ⁺	293	1636	4.50	
	313	1670	4.55	
	277	341	2.54	
²⁰⁴ Hg ²⁺	293	627	3.85	
	313	808	4.45	

The distribution coefficients of ⁶⁰ Co ²⁺ , ^{115m} Cd ²⁺ and ²⁰³ Hg ²⁺ on			
ETS-4 and ETS-10 titanosilicates.			

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For all sorption systems and the considered radiocations, a rapid increase of K_d in the first minutes of contact can be observed. The higher the exchange temperature, the higher the value of K_d ; further, the equilibrium value is established more quickly. The maximum sorption capacity at equilibrium, stated in ml/g (K_d) and meq/g (exchange capacity – R), are summarised in Table 1. The values of ΔH^0 , ΔS^0 and ΔG^0 for the considered sorption systems were given in Table 2.

Table 2.

Thermodynamic parameters for the sorption of ${}^{*}M^{n^{+}}$	radiocations on
ETS-4 and ETS-10 titanosilicates.	

Cotion Titoposilioot		K _d , cm ³ /g			
Cation	manosilicate	277 K	293 K	313 <i>K</i>	333 K
Co ²⁺	ETS-4	536	593	625	649
0	ETS-10	436	733	760	_ ^b
Cd ²⁺	ETS-4	583	651	693	750
Ca	ETS-10	631	791	811	_ ^b
1.1m ²⁺	ETS-4	644	778	801	810
Hg⁻	ETS-10	341	627	808	_ ^b

3.2. Use of some calixarenes as cleaning agents for low radioactive waste waters Calixarenes are products of the condensation of phenol *p*-substituited with formaldehyde in an alkaline medium. One of the most important practical applications of this class of compounds is the environmental decontamination of waters containing ions of the heavy metals like Cs⁺, Ag⁺, Au⁺, Hg²⁺, Pb²⁺, Cd²⁺ etc.

In order to clean some low radioactive contaminated waters containing $\beta+\gamma-active$ cations (⁵⁵⁺⁵⁹Fe³⁺, ⁶⁰Co²⁺, ⁶⁵Zn²⁺ and ¹³⁷Cs⁺), calix[4]arene and *p*–(*tert*–butyl)calix[4]arene has been used [9]. Experiments were performed in the absence of ionic competition, at 277, 293 and 313 K. Whatever the temperature and the sorbent used, the capacity of retaining the $\beta+\gamma-active$ cations varies as follows: ⁵⁵⁺⁵⁹Fe³⁺>⁶⁰Co²⁺>⁶⁵Zn²⁺>¹³⁷Cs⁺ (the capacity of sorption decreases with the increasing ionic radius and decreasing cationic valence).

In the uranium case, the interaction is a chemical one, a new 1:1 $UO_2^{2^+}$: *p–tert–*butylcalix[4]arene complex being synthesised [10] in acetone. The combination metal: ligand ratio was determined by mass spectrometry and by the Job method. The chemical binding of uranium was proved by FT–IR spectroscopy and by a leaching study. The mass spectrum of the $UO_2^{2^+}$:*p–tert–*butylcalix[4]arene complex (Fig. 2) indicates that the molecular peak appears at m/z = 908 a.m.u. which correspond to a combination ratio M:L = 1:1 (650÷700 K). The most intense peaks appears at m/z = 56 a.m.u. (solvent) and m/z = 648 a.m.u. (*p–tert–butylcalix*[4]arene ligand). Although other significant peaks do not appear in the 648÷908 u.a.m. m/z range, we cannot conclude that the prepared compound is pure, because a part of ligand may be still unreacted.

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Fig. 2. Mass spectrum of the 1:1 UO₂²⁺:p-tert-butylcalix[4]arene complex.

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INHIBITORY EFFECT OF METOPROLOL UPON CATALASE-H₂O₂ DECOMPOSITION, USED AS A POTENTIAL KINETIC METHOD TO DETERMINE THE DRUG CONCENTRATION

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ABSTRACT. The enzyme catalyzed process of decomposition of hydrogen peroxide has been investigated by means of a Clark oxygen sensor, in the presence and absence of various concentrations of Metoprolol – a β -blocker drug - having an inhibiting role. The Michaelis – Menten kinetic parameters of H₂O₂-catalase reaction have been determined from Lineweaver-Burk plots. The inhibition pattern we have deduced, suggested by the Lineweaver - Burk plots, corresponds to a fully mixed inhibition mechanism. Inhibition constants K₁ and K₁ were determined. The inhibitory effect can be used to determine metoprolol in low concentrations by a kinetic method.

Keywords: Metoprolol, Catalase, Hydrogen peroxide decomposition, Inhibition mechanism, Kinetic methods of analysis.

INTRODUCTION

Inhibitors of enzymatic reactions have acquired large applications in medical and pharmaceutical research [1]. The catalase mediated decomposition of H_2O_2 :

$$2H_2O_2 \xrightarrow{catalase} 2H_2O + O_2 \tag{1}$$

has been the subject of extensive investigations both with regard to the kinetics and to the mechanism of the reactions.[2, 3, 4, 5]

Similar reactions were also observed for other organic peroxides [6, 7] Well-known inhibitors of these reactions are ions like azide, cyanide and fluoride [8, 9, 10], or some organic compounds like 3-amino-1, 2, 4-triazole [11] atenolol [12] and metoprolol [13].

Catalase is widely distributed in nature. It is found in all aerobic microorganisms, in plant and animal cells [14]. The catalase activity of mammalian tissues varies greatly: it is highest in liver and kidney and low in connective tissues. The enzyme, when located in organelles, acts as a regulator of the H_2O_2 level, while in erythrocytes, catalase provides protection for hemoglobin against the oxidizing agents like H_2O_2 together with gluthathione peroxidase [16].

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Inhibitors are compounds that slow down the activity of the enzyme by preventing either the formation of substrate-enzyme or breaking down enzyme-product complexes.

Metoprolol ($C_{34}H_{56}N_2O_{12}$) is a cardioselective β -adrenoceptor blocking agent (β -blocker). Its formula is (±)-1-isopropyl-amino-3-p-(2-methoxyethyl)- phenoxypropan-2-ol(2R,3R)-tartrate with number 56392-17-7, in accordance to British Pharmacopeia[17],



This agent is used in clinical medicine for the treatment of various diseases including hypertension, pectoral angina, cardiac aritmia, and especially ventricular tahicardic and hart attract, [18, 19, 20]

A number of analytical methods have been developed for its determination including spectrophotometrical [21], potentiometrical and amperometric methods [22, 23, 24]. The purpose of this work is to establish the type of inhibition for this reversible inhibitor by it effect on the rate of hydrogen peroxide decompositions catalyzed by catalase.

EXPERIMENTAL

Equipment. The measurements were undertaken with a Clark oxygen sensor, attached to a Multiline P4 multimeter with automatic data acquisition on a PC. The sensors cover a measuring range from 0 up to 19.99 mg/L for the dissolved oxygen, with a resolution of 0.01 mg/L and an accuracy of $\pm 0.5\%$ from 5.30 °C. All experiments were performed in a vessel provided with a water jacket. To maintain a constant temperature value (20 \pm 0.1 °C) it was connected to a Falc 90 recirculation water bath. The reaction mixture was stirred during the run with a magnetic stirrer, always with the same frequency.

Reagents and solutions. We used bacterial catalase from *Micrococcus Lysodeikticus* 176340 U/ml (where the enzyme unit 1U is the amount of enzyme needed to transform 1 µmol of the substrate within 1 min, under standard conditions), from FLUKA with a purity index of 0.85, which came from the ratio of absorbance at 405 nm and 280 nm, A_{405}/A_{280} . The molar concentration of catalase was determined spectrophotometrically at 407 nm, where the molar absorptions coefficient is known to be $4x10^5 \text{ mol}^{-1} \text{ dm}^3 \text{ cm}^{-1}$ [21].

All the other reagents were of analytical reagent grade. The solutions were prepared with de-ionized, four-times distilled water in order to avoid the interference of heavy metals. Stock solutions of catalase (9x10⁻¹⁰ mol/L), H₂O₂ (4.9 x 10⁻² mol/L) and metoprolol (10⁻⁵ mol/L) in phosphate buffer of pH=7.0 were freshly prepared before

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each set of runs. Hydrogen peroxide has been standardized against potassium permanganate in acidic media.

Procedure. The reaction mixture of 10 mL volume was prepared directly in the reaction vessel connected to a thermostat. All the solutions of the reagents were kept in the water bath. Measured volumes of buffer and metoprolol stock solution were placed in the vessel, and the change in the concentration of oxygen was monitored while the mixture was continuously stirred. The oxygen content of the solutions became constant after about 50 s, when a known volume of H_2O_2 stock solution was added. After about 50 s the concentration of oxygen reached a constant value. Now, the catalyzed reaction was started by a quick adding of 2 mL of the enzyme stock solutions by means of syringe. A typical oxygen concentration versus time curve obtained by monitoring the reaction with a Clark sensor is presented in Fig. 1. The rate has been measured after the introduction of the enzyme solution into the mixture as the slope of the oxygen concentration increase with time. Only the starting period, within 25 - 30 % of reaction, with a linear dependence has been considered. The degree of H_2O_2 transformation was calculated from the experimental measured oxygen concentration as

$$X = \frac{2[O_2]_t - [O_2]_0}{[H_2O_2]_0}$$
(2)

where $[O_2]_t$ is the actual concentration of O_2 , and $[O_2]_0$ is the constant value of $[O_2]$ obtained after the addition and consumption of the whole amount of H_2O_2 .



Fig. 1. A typical curve for O_2 evolution, recorded with the Clark sensor in the reaction mixture pH=7 at $20^{\circ}C$

RESULTS AND DISCUSSION

A large amount of information is available in the literature concerning the kinetics and the mechanism [4, 26, 27, 28] of the reaction in the absence of inhibitor. Although the catalase- H_2O_2 complex formed in the first step of the reaction interacts with another hydrogen peroxide molecule [29] and some exchange of the valence state

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of iron ion takes place, we obtained a Michaelis behavior of the over-all reaction (1) with the initial rates.

The initial reaction rates were determined from the slopes of the early part of the O_2 evolution curves, after catalase addition. According to the well-known Michaelis-Menten equation [30]

$$\mathbf{r}_{0} = \frac{\mathbf{k} \begin{bmatrix} \mathbf{E}_{0} \end{bmatrix} \begin{bmatrix} \mathbf{S} \end{bmatrix}}{\mathbf{K}_{M} + \begin{bmatrix} \mathbf{S} \end{bmatrix}} = \frac{\mathbf{r}_{\max} \begin{bmatrix} \mathbf{S} \end{bmatrix}}{\mathbf{K}_{M} + \begin{bmatrix} \mathbf{S} \end{bmatrix}}$$
(3)

where $[E]_0$ and [S] stand for the initial concentration of the enzyme and for the substrate concentration respectively, K_M and r_{max} are the Michaelis-Menten parameters, and k is the rate constant of breakdown of the enzyme substrate complex to the product. This equation can be brought into the double-reciprocal form:

$$\frac{1}{r_0} = \frac{1}{r_{max}} + \frac{K_M}{r_{max}} \cdot \frac{1}{[S]}$$
(4)

and used to obtain the Lineweaver-Burk plots [31]. The value of the Michaelis-Menten constant $K_{\rm M}$ = 4.55x10⁻⁴ mol/L and the maximum velocity $r_{\rm max}$ =5.11x10⁻² mol L⁻¹ s⁻¹, obtained by us, are comparable to those mentioned in literature[32]. And dose to those determined in our previous study. [12]

The influence of metoprolol on the enzyme-catalyzed decomposition of hydrogen peroxide .Several kinetic runs performed in the presence of different concentrations of metoprolol proved the inhibitory effect of this compound. When the concentration of H_2O_2 was varied at several chosen concentrations of metoprolol, Lineweaver-Burk plots for the inhibited reaction

$$\frac{1}{r_{o(I)}} = \frac{1}{r_{max(I)}} + \frac{K_{M(I)}}{r_{max(I)}} \cdot \frac{1}{[S]}$$
(5)

Table 1

resulted in a family of straight lines (Fig. 2), which have a common intersection point on the left side of the ordinate and below the abscissa.

•		
[Metoprolol] ₀ x10 ⁷ (mol/L)	Intercept (L.s/mol)	Slope (s)
0	19.538	0.0089
1	23.147	0.0095
2	27.150	0.0099
3	37.120	0.012

Equations of Lineweaver-Burk plots for different concentration of metoprolol

As the slopes of Lineweaver- Burk plots depend on the inhibitor concentration, a systematic study could lead to a kinetic method of determination of this inhibiting agent concentration. The Michaelis-Menten parameters in the presence of the inhibitor were determined from the parameters of the lines in figure 2. Their values are given by the equations (6):

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$$\mathbf{r}_{\max(\mathbf{I})} = \mathbf{r}_{\max} \quad \frac{1 + \beta \frac{[\mathbf{I}]}{\mathbf{K}'_{\mathbf{I}}}}{1 + \frac{[\mathbf{I}]}{\mathbf{K}'_{\mathbf{I}}}} \quad \text{and} \quad \mathbf{K}_{\mathbf{M}(\mathbf{I})} = \mathbf{K}_{\mathbf{M}} \left(\frac{1 + \frac{[\mathbf{I}]}{\mathbf{K}_{\mathbf{I}}}}{1 + \frac{[\mathbf{I}]}{\mathbf{K}'_{\mathbf{I}}}} \right)$$
(6)

where: K'₁ and K₁ are the dissociation constants of the enzyme – substrate - inhibitor complex, ESI, and the enzyme. inhibitor complex, EI. The constant β is equal to zero for full inhibition, while for partial inhibition 0< β ≤1. Lineweaver- Burk plots of Fig. 2 correspond to the inhibition pattern of either full or partial mixed inhibitors [33].



Fig.2. Lineweaver Burk plots obtained for different concentration of metoprolol

In order to distinguish between full and partial inhibition, the slopes and intercepts from the primary Lineweaver- Burk plots were re-plotted against the corresponding inhibitor concentration. They are presented in figure 3. Both plots gave straight lines. This behavior is considered as typical for full inhibition [24]. Considering the possibility of a partial inhibition, a non-linear fit of the intercept 1/ $r_{max(I)}$ =f([I] and the slope K_{M(I)}/ $r_{max(I)}$ =f [I] gave negative values for β . Therefore it is very likely that metoprolol acts as a reversible full mixed inhibitor according to the reaction scheme 1

The plot (Fig.3) of slopes and intercepts respectively, against the inhibitor concentration were used to determine the dissociation constants mentioned in the scheme above, $K_I=8.86 \times 10^{-7}$ M for EI complex and $K'_I=3.04 \times 10^{-7}$ M for ESI complex . The constant K_I was obtained from the slope of the $K_{M(I)}/r_{max(I)}$ versus [Metoprolo]] graph:

$$K_{I} = \frac{K_{M}}{r_{\text{max \cdot slope}}}$$
(7)

while the K_1 constant was determined from the slope of the $1/r_{max(1)}$ versus [metoprolol] graph:

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Scheme 1. Mixed inhibition mechanism for metoprolol, where E stands for catalase, S for hydrogen peroxide and I for metoprolol.





$$K'_{I} = \frac{1}{r_{\text{max} \cdot \text{slope}}}$$
(8)

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Table 2

Mean values of $r_{max(I)}$ and Michaelis constant in the presence of the inhibitor $K_{M(I)}$

- ()			
[Metoprolol] ₀ x 10	′(mol/l) r	max(I) x 10 ² (mol/Ls)	K _{M(I)} x 10 ⁴ (mol/L)
0.00		5.11	4.55
1.00		4.32	4.10
2.00		3.68	3.64
3.00		2.69	3.23

Kinetic method for the determination of metoprolol

We tried to exploit the inhibitory effect of metoprolol, upon the catalytic reaction of catalase, for the determination of this compound by means a kinetic method. The method consist in the monitoring of the oxygen evolution by means of Clark sensor The results were employed to obtain a calibration graph of initial rate against metoprolol concentration at fixed concentration of H_2O_2 and catalase. The graph exhibit a non linear aspect as described by the equation (9) and presented in figure 4.



Fig 4. Calibration graph for metoprolol

$$\mathbf{r}_{(\mathbf{I})} = \frac{\mathbf{r}_{\max} \cdot [\mathbf{S}]}{\frac{\mathbf{K}_{\mathbf{M}}}{\mathbf{K}_{\mathbf{I}}} \cdot [(\mathbf{K}_{\mathbf{I}} + [\mathbf{I}]) + [\mathbf{S} \cdot (\mathbf{K'}_{\mathbf{I}}) + [\mathbf{I}]]]}$$
(9)

Its linear form is presented in figure 5.

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Fig 5. Linear calibration for metoprolol

It is obvious that heavy metals ions (Cu ²⁺, Fe ³⁺, Zn ²⁺, Ni ²⁺) will influence the reaction rate, because they have a catalytic effect on the decompositions of H₂O₂, but according to the literature there are many organic compounds (especially aromatic compounds) that may interfere too, because they act as inhibitors of catalase, including: naphthalene, 2-naphthol, 2- naphthalene-sulfonic acid, benzene, phenol, benzenesulfonic acid, menthol, inositol, biotin, procaine, sulfanilamide, pyridoxine, folic acid, aminopterin, and riboflavine [34]

Further sturdy will elucidate the potential interference on this kinetic method.

CONCLUSIONS

The catalase –catalyzed decomposition of hydrogen peroxide in phosphate buffer was studied in the presence of metoprolol. The metoprolol acts like a fully mixed inhibitor, as observed from Lineweaver-Burk plots.

Michaelis Menten parameters were determined for the decomposition of H_2O_2 in the present of catalase yielding the values of K_M =4.55x10⁻⁴ mol/L and

r_{max}=5.11x10⁻²mol/L⁻¹s⁻¹

Linear dependence of the slope and respectively intercept of Lineweaver-Burk linearisations from metoprolol concentration indicates a fully mixed inhibition mechanism.

A kinetic method was suggested for the determination of metoprolol according to its inhibitory effect.

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STUDIES ON THE INFLUENCE OF FLUX NATURE ON THE PROPERTIES OF NIOBIUM ACTIVATED YTTRIUM TANTALATE PHOSPHOR

AMALIA MESAROS^{1,2}, ELISABETH - JEANNE POPOVICI¹, LAURA MURESAN¹, MARIA ŞTEFAN^{1,2}, RODICA GRECU¹ AND MARILENA VASILESCU³

ABSTRACT. Niobium activated yttrium tantalate (YTaO₄:Nb) presents good X-ray absorption and emits in the blue region of the spectrum. The goal of the paper is to study the influence of flux nature on the crystalline structure, morphology and luminescent characteristics of YTaO₄:Nb powders. Phosphors samples were prepared by solid state reaction route and theirs properties were investigated by X-ray diffraction (XRD), Fourier transform infrared spectroscopy (FTIR), photoluminescence (PL) measurements and scanning electron microscopy (SEM).

Keywords: yttrium tantalate, phosphors, luminescence, X-ray imaging

INTRODUCTION

Niobium activated yttrium tantalate, YTaO₄:Nb is an efficient luminescent material used in medical X-ray imaging applications[1,2]. The characteristic emission spectra consist in a broad band situated in the UV-blue domain of the electromagnetic spectrum with a maximum at 390-410 nm. Performances of YTaO₄:Nb phosphor powder are correlated with the crystalline structure, particle size, morphology and luminescence properties.

It is well known that the emission intensity and colour purity of $YTaO_4$ -based phosphor are extremely sensitive to crystalline phase composition of the materials [3-5]. Depending on the synthesis conditions yttrium tantalate present different crystalline structures. There are two polymorphs, i.e. high temperature tetragonal (T-YTaO₄ phase, scheelite structure) and low-temperature monoclinic (M-YTaO₄ phase, fergusonite structure) forms. There is an additional monoclinic phase, designed as M-prime form (M'-YTaO₄) that can be obtained in appropriate conditions, namely below 1450°C. The high luminescence performances of niobium activated

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yttrium tantalate phosphor are associated with the monoclinic crystalline structure, where M'-YTaO₄ represents the equilibrium phase at room temperature.

The preparation of $YTaO_4$: Nb phosphor is usually achieved by solid state reaction route from synthesis mixtures containing different metallic oxide sources. The monoclinic M' crystalline structure of the $YTaO_4$ host lattice, as well as the emission centre formation is substantially improved when the thermal synthesis is flux-assisted by some inorganic salts.

The paper presents several aspects referring to the synthesis of niobium activated yttrium tantalate phosphor (YTaO₄: Nb). The influence of the flux nature on the crystalline structure, particle morphology and luminescent characteristics of $YTaO_4$: Nb phosphor is investigated in order to identify an optimal flux reagent that could generate a high performing material.

EXPERIMENTAL PART

Niobium activated yttrium tantalate samples were prepared by solid state reaction route from homogeneous mixtures consisting of raw oxide precursors Y_2O_3 (99.9%), Ta_2O_5 (Optipur), Nb_2O_5 (99%) and Li_2SO_4 (99%) and/or Na_2SO_4 (99%) as flux. The stoichiometric amounts of Y_2O_3 , Ta_2O_5 , Nb_2O_5 and 30 wt % alkaline sulphates were ball-milled with acetone and dried at 70°C. The powders mixture was calcined in air at 1200°C, for 4 h and slowly cooled to the room temperature. Finally, phosphors samples were water washed, dried and sieved.

The as-prepared phosphors were characterized by fluorescence and FTIR spectroscopy, X-ray diffraction and scanning electronic microscopy.

Photoluminescence (PL) measurements were performed at room temperature, with a Perkin-Elmer 204 Fluorescence Spectrophotometer. The emission spectra were registered under 254 nm excitation and were normalised in comparison with an internal standard. X-ray Diffraction (XRD) analysis was performed on SIEMENS D5000 diffractometer (CuK α radiation). IR absorption spectra were registered on JASCO 610 FTIR Spectrometer (KBr pellets technique). Scanning Electron Microscopy (SEM) analysis was performed using a LEO 1550 microscope.

The interpretation of XRD data was achieved on the basis of powder diffraction files namely, PDF 00-024-1425 for M'-YTaO₄, PDF 00-048-0265 for orthorombic- Y_3TaO_7 .

RESULTS AND DISCUSSION

Niobium activated yttrium tantalate phosphors (YTaO₄:Nb) were obtained by the classic solid state reaction route, from synthesis mixtures containing yttrium oxide and tantalum oxide as generators of the host matrix, niobium oxide as generator of activator ions, and lithium sulphate and/or natrium sulphate as flux. The formation of yttrium tantalate activated with 15 mole % niobium phosphors could be described by the equation (1):

$$Y_2O_3 + (1-x)Ta_2O_5 + xNb_2O_5 \xrightarrow{M_2SO_4,T} 2YTa_{1-x}Nb_xO_4 (1)$$

where: x=0.15

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During the thermal synthesis stage, the flux reacts with the mixture of oxides to produce intermediate compounds that are more reactive than the starting oxides. According to the literature, a high calcination temperature facilitates their mutual interaction to give the final product and to regenerate the flux [1,2]. For this reason, the flux compound is considered as catalyst or reactive flux [6].

Photoluminescence (PL) characteristics, crystalline structure, particle morphology and sizes of $YTa_{0.85}Nb_{0.15}O_4$ samples were determined in order to establish the correlation between the phosphors properties and their synthesis conditions.

Luminescence properties

All powder samples are white coloured and exhibit blue luminescence during a 254 nm excitation.

The emission spectra show broad bands with a maximum at around 390 nm (**Fig. 1**). The sample, which was prepared at 12000C with Li2SO4 as flux, presents the highest luminescent intensity. Using a Li_2SO_4 - Na_2SO_4 mixture or Na_2SO_4 as flux, an obvious decrease of the YTaO₄: Nb phosphor emission intensity can be observed.



Fig. 1. Emission spectra of YTa_{0.85}Nb_{0.15}O₄ samples (λ_{exc}= 254 nm)

The flux-dependence of the luminescence intensity can be partially explained by the crystalline structure or particle morphology and the size of $YTaO_4$ -based phosphors obtained in different conditions.

Crystalline structure

The crystalline structure and order degree of phosphors were evaluated on the basis of X-ray diffraction patterns (XRD) and FTIR spectra (**Fig. 2**).

The most homogeneous crystalline phosphor powder was obtained at 1200^oC, by using Li₂SO₄ as flux. Sample T7, YTaO₄:Nb [Li₂SO₄; 1200^oC] is a single

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phase material of monoclinic M' polymorph form. Replacing Li_2SO_4 with Na_2SO_4 (sample T9), additional reflections are observed that can be ascribed to some intermediate compounds, such as the orthorhombic Y_3TaO_7 . This shows that at 1200°C, sodium sulphate does not assure the complete conversion of oxides into yttrium tantalate phase.

Mention has to be made that our experimental data for the compounds obtained in the Y_2O_3 -Ta₂O₅-Nb₂O₅ system are in good agreement with the literature XRD data for the Y_2O_3 -Ta₂O₅ system. This fact evidences that niobium oxide is well dissolved into the YTaO₄ crystalline lattice to form the niobium activated yttrium tantalate phosphor, as suggested by the PL spectra.



Fig. 2. XRD patterns (left) and FT-IR (right) spectra of YTa_{0.85}Nb_{0.15}O₄ samples prepared with different fluxes,T7= Li₂SO₄]; T8= Li₂SO₄+Na₂SO₄; T9= Na₂SO₄ (M'=M'-YTaO₄ and O= orthorhombic Y₃TaO₇ structures)

The infrared spectra confirm that flux nature influences the crystalline order degree of phosphors, as it was already illustrated by the XRD patterns. Sample T7 prepared with Li_2SO_4 has a single phase M'- structure, as shown by the well formed 439 and 808 cm⁻¹ bands. In sample T9, obtained with Na₂SO₄ as a flux, the M' phase still exists, but the crystalline structure is less organised, as suggested by the FTIR spectra.

For the Y_2O_3 -Ta₂O₅-Nb₂O₅ system, Na₂SO₄ shows reduced flux reactivity as compared to Li₂SO₄. This behaviour is responsible for the weakest luminescence emission observed for T9 samples.

Particle morphology and sizes

SEM investigations were performed with the aim to characterize and compare the particle morphology and sizes for $YTaO_4$:Nb powders obtained in different synthesis conditions (**Fig. 3**)

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Fig. 3. SEM images of $YTa_{0,85}Nb_{0.15}O_4$ samples prepared with different fluxes: T7 - Li_2SO_4 (left) and T9 - Na_2SO_4 (right)

Using Li₂SO₄ as flux (sample T7), non-agglomerated particles with regular shape are obtained. Most of the polyhedral, elongated crystals are of $2 \div 5 \mu m$ in length and about $1\mu m$ in thickness. The partial or the total replacement of Li₂SO₄ by Na₂SO₄ strongly decreases the particle dimensions. In these conditions, particles with less regulated shape and with aggregation tendency are formed.

The most heterogeneous crystalline powder was prepared with Na_2SO_4 , while the most homogeneous powder was obtained with Li_2SO_4 as flux.

Even if Li₂SO₄ (m.p.=884°C) and Na₂SO₄ (m.p.= 860°C) show close melting point values, their behaviour as flux is very different due to the large difference between the two cation sizes. The fact that Li⁺ (0.073 nm) is much smaller than Na⁺ (0.113 nm) is in the favour of the total conversion of oxides into the M'-YTaO₄ crystalline phase.

The general characteristics of $YTa_{0.85}Nb_{0.15}O_4$ phosphors prepared with different fluxes, i.e. the photoluminescence (PL) intensity at peak position, the main crystalline phase and the powder particle dimensions are summarized in Table 1.

Table 1

Phosphor Elux pature		General properties		
sample	(w/w)	PL intensity I ₃₉₀ (%)	Crystalline phases	Particle size (µm)
T7	30% Li ₂ SO ₄	125	M'	2.0 ÷ 5.0
Т8	15% Li ₂ SO ₄ -15% Na ₂ SO ₄	83	M'	0.5÷3.0
Т9	30% Na ₂ SO ₄	61	M' + O	0.2÷1.0

General characteristics of YTaO₄: Nb samples

where: $M' = M'-YTaO_4$; $O = orthorombic-Y_3TaO_7$

One can conclude that the highest luminescence intensity of $YTa_{0.85}Nb_{0.15}O_4$ [1200°C;Li₂SO₄] is associated with the high order degree of the crystalline lattice with M'-YTaO₄ structure phase. At 1200°C, Li₂SO₄ shows the strongest flux reactivity for the Y₂O₃-Ta₂O₅-Nb₂O₅ system. AMALIA MESAROŞ, ELISABETH - JEANNE POPOVICI, LAURA MURESAN, MARIA STEFAN, ET AL.

CONCLUSIONS

The flux nature is an important factor that determines luminescence properties, structural and morphological characteristics of niobium activated yttrium tantalate phosphor. Morphological and structural investigations put in evidence the high crystalline order degree of $YTaO_4$:Nb powders prepared with lithium sulphate as flux. The photoluminescence intensity is strongly affected by the crystalline homogeneity and particle dimensions of the phosphor powders.

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Dedicated to professor Gh. Marcu at his 80th anniversary

SPECTRAL INVESTIGATIONS OF EUROPIUM ACTIVATED YTTRIUM OXIDE PHOSPHOR PREPARED BY COPRECIPITATION METHOD WITH OXALIC ACID

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ABSTRACT. Europium activated yttrium oxide phosphors were prepared by reagent simultaneous addition technique, using oxalic acid as precipitating reagent. The aim of the paper was to study the influence of the firing conditions on luminescent and morpho-structural properties of Y_2O_3 :Eu phosphors. Thermal analysis, X-ray diffraction, SEM, FTIR and fluorescence spectroscopy were used to investigate precursor and phosphor powders. The correlation between the phosphor properties and precursor quality enabled us to select the optimal synthesis conditions.

Keywords: yttrium oxide, phosphors, luminescence; wet chemical method

INTRODUCTION

Europium activated yttrium oxide (Y_2O_3 : Eu) is a well-known red emitting phosphor used for applications in displays, optoelectronic devices and fluorescent lamps [1-3]. Phosphor utilisations depend on luminescence performances and powder characteristics that are defined during the synthesis stages. Luminescence characteristics, particle morphology and size or crystalline order degree are factors that determine the use of phosphor powders in optoelectronic devices.

Conventionally, Y_2O_3 : Eu powders are prepared from yttrium/europium oxide mixture, by the ceramic method that implies high firing temperature and long thermal treatment periods [4]. Mild thermal synthesis conditions could be used when oxides species are generated from precursors obtained through the wetchemical preparation route.

The paper presents several results referring to the synthesis of fine powders of Y_2O_3 : Eu phosphor from yttrium/europium oxalate precursor obtained by the wet chemical route. The co-precipitated precursor was prepared by using the reagent simultaneously addition technique –*SimAdd* developed in our previous works [5-7].

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In this purpose, europium containing yttrium oxalate based precursor was obtained under well controlled precipitation conditions from yttrium-europium nitrate and oxalic acid solutions and was converted into phosphors in different thermal treatment conditions. Spectral investigations as well as scanning electron microscopy (SEM) and thermal analysis were used for precursor and phosphors characterisation. A correlation between the preparation conditions and precursor and phosphors characteristics was established in order to improve the *SimAdd* technique for the obtaining of fine powders of europium activated yttrium oxide phosphors.

EXPERIMENTAL PART

Europium activated yttrium oxide phosphors (Y_2O_3 : Eu (3 mol%)) were prepared from yttrium-europium oxalate precursor.

Yttrium-europium precursor (sample code PG26) was prepared by the wet chemical method, using the simultaneous addition (*SimAdd*) technique of reagents into a bottom solution. Equal volumes of oxalic acid (0.45M) as precipitating reagent and yttrium-europium nitrate solution (0.30M) were simultaneously added, with equal constant flow (~5.7 mL min⁻¹), into a diluted solution of oxalic acid. Precipitation was carried out at 80^oC, under continuous stirring and pH monitoring. During the precipitation stage, the pH of the bottom solution was adjusted with ammonia to 2. The precursor post-precipitation treatment consisted of 24 h aging (maturation stage), water wash and drying. Yttrium-europium oxalate precursor was fired for 2 hrs in air, at 1100°C, 1200°C and 1300°C to provide the corresponding phosphor powder, P19, P27 and P28, respectively. No flux was used for the phosphor thermal synthesis stage. The as obtained materials were water washed, dried and sieved.

Precursors and phosphors were characterised by thermal analysis-TG-DTG-DTA curves (Paulik –Erdely Derivatograf OD-102; heating rate 6° C/min), infrared absorption spectroscopy -FTIR spectra (JASKO 610 FTIR Spectrometer; KBr pellets technique), photoluminescence measurements –PL spectra (JASKO FP-6500 Spectrofluorimeter Wavel; λ_{exc} =254nm), X-ray diffraction–XRD patterns (D8Advanced Brucker Diffractometer, CuKα radiation) and Scanning Electronic Microscopy-SEM images (JEOL –JSM 5510LV Microscope; Au-coated powders).

RESULTS AND DISCUSSION

This main aim of the study was to clarify some aspects concerning the synthesis of very fine particles of Y_2O_3 : Eu phosphors by wet chemical method, using the reagent simultaneous addition technique -*SimAdd*. The precursor was prepared by co-precipitation, from yttrium-europium nitrate mixture and oxalic acid.

Mention has to be made that in precursor and phosphor samples, about 3 mole% of yttrium was replaced by europium. It is supposed that, excepting the PL properties, the isovalent substitution of Y^{3+} with small amounts of Eu³⁺ does not influence the general characteristics of precursor and phosphors so that its presence is neglected during the discussion.

For the beginning, the precursor composition was estimated on the basis of the thermal analysis. TG-DTG-DTA curves of the PG26 precursor show that the thermal decomposition proceeds in three major steps. These steps can be associated SPECTRAL INVESTIGATIONS OF EUROPIUM ACTIVATED YTTRIUM OXIDE PHOSPHOR

with water loss (~ 191° C), decomposition of the metallic oxalate (~ 408° C) and probable, decomposition of the metallic oxy-carbonate intermediate (~ 530° C).

Assuming that the precursor is a metallic basic oxalate and taking into consideration the literature data [8], the yttrium oxide phosphor formation could be described by the following conversion path:

 $Y_2(C_2O_4)_3 \cdot xY(OH)_3 \cdot yH_2O \rightarrow Y_2(C_2O_4)_3 \cdot xY(OH)_3 \rightarrow Y_2O_2CO_3 \rightarrow Y_2O_3$

It is well known that the thermal synthesis conditions determine the phosphor characteristics [9]. In the present study, three phosphor samples were prepared in different thermal regime, namely P19 (1100° C), P27 (1200° C) and P28 (1300° C).

In order to correlate the precursor characteristics with the properties of the corresponding Y_2O_3 : Eu phosphors, FTIR spectra and XRD spectra were registered and comparatively analysed. **Fig. 1** and **2** presents FTIR and XRD spectra for PG26 and P27 samples.

The precursor vibrational spectrum consists of some characteristic bands i.e. v (H-O) ~3470, v (C=O) ~1634, v (C-O) ~1324, δ (O-C=O)+(Y-OH)~814-790. After the thermal treatment, the precursor specific FTIR bands disappear and the characteristic vibration of Y-O bond, at ~560cm⁻¹ can be observed. FTIR spectra suggest the complete conversion of yttrium oxalate-based precursor into yttrium oxide phosphors.

XRD spectra illustrate that all Y₂O₃: Eu phosphor samples as well as the parent precursor consist of well-formed crystalline phases. According to the literature, the yttrium oxalate-based precursor contains mostly the crystalline phase of yttrium ammonium oxalate (PDF 221047). All Y₂O₃: Eu powders are homogeneous and well crystallized, which demonstrate the formation of Y₂O₃-Eu₂O₃ solid solutions with cubic structure (PDF411105).

The XRD measurements suggest that yttrium-based precursor contains NH_4^+ ions. One can suppose that the specific ammonium infrared vibrations, i.e. v (N-H) and δ (N-H), are covered by those of the basic yttrium oxalate.

According to the XRD and FTIR spectra, the precursor can be formulated as: $[Y_{2-2x}(NH_4)_{2x}][(C_2O_4)_{3-3y}(OH)_{3y}]\cdot zH_2O$ where 0<x<1; 0<y<1.

The thermal treatment of yttrium-oxalate based precursors at $1100 \div 1300^{\circ}$ C ensures the incorporation of Eu³⁺ ions into the Y₂O₃ cubic crystalline lattice and the formation of luminescent centres.



Wavenumber (cm⁻¹)

FIG. 1. FT-IR SPECTRA FOR PRECURSOR (PG26) and phosphor (P27) prepared at 1200 °C



Fig. 2. XRD patterns for precursor (PG26) and phosphor (P27) prepared at 1200 °C

Fig. 3 presents the emission spectra of yttrium oxide phosphor (P19) registered for different excitation radiations.

On the basis of phosphor emission spectra, the PL intensity of the main emission peak was evaluated in comparison with the standard (Kemira, I_{611nm}=100%). Under 254 nm excitation, the phosphor emits mainly in the red domain, i.e. at ~ 590 nm (${}^{5}D_{0} \rightarrow {}^{7}F_{1}$ electronic transition) and ~611 nm (${}^{5}D_{0} \rightarrow {}^{7}F_{2}$

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transition). The excitation radiation influences the emission intensity and even the ratio between the different emission bands.

The phosphor thermal synthesis regime determines the red emission intensity of Y_2O_3 : Eu phosphors (**Fig. 4**). The emission intensity slightly increases with the firing temperature from 74.3% (P19) to 80.2% (P27) and 84.7% (P28) for 254 nm excitation and from 31.4% (P19) to 47.5% (P27) and 48.0% (P28) under 365 nm radiation excitation.

Phosphor morphology and particle size depend on the precursor characteristics and the thermal synthesis regime. The SEM images of phosphor P27 prepared at 1200° C and of the corresponding precursor PG26 are presented in **Fig. 5**.





Fig. 3. Emission spectra of P19 phosphor samples

Fig. 4. Influence of synthesis temperature on the PL emission intensity



Fig. 5. SEM images of precursor (left) and phosphor (right)

Platelets like particle of precursor are converted into conglomerates, with irregular shape, of small particles. As illustrated by the SEM images, the thermal treatment performed without flux leads to a partial conversion of precursor platelets of 1 \div 3 µm to more or less dispersed spherical primary phosphor particles of ~0.5 µm.

The thermal treatment induces a decrease of the particle size due to the decomposition. The incomplete morphological conversion suggests that, during the

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phosphor thermal synthesis, some mineralising agents (fluxes) need to be added to the yttrium-based precursor.

CONCLUSIONS

The wet chemical synthesis method, especially the SimAdd technique, facilitates the control of some preparative parameters for obtaining the yttrium oxalate based-precursor for Y2O3: Eu powders with small particle sizes. XRD and FTIR spectral investigations suggest that the precursor is mainly a hydrated yttrium-ammonium oxalate. Additional PL and SEM measurements revealed that the firing temperature influences the luminescence characteristics and the morpho-structural properties of phosphors particles obtained by the *SimAdd* technique, from precursor prepared with oxalic acid as precipitating reagent. Further experiments are to be performed in order to improve the PL characteristics and powder morphological characteristics.

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Dedicated to professor Gh. Marcu at his 80th anniversary

GROWTH AND CHARACTERISATION OF ZINC SULPHIDE THIN FILMS DEPOSITED ON ITO COATED GLASS

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ABSTRACT. ZnS thin films were grown onto ITO coated glass by chemical bath deposition. The multilayer technique was used in order to prepare ZnS/ITO/glass/ ZnS heterostructures with variable film thickness. Optical properties were investigated by UV-VIS absorption/reflection and fluorescence spectroscopy. The main ZnS films characteristics were correlated with the growing conditions and the annealing regime.

Key words: zinc sulphide; thin films; ITO coated glass; chemical bath deposition

1. INTRODUCTION

Zinc sulphide (ZnS) is a wide, direct band gap semiconductor with interesting optoelectronic properties. A large range of applications exists for thin films of ZnS such as n-window layers of solar cells, electroluminescent displays and other optoelectronic devices [1]. Different methods could be used for ZnS thin films preparation including sputtering, metal organic chemical vapour deposition (MOCVD), atomic layer epitaxy (ALE), pulsed laser deposition, chemical bath deposition (CBD) or electrodeposition [2,3]. Among them, CBD is a simple and inexpensive method and produces uniform, adherent and reproducible films. Moreover, CBD is a low temperature technique and can be used for ZnS deposition onto a variety of substrates.

The aim of this work is to study the influence of different CBD preparative parameters on the quality of ZnS thin films deposited onto ITO coated glass and, especially, on some of their optical characteristics, including their luminescence ability.

2. EXPERIMENTAL PART

ZnS thin films have been grown by CBD method on ITO (indium tin oxide) coated glass pieces provided by Optical Filters Ltd.(UK). The deposition of ZnS was carried out from a mixture of zinc acetate, thiourea, NH_3 aqueous solution, and sodium citrate and twice distilled water. Prior the deposition, the platelets (50mm x 25mm x 1mm) were ultrasonically cleaned with acetone/ethanol mixture and dried. Moreover, all the deposition reagents were purified by some characteristic procedures. The multilayer deposition technique was used in order to prepare

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ZnS/ITO/glass/ZnS heterostructures with variable film thickness. The deposition temperature was 82-86°C and pH mixture was 9.5-10.5. The details of experimental technique have been previously described [4,5]. Most of the heterostructures were obtained in a chemical bath with the standard composition: $[Zn^{2+}] = 0.015$ M; $[C_6H_5O_7^{-3}] = 0.060$; $[NH_3] = 0.300$ M; [thiourea] = 0.15 M. Several samples, for luminescence investigation purpose, were prepared from chemical bath containing a smaller concentration of sodium citrate, namely 0.045 M.

The samples were washed, dried ($100^{\circ}C$) and annealed in nitrogen ($500^{\circ}C$) or air ($400 \div 550^{\circ}C$), using a special protecting system. In the later case, the thermal treatment was performed in special to ZnS-based mixtures containing Cu and Mn salts to generate luminescent ZnS thin films.

Zinc sulphide thin films were characterised by thickness and UV-Vis transmittance and reflectance spectra, as well as some photoluminescence investigations. Prior performing the measurements, ZnS films were partially removed with HCl (1:1) from some ZnS/ITO/glass/ZnS heterostructures to give the corresponding ZnS/ITO/glass heterostructures. Optical investigations were performed by using a UNICAM Spectrometer UV4 and photoluminescence measurements by means of a Perkin Elmer 204 Fluorescence Spectrophotometer (λ_{exc} = 365 nm). The film thickness was evaluated by the microweighing, as indicated in our previous work [6].

3. RESULTS AND DISCUSSION

ZnS/ITO/glass/ZnS heterostructures were prepared from chemical bath containing zinc acetate as zinc source, thiourea as sulphur source, aqueous solution of NH_3 as chelating and pH regulating agent, sodium citrate as chelating agent and doubly distilled water.

The chemical bath deposition process uses a controlled chemical reaction to effect the slow formation of ZnS thin films deposited onto ITO coated glass. The general chemical equation is:

 $Zn(CH_3COO)_2 + (NH_2)_2CS + 2OH \rightarrow ZnS + H_2CN_2 + 2H_2O + 2CH_3COO^{-1}$

CBD method was adapted for multilayer ZnS film formation onto ITO coated glass platelets. With this deposition technique one to five superposed ZnS layers were deposited onto ITO coated glass platelets.

In the above-mentioned conditions, adherent and homogeneous ZnS thin films were grown on both ITO coating and glass substrates to give the ZnS/ITO/glass/ZnS heterostructures. Some ZnS/ITO/glass heterostructures were obtained by the removal of ZnS films from the glass side opposite to the ITO coating

The growing parameters of the as prepared heterostructures are presented in **Table 1**. As expected, increase of zinc sulphide film thickness with the total deposition time could be noticed. The use of a high number of successively deposited layers (coatings) determines the increase of the film thickness. One can note that, the growing rate, calculated by dividing the ZnS film thickness to the total deposition time is increases with the number of coatings. According to our previous work [5,7,8], for the same deposition time, the multilayer grown film is thicker when it deposited onto the glass substrates in comparison with the film formed onto the ITO coated face of the glass platelet. Mention has to be made that, according to the manufacturer measurement, the thickness of the ITO coating is ~20 nm. GROWTH AND CHARACTERISATION OF ZINC SULPHIDE THIN FILMS DEPOSITED ON ITO ...

Table 1.

The growing parameters of some ZnS/ITO/glass heterostructures prepared by
multilaver technique

Samples	Total deposition time	ZnS packing density	ZnS film thickness**	ZnS film growing rate
code	n x m*	(mg/cm ²)	(nm)	(nm/min)
ITO 3.0	0	0	0	0
ITO 3.1	1 x 60 = 60 min	0.13	33	0.55
ITO 3.2	2 x 60 = 120 min	0.33	80	0.67
ITO 3.3	3 x 60 = 180 min	0.60	150	0.83
ITO 3.4	4 x 60 = 240 min	0.80	200	0.83
ITO 3.5	5 x 60 = 300 min	1.06	260	0.87

*Where: n = number of layers, m = deposition time; ** on the ITO coated glass face

UV-Vis absorption /reflection and photoluminescence spectroscopy were used to investigate the optical properties of different ZnS/ITO/glass or ZnS/ITO/glass/ZnS heterostructures. Moreover, in order to prove the ZnS film ability to develop luminescent properties, some of the heterostructures containing relatively thick films were annealed in special ZnS-based mixtures containing copper and manganese doping ions. The photoluminescence behaviour of the as obtained copper and manganese doped ZnS/ITO/glass/ZnS heterostructures, notated ZnS:Cu,Mn /ITO/glass/ZnS. SnS:Cu,Mn, was evaluated from the emission spectra registered under UV excitation.

The transmission spectra (**Fig.1**) of the heterostructures that contains multilayer ZnS films illustrate the high transparency of the ITO coating over the entire visible domain (over 90%). The transparency decreases parallel with the increase of the film thickness. The multiple maxima on transparency curve of some ZnS/ITO/glass heterostructures reveal a good quality of the deposited thin film (**Fig.1**).



Fig. 1. Transmission spectra of some ZnS/ITO/glass heterostructures

Fig. 2. Specular reflectance of ZnS films deposited onto ITO coated glass face

The reflectance spectra of ZnS/ITO/glass heterostructures were registered in order to evaluate the ZnS film optical quality (**Fig. 2**). The spectra registered at 0° incidence indicate the diffuse reflection of the films whereas the spectra obtained at 8° incidence of the visible light illustrate both the specular and diffuse

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reflection properties of the investigated heterostructure. The specular reflectance of some samples, evaluated as the difference between the reflection measured at 8° and 0° incidence, decreases as the number of ZnS layers increases One can note the strong specular reflection properties of the ITO coated glass. A thin film of ZnS (ITO3.1) attenuates the ITO layer reflection and shows only a weak reflection characteristic to the ZnS material. This one is clearly put in evidence for the relatively thick layers (ITO3.5).

Some of the ZnS/ITO/glass and ZnS/ITO/glass/heterostructures were treated thermally either in a nitrogen atmosphere, or in some peculiar conditions that allows the evaluation of the luminescence ability of the CBD obtained ZnS films. **Table 2** presents the thickness and the thermal treatment conditions of some heterostructures containing un-doped or copper-manganese doped ZnS thin films.

The transmittance spectra of the ZnS/ITO/glass heterostructure obtained in 300 min deposition was measured before and after the thermal treatment (**Fig.3**). It is obvious that the post-growing thermal treatment decreases the film transmittance and diminishes the film optical quality.

The emission spectra of copper-manganese doped ZnS/ITO/glass/ZnS heterostructures are depicted in **Fig. 4**. The sample prepared at 400°C shows a very weak luminescence with relatively well evidenced emission bands situated at about 467, 524 and 560 nm. These bands are correlated with the presence of some specific emission centres associated with the self-activated SA-, Cu- and Mn-luminescence in zinc sulphide layer, respectively. The weak luminescence observed for this structure could be explained either by an incomplete incorporation of the activators into the ZnS lattice generating a small number of luminescence centres, or to a poor quality of the films deposited onto ITO substrate.

In order to verify the luminescence ability of ZnS thin films, attempts were made to incorporate Cu-Mn activators by annealing the system at 550°C. The sample obtained using this thermal synthesis regime, shows a relatively strong luminescence situated mostly into the yellow spectral region. The main emission peak is observed at 577 nm, close to the theoretical value (580 nm) originated into the Mn-centres. The higher annealing temperature is in the favour of Cu-and Mn activator incorporation.

Table 2.

Sample code	Thickness (nm)**	Doping and thermal treatment	Heterostructure
ITO3.5TT	260	500°C, N ₂	ZnS/ITO/glass
ITO11.3	320	Cu-Mn doping, 400°C	ZnS:Cu,Mn/ITO/glass/ZnS:Cu,Mn
ITO12.1	310	Cu-Mn doping, 550°C	ZnS:Cu,Mn/ITO/glass/ ZnS:Cu,Mn

Thickness and thermal treatment conditions of some ZnS containing heterostructures

*where: n = number of layers, m = deposition time; ** mean value

The better yellow-orange emission observed for the heterostructure prepared at relatively high temperature reveals a good luminescence ability of zinc sulphide layer.

The luminescence performances of the ZnS/ITO/glass/ZnS heterostructures doped with copper and manganese are predominantly sensitive to the annealing regime and the doping conditions.

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Fig. 3. Transmission spectra of some ZnS/ITO/glass samples before and after the thermal treatment



4. CONCLUSIONS

The bath composition and the special CBD technique used in our experiments proved to be convenient for the deposition of adherent and homogeneous ZnS films with controllable thickness onto the ITO coated glass substrate. The growth parameters influence the optical properties of the as deposited thin films.

UV-Vis absorption /reflection investigations illustrated the quality of the as prepared ZnS/ITO/glass/ZnS heterostructures. The photo-luminescence measurements on some copper and manganese doped ZnS/ITO/glass/ZnS heterostructures proved the ability of ZnS films to develop good light emitting properties.

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Dedicated to professor Gh. Marcu at his 80th anniversary

BIS-PHENOTHIAZINYL-PHENYL-METHANE DERIVATIVES

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ABSTRACT. The synthesis of some *bis*-(phenothiazin-3-yl)-phenylmethane derivatives by the condensation of phenothiazine with aromatic aldehydes was investigated. The structural assignments for the new compounds were based on NMR, IR and UV-Vis spectroscopy.

INTRODUCTION:

The chemical reactivity of phenothiazine towards electrophilic substitution reactions was demonstrated in numerous examples of N- or C- substituted derivatives preparations. The transmission of the electronic effects between the two heteroatoms and the two benzene rings is very efficient and electrophilic substitution occurs easily. Rather mild electrophiles can be used to accomplish the substitution reaction in position 10 (N-alkylphenothiazines [1], N-acylphenothiazines [2]) or in positions 3 and 7 (C-halogenophenothiazines [3-5], C-nitrophenothiazines [6.] C-alkylphenothiazines [7,8]). Dyes formation by the condensation of phenothiazine with aromatic ketones (such as Michler's ketone) was also reported [9].

The paper presents the condensation reaction of phenothiazine with aromatic aldehydes in the presence of acid catalysts.

RESULTS AND DISCUSSIONS

The condensation reaction of phenothiazine **1** with aromatic aldehydes (benzaldehyde, *o-, m-* and *p*-nitrobenzaldehyde) in acid media was performed. The mild electrophile generated by the aldehyde in the presence of methanesulfonic acid determines the substitution of the phenothiazine ring in position 3 (activated by the electron donor effect of the nitrogen heteroatom). In boiling ethanol solvent the reaction product **4** has a low solubility and easily separates from the reaction mixture. Scheme 1 presents an overview of the chemical reactions involved.

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Scheme 1.

The structure assignment for compounds **4a-d** was performed by H-NMR, FTIR and UV-Vis spectroscopy.

The IR spectroscopic investigations of the reaction products showed that the alcohol intermediate **2** was not separated under the reaction conditions employed (no characteristic absorption bands due to C-O and O-H bonds stretching vibrations were recorded for the separated reaction product).

During the reaction progress, a green color appears in to the reaction mixture, possibly due to the competing elimination reaction generating the compounds **3** (characterized by a fully conjugated π electrons system). The compounds **3** are soluble in ethanol.

The *bis*-(phenothiazin-3yl)-phenylmethane **4a** and its nitro-substituted derivatives **4b-d** separated from the boiling reaction mixture as precipitates which were filtered. The H-NMR spectra, showed for the aliphatic proton a characteristic chemical shift situated in the range of 5.1-5.8 ppm. The most deshielded aliphatic proton appears in the structure of the *orto*-nitro-substituted derivative **4b** (δ =5.8 ppm), while the signal of the same proton in the structure **4a** is situated at 5.1 ppm. Intermediate values (5.4 ppm) were recorded for the compounds **4c-d** containing a nitro group situated in position *meta* or *para*.

EXPERIMENTAL:

The chemical reagents and the solvents were purchased from Merck (for synthesis purity)

The IR spectra were recorded on a FTIR Bruker Vector 22 spectrometer.

The UV-Vis spectra were recorded on a UNICAM Helios β spectrometer.

The H-NMR spectra were recorded on a Bruker Avance 300 MHz spectrometer.

General procedure for the condensation of phenothiazine with aromatic aldehydes.

Phenothiazine (2g, 1 mmol) was solved in ethanol (120 mL), methanesulphonic acid (0.5 mL) and an ethanolic solution of the aromatic aldehyde (0.5 mmol) were added to the clear phenothiazine solution. The reaction mixture was heated under

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vigorous stirring to reflux for several hours. During the reaction progress, a green color appears and a precipitate starts to accumulate.

The reaction mixture thus obtained was cooled down at room temperature, the precipitate was filtered out and then washed several times with cool ethanol. The reaction product is a grey-green powder highly insoluble in toluene, acetone and chloroform, soluble in THF.

Bis-(phenothiazin-3-yl)-phenylmethane 4a

Reaction time: 12 hours, green powder , m.p. 251 $^{\circ}\text{C}$, yield 60 %. IR [cm⁻¹]: 3400, 3100, 1600, 1462, 1300, 803, 741, 604

UV [nm]: 248, 320.

H-RMN (300 MHz, DMSO-d₆) δ : 7.3-6.1 ppm, m 19H (phenyl and phenothiazine unit), 5.1 ppm, s, 1H (CH)

Bis-(phenothiazin-3-yl)-2-nitrophenylmethane 4b

Reaction time: 40 hours, grey powder , m.p. 278 °C, yield 48%.

IR [cm⁻¹] 3400, 3100, 1514, 1339, 810, 742, 650

UV [nm]: 248, 320.

H-RMN (300 MHz, DMSO-d₆) δ : 8.1-6.6 ppm, m 18H (*o*-phenylene and phenothiazine unit, 5.8 ppm, s, 1H (CH).

Bis-(phenothiazin-3-yl)-3-nitrophenylmethane 4c

Reaction time: 20 hours, brown powder, m.p. 264 °C, yield 55 %.

H-RMN (300 MHz, DMSO-d₆) δ: 8ppm, d, 2H, 7.4 m 2H, (*m*-phenylene unit),

6.1ppm, m, 14 H (phenothiazine unit), 5.3 ppm, s, 1H (CH).

Bis-(phenothiazin-3-yl)-4-nitrophenylmethane 4d Reaction time: 15 hours, grey powder , m.p. 255 °C, yield 61%. IR: [cm⁻¹]: 3400, 3100, 1600, 1514, 1339, 809, 742, 690. UV: 250 nm, 324 nm

H-RMN (300 MHz, DMSO-d₆) δ : 8.6 ppm s, 2H (NH), 7.34 d 2H, 8.1, d, 2H (*p*-phenylene unit), 6.6–6.9 ppm, m, 14 H (phenothiazine unit), 5.4 ppm, s, 1H (CH).

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Dedicated to professor Gh. Marcu at his 80th anniversary

MOLECULAR PACKING OF SOME CAROTENOIDS IN LANGMUIR MONOLAYERS AT THE AIR/WATER INTERFACE

OSSI HOROVITZ*, MARIA TOMOAIA-COTISEL*

ABSTRACT. The characteristic molecular areas were determined from surface pressure versus molecular area isotherms of three carotenoids: β , β -carotene-4-one (echinenone, ECH), β , β -carotene-4,4'-dione (canthaxanthin, CAN) and 4,4'-diapo- ψ , ψ -carotene-4,4'-dial (APO), spread and compressed as Langmuir monolayers at the air/water interface. Quantum chemical semi-empirical SCF MO calculations (AM1 and PM3) are performed for the optimized geometries of carotenoid molecules and similar theoretical results are obtained by both methods. The characteristic surface molecular areas are discussed in terms of molecular packing in Langmuir monolayers and intermolecular interactions, respectively. The orientation of these carotenoid molecules in the Langmuir monolayers is discussed by using a rotating rigid plate model and optimized molecular geometries.

Keywords: carotenoids, Langmuir monolayers, molecular packing, molecular structure, semi-empirical MO calculations.

INTRODUCTION

Langmuir monolayers are obtained by spreading amphiphilic organic molecules using an appropriate solvent or a mixture of organic solvents at the air/water interface [1-3]. Generally, these monolayers are compressed up to their collapse and they are employed as model systems in studying surface phenomena, associated with various applications in science and technology [4]. In particular, carotenoids can be potential key elements for molecular electronic devices. Also, carotenoids are essential biological active molecules, which can influence or modulate the properties of cellular and subcellular structures [5, 6]. Therefore, the behavior of Langmuir monolayers of carotenoids is of interest and is investigated at the air/water interface both in pure state [7-10] and in mixtures with lipids [11-14] or electrolytes [15-18]. The obtained data show that the local ordering and molecular packing of carotenoids in Langmuir monolayers at the air/water interface behavior of the same aqueous phase and at similar conditions for compression rate and spreading rate.

The objective of this paper is to investigate experimentally and theoretically how the equilibrium intermolecular interactions and hydrophilic and hydrophobic forces affect the molecular packing of some carotenoids in Langmuir monolayers at the air/water interface. In the following the rotating rigid plate model previously proposed by us [19] for carotenoid molecules is reconsidered, and used to explain the molecular

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packing of the three chosen carotenoids for this study. Optimized geometries of the molecules are used, as resulted from semi-empirical MO computations. Finally, the obtained results are summarized in the conclusion section of this paper.

THE ROTATING RIGID PLATE (RRP) MODEL OF CAROTENOIDS

In order to correlate the structural and geometrical characteristics of carotenoid molecules with their surface properties as monolayers at the air-water interface, the rotating rigid-plate model of carotenoid molecules was previously proposed [19-21]. In the framework of this model, the molecule (all-*trans* isomer) is considered to be rigid (due to its delocalized π -bond system), but free rotation

about the simple o-bonds has also been taken into account, the most stable conformation being assumed. For the bond lengths and bond angles, experimental data (from X-ray analysis) have been used, when available (for canthaxanthin [22]); otherwise they have been approximated from the bond lengths and covalent radii given in chemical tables and the bond angles from the hybridization types.

The hydration of the polar headgroup of the molecule, immersed in water, was modeled by the inclusion of hydrogen-bridged water molecules. Two H_2O molecules are considered to be linked to the C=O group, thus anchored in the water phase.

The carotenoid molecule is seen as a parallelepiped; its length, c, is taken as the length of the chain axis (c'), including the hydration water molecules (see Fig.1 for canthaxanthin molecule). The length of the headgroup axis (a'), and also including hydration water, a, is the width of the molecular plate, while b represents its thickness – given mainly from the out-of-plane hydrogen and carbon atoms [21].

In the estimation of the molecular areas in the monolayer at the air/water



Fig.1. Rigid plate model of the canthaxanthin molecule [1]

interface, arguments were given for a calculation in which the molecule is assumed to perform a free rotation about the vertical chain axis, and the resulting vertical cylinders adopt a tetragonal close packing [19, 21]. This does not necessarily imply a real rotation, but also a random orientation of the head group axis in the interface plane. Therefore, since b < a, the mean area occupied by a molecule in the interface layer should be $A_4 = a^2$, instead of $A_p = ab$, corresponding to a close packing of vertical rigid plates with parallel orientation.

Another estimated characteristic of the monolayer was the immersed fraction α of the molecules, defined as the ratio between the length of the part of the chain immersed in water, as determined by the hydration, c_w (Fig.1) and the total chain length, c:

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$$\alpha = c_{\rm w}/c \tag{1}$$

The free part of the chain, c_f is therefore $c-c_w$ For echinenone and canthaxanthin, the values obtained from this model [19] were: a = 0.78 nm, b = 0.49 nm, c = 3.2 nm, $c_w = 0.7$ nm, $\alpha = 0.22$ and $A_4 = 0.61$ nm², $A_p = 0.38$ nm².

The present paper deals with the application of the RRP model to three carotenoid pigments, namely β , β -carotene-4-one (echinenone, ECH), β , β -carotene-4,4'-diane (canthaxanthin, CAN) and 4,4'-diapo- ψ , ψ -carotene-4,4'-dial (4,4'-diapolycopenedial, APO) (Fig. 2), using their molecular structure, as resulted from semi-empirical MO calculations (AM1 and PM3).



EXPERIMENTAL PART

 $C_{30}H_{36}O_2$

Fig.2. Molecular structure of carotenoids: β , β -carotene-4-one: ECH, β , β -carotene-4,4'-dione: CAN and 4,4'-diapo- ψ , ψ -carotene-4,4'-dial: APO.

The three natural carotenoid pigments (ECH, CAN and APO) used here were supplied by Hoffmann-La Roche and present all-trans configuration. The high purity spreading solvents, benzene pro-analysis or a mixture of benzene and 2-4% absolute ethanol, were purchased from Merck. Known amounts of the compounds dissolved in the spreading solvents were placed by means of a Hamilton syringe or a micropipette on the air/water interface. The water subphase was double distilled water with a resistivity of 18 Mohm cm⁻¹. Compression of the monolayer was started after about 10 minutes, allowed for solvent evaporation. Then, the spread monolayer of each carotenoid compound was compressed at a chosen rate of compression in the interval from 0.01 to 0.03 nm²/(molecule min), by using the Langmuir equipment KSV 5000. The reproducible compression isotherms, in terms of surface pressures versus molecular areas, were recorded at 20 °C as described elsewhere [5]. The surface pressure was measured by the Wilhelmy method within the error of \pm 0.2 mN/m. The compression curves were reproducible within 0.02 - 0.04 nm²/molecule. The results do not depend on the compression rate in the region used. Each isotherm given in this paper represents the mean of at least 10 different recordings.

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COMPUTATIONAL PART

MO calculations were performed, regarding σ and π electrons, on the three carotenoids (ECH, CAN and APO) in their all-trans forms [23]. Computations were made at the restricted Hartree-Fock (RHF) level using two semi-empirical SCF MO methods: Austin Model 1, AM1 [24] and the Parametric Model number 3, PM3 [25], by means of HyperChem 7.5 software package [26]. The two methods are based on the Neglect of Differential Diatomic Overlap (NDDO) integral approximation, and differ by their parameterization. The computation options were: total charge: 0; spin multiplicity: 1 (singlet); state: lowest. The geometries of the molecules were optimized by the Polak-Ribiere (conjugate gradient) algorithm approach. The SCF convergence limit was of 0.042 kJ/mol, and RMS gradient was of 4.18·10⁹ kJ/(m×mol).

For the optimized geometry, energetical parameters (total electronic energies, enthalpies of formation), electron distributions (charge densities and bond orders) and geometrical parameters (bond lengths, bond angles, torsion angles) were obtained [23]. Both computational methods gave quite similar results for the geometry of the molecules. The first inertial axis of the molecule is roughly going along the conjugated chain of the molecules, while the second axis is perpendicular to it in the molecular plane.

RESULTS AND DISCUSSION

The **compression** isotherms, in terms of surface pressure (π , mN/m) versus mean molecular area (A, nm²) curves for three carotenoids, viz. β , β -carotene-4-one (ECH), β , β -carotene-4,4'-dione (CAN) and 4,4'-diapo- ψ , ψ -carotene-



Fig.3. Compression isotherms: surface pressure versus mean molecular area for carotenoids: β_{β} -carotene-4-one (ECH), β_{β} -carotene-4,4'-diane (CAN) and 4,4'-diapo- ψ , ψ -carotene-4,4'-dial (APO) spread as monolayers at the air/water interface, at 20 °C.

4,4'-dial (APO), spread at the air/water interface, are presented in Fig. 3.

From these compression isotherms, surface properties were determined, namely: the limiting molecular area A_0 (see Fig. 3), obtained by extrapolating to π = 0 the high pressure linear portion of the compression isotherm; A_c , is the collapse area and the corresponding surface pressure noted π_c , the collapse pressure (Table 1).

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Table 1.

0.40

Carotenoid	$\pi_c \ (mN \ m^{-1})$	$A_0^{}(\text{nm}^2)$	$A_{\rm c}({\rm nm^2})$					
ECH	32	0.60	0.48					
CAN	36	0.60	0.44					

32

APO

Surface characteristics of the three carotenoids studied.

The **optimized molecular geometry**, as resulted from the PM3 calculations, is visualized for the three molecules in the plane of the first two inertial axes in Fig.4.

0.60



Fig. 4. Optimized geometries (PM3 calculation) for the three carotenoid molecules, represented in the plane of the first and second inertial axes.

For a discussion of the **molecular orientation and packing at the air/water interface,** on these models the length of the chain was measured, as the maximum size in the direction of the first axis: 2.93 nm (ECH), 2.94 nm (CAN), and 3.02 nm (APO). Since in these representations, distances between atomic nuclei are obtained, the covalent radii of the end atoms (H for ECH and CAN, O for APO) should be added, what gives for the lengths c' (Fig. 1) the values: 2.99 nm (ECH), 3.00 nm (CAN), and 3.15 nm (APO).

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Following the considerations given in the earlier treatment [19], about 0.1 nm have to be added to *c*' for the lower water molecule bonded to the C=O group in ECH and CAN, thus obtaining a *c*-value of about 3.1 nm for these two molecules. As for the immersed part of these chains, given by the vertical projection of the two O····H-O bridges and of the other two O-H bonds, it was estimated to be $c_w = 0.7$ nm, the immersed fraction (eq. 1) being $\alpha = 0.23$.



Fig.5. The shape of the lower part of the APO-molecule and the size of its hydrated polar head group for a vertical orientation of the molecular axis – 1st inertial axis (A) and for an angle of 20° to the vertical (B)

For the APO molecule, a similar model was sketched in Figure 5A, for the lower part of the molecular chain. An angle of 60° of the C=O bond to the air-water interface was assumed, corresponding to a vertical orientation of the conjugated chain, as in earlier MO studies on this compound [27, 28]. Adding 0.23 nm for the vertical projection of the lower O····H-O bridge, we obtain for the total (hydrated) chain length *c* = 3.38 nm. As for the immersed part of the chain, besides the contribution of the lower water molecule (0.23 nm + oxygen radius), we have to consider the vertical projection of the O-H bond of the second water molecule and the H radius. This gives a *c*_w value of about 0.43 nm and an immersed fraction $\alpha = 0.13$, less than for the other two molecules.

A previous investigation on these three carotenoid molecules [23], correlating surface properties with MO calculated quantities, viz. dipole moments, has suggested that the APO molecules should deviate from the vertical, i.e. to be inclined in order to decrease the angle of the C=O bond to the air/water interface. This orientation would allow for an increased interaction with the water molecules. The angle should be about 40°, i.e the deviation of the molecule axis from the vertical, of about 20° (Fig. 5B). For this orientation, the depth of the hydration layer, given by the two hydrogen-bonded water molecules, would be $c_w = 0.56$ nm and the length of the immersed part of the molecular chain about 0.6 nm. The total length of the hydrated chain is given by adding to *c*' the contribution of the lower water molecule

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to the hydration, about 0.29 nm, and is therefore c = 3.44 nm. The immersed fraction α becomes now 0.17. Thus the inclination allows for a better hydration of the APO molecule.

The areas occupied by the molecules in the monolayer can be estimated from the molecular volumes, *V*, calculated as QSAR properties by the HyperChem software, divided by the calculated length of the molecule *c*'. These values are given in Table 2 and are in quite good agreement with the experimental values of extrapolated molecular areas, A_0 [19], obtained by extrapolating the high pressure linear portion of the surface pressure – area curves to pressure $\pi = 0$.

Table 2.

Molecule	<i>c</i> ' (nm)	$V(nm^3)$	V/c'	<i>a</i> (nm)	b	A ₄	Ap	Ao	Ac
			(nm ²)		(nm)	(nm ²)	(nm ²)	(nm ²)	(nm ²)
ECH (1)	2.99	1.8334	0.613	0.74	0.56	0.548	0.414	0.60	0.48
(2)	3.10			0.87		0.757	0.487		
(3)	3.2			0.78	0.49	0.608	0.38		
CAN (1)	3.00	1.8292	0.610	0.75	0.52	0.562	0.39	0.60	0.44
(2)	3.10			0.88		0.774	0.458		
(3)	3.2			0.78	0.49	0.608	0.38		
APO (1)	3.15	1.4923	0.474	0.60	0.24	0.36	0.144	0.60	0.40
(2)	3.38			0.585		0.342	0.14		

Calculated and experimental molecular areas in monolayers

(1) without hydration; (2) with hydration; (3) estimated in [1]

On the assumption that the dimensions of the "molecular plate", are in first approximation colinear with the inertial axes: *c* with the 1st, *a* with the 2nd and *b* with the 3rd axis, we can evaluate these dimensions. As above for *c*, from the representations of the optimized geometries of the molecules in the planes of the first and second inertial axes (Fig. 4), in the plane of the first and third axes or of the second and third inertial axes (Fig. 6) the other two dimensions of the molecules, *a* and *b*, can be measured. In Table 2, these values are given as read from the diagrams with addition of the radii of the end atoms (1), and with addition of the hydration water molecules (2). The values estimated in [19] are also given (3). For the APO molecule, the value (2) was obtained from the model (Fig.5A) of the hydrated head group.

The molecular areas have been calculated from these 3 kinds of molecular dimensions:

(i) from the model of the freely rotating molecular rigid plates, with tetragonal close packing, $A_4 = a^2$;

(ii) from the model of close packing of parallel oriented rigid plates, $A_p = ab$ These values are also given in Table 1. The A_c values are the molecular areas at the collapse of the carotenoid monolayer.

The A₄ values estimated in [19] for ECH and CAN are in good agreement with the experimental A₀ values. The A₄ values calculated form the values (2) are larger than the values reported in [19] (3). The explanation is that the *a*-values (3) were estimated for the head group only, while the *a*-values (2) refer to the "width" of the entire molecule, in its real geometry. The same does apply to the "thickness" *b*-values (1) and (3).

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Fig.6. The carotenoid molecules, represented in the plane of the first and third inertial axes (A) and of the second and third inertial axes (B). The scales of the representations are different.

The A_p values (2), calculated from the dimensions of the hydrated ECH and CAN molecules, are in much better agreement with the molecular collapse areas than the estimated values (3); moreover, they give also the right order (ECH > CAN), while the values estimated in [19] make no difference between the two molecules. It can be assumed that near the collapse pressure, the free rotation of the molecules is hindered and the head group axes tend to assume a parallel orientation.

The APO molecule represents a different case. The lower immersed fraction for this substance brings about that higher energies are necessary to perform a vertical orientation of the chain axes. The molecules seem to keep their inclined orientation even at the collapse, due to the weak intermolecular interactions, as suggested in [23]. Therefore, the molecular areas calculated for a vertical orientation are much lower than the experimental A_o and A_c values.

CONCLUSIONS

Three carotenoid molecules, ECH, CAN, and APO, have been studied in Langmuir monolayers at the air/water interface. The surface characteristics, like limiting molecular areas, collapse areas and collapse pressures are well correlated with the carotenoid molecular structures and the molecular orientation at the air/water interface. The obtained data are explained by the interplay of the intermolecular hydrophilic and hydrophobic forces which lead to stable Langmuir monolayers, with

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the rotating molecules tetragonally packed in the liquid state of the monolayers (rotating rigid plate model of the carotenoid molecules). At collapse, the non-rotating carotenoid molecules are close packed in Langmuir monolayers.

Quantum chemical semi-empirical SCF MO calculations (AM1 and PM3) are performed for the optimized geometries of carotenoid molecules. From the obtained geometry of the molecules, the parameters for the rotating rigid plate model of the molecules could be estimated, in good agreement with the experimental values for the limiting molecular area A_0 and the collapse area A_c . As regards the orientation of these carotenoid molecules in the monolayer, for the echinenone and canthaxanthin molecules a perpendicular direction of the molecular axis to the air/water interface is found, while for the 4,4'-diapo- ψ , ψ -carotene-4,4'-dial (APO) molecule a deviation from the vertical is suggested.

It would be of interest to study other molecules also having the poly-ene chain attached to beta-ionone rings or to a flexible moiety like an alkyl chain.

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Dedicated to professor Gh. Marcu at his 80th anniversary

HEMES REVISITED BY DENSITY FUNCTIONAL APPROACHES. 2. A PARADIGM FOR AXIAL LIGATION IN HEMOPROTEINS

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ABSTRACT. Reported here are DFT descriptions of dioxygen reduction at the active site of cytochr]ome cd_1 nitrite reductase, an enzyme with a histidine-ligated heme active site, known to act as cytochrome oxidase. Also explored is the possibility of nitric oxide reduction by histidine-ligated hemes. The energetics of these two processes correlate well with previous findings on other hemoproteins, and allow further elaboration on the newly proposed "thiolate obstruction" theory [Silaghi-Dumitrescu, Eur. J. Inorg. Chem. 2003, 1048]. This theory, orthogonal if not opposed to the classical "thiolate push" dogma [Dawson et al., J. Am. Chem. Soc. 1976, 98, 3707], argues that efficient reduction of diatomics such as dioxygen and nitric oxide is more readily accomplished by histidine-ligated hemes than by thiolate-ligated hemes. Generalizing, for reductive processes involving a small diatomic as the sole substrate, neutral ligands (e.g., histidine in cytochrome oxidases and heme oxygenase, lysine in cytochrome c nitrite reductase) are found to always be preferable over anionic ligands. By contrast, in enzymes designed to deal with more than one substrate, anionic ligands are preferable (e.g., cysteine or tyrosine in monooxygenases), since they allow the safety switches needed to avoid uncoupling in their race against entropy.

INTRODUCTION

Proteins such as cytochrome P450 (P450), horseradish peroxidase (HRP) or hemoglobin are all known to bind and/or reduce (activate) dioxygen at their heme active sites, in the ferrous form.[1-11] The thiolate-ligated ferrous heme of P450 binds dioxygen and promotes proton-dependent O-O bond cleavage following a one-electron reduction.[1] By contrast, the ferrous histidine-ligated heme of hemoglobin only binds dioxygen in a reversible manner.[12] Hemoprotein ferric-hydroperoxo complexes decay via proton-assisted heterolytic cleavage of the O-O bond to yield water and a [Fe=O]³⁺ unit (known as Compound I), where the iron is considered to be in the formal oxidation state +4 and a further oxidizing equivalent is proposed to be delocalized onto the porphyrin. Two active-site "distal" residues, a histidine and an arginine, are crucial in promoting O-O bond cleavage in the ferric-hydroperoxo complex of canonical (histidine-ligated) peroxidases. The hemoglobin active site lacks the distal arginine and is much less efficient than HRP at activating peroxide.[1-10] The heme-thiolate active site of P450 does not contain the histidine-arginine catalytic pair of HRP, yet still cleaves the O-O bond of its ferric-hydroperoxo complex

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(using water molecules as the source of protons). These differences between thiolate and histidine-ligated hemes are traditionally rationalized in terms of the thiolate ligand being able to push electron density more efficiently onto the iron-dioxygen moiety, thereby facilitating O-O bond cleavage ("the push effect").[10, 11, 13]

Notable exceptions to the thiolate/histidine rule also exist. Cytochrome oxidases, which accomplish four-electron reduction of dioxygen to water, feature histidine and not thiolate ligands.[14] Heme oxygenase, which reduces dioxygen to hydroperoxide, also features a histidine ligand.[15]

Nitric oxide reduction by thiolate-ligated heme (in cytochrome P450 nitric oxide reductase, P450nor) and by lysine-ligated heme (in cytochrome c nitrite reductase, CcNIR) may either involve two one-electron reduction steps, $[FeNO]^{3+} \rightarrow [FeNO]^{2+} \rightarrow [FeNO]^{+}$, or one two-electron reduction step, $[FeNO]^{3+} \rightarrow [FeNO]^{+}$; both mechanisms involve concomitant or subsequent protonation of the NO ligand.[16] The one-step mechanism is now generally accepted for P450nor (where subsequent addition of a second NO molecule results in generation of N_2O), whereas the twostep mechanism is active with CcNIR (where the NO ligand is subsequently further reduced to NH₃ and H₂O).[16, 17] The choice for a one-step, 2-electron mechanism in P450nor was rationalized by differences in thermodynamics: the thiolate heme-NO adduct is more electron rich and therefore a second one-electron reduction would be harder to accomplish than with a lysine-ligated heme. Indeed, this second one-electron reduction was found to be, at least in gas-phase calculations, energetically uphill with thiolate and energetically downhill with lysine. Thus, although the "thiolate push" dogma[1, 13] would link thiolate axial ligands to more facile activation of diatomics, we found the opposite to be true for nitric oxide activation ("thiolate obstruction"). Accordingly, the electrons required for nitric oxide reduction in thiolate-ligated P450nor are directly supplied by the hydride (2-electron) donor NADH, whereas in CcNIR the electrons are supplied by neighboring hemes, in a one-electron fashion.[16, 17]

The "thiolate obstruction" theory further developed upon comparing oneelectron reduction of formally ferrous-dioxygen species to formally ferric-peroxo.[11, 16] There too, gas-phase reduction was thermodynamically uphill with anionic ligands (thiolate, phenoxide, imidazolate), but thermodynamically downhill with a neutral ligand (imidazole, or no ligand at all). When the same calculations were performed in a solvent of ε -4.3 rather than in gas-phase, thus mimicking an enzyme active site, the differences between histidine and anionic models were diminished to some extent, but the histidine still remained clearly more adept at favoring one-electron reduction of the ferrous-dioxygen adduct.[11] P450, which features an anionic thiolate ligand to the heme, activates dioxygen specifically for oxygen atom insertion into an organic substrate (monooxygenase chemistry). It is then desirable that dioxygen reduction only occur in the presence of substrate, to avoid the wasteful/toxic reduction of dioxygen known as uncoupling.[1] The thiolate, and anionic ligands in general, thus offers the possibility of an extra safety switch, that ensures maximum catalytic activity in systems fighting entropy (i.e., relying on the concomitant presence of more than one substrate at the active site).

Cytochrome cd_1 nitrite reductase (cd1NIR) is known to efficiently reduce dioxygen to water at its heme d_1 active site, thus acting as a soluble cytochrome oxidase.[18] The axial heme ligand in cd1NIR is a histidine. In line with our previous findings on heme *b* models, we expect one–electron reduction of the cd1NIR ferrous-

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dioxygen adduct to be energetically very favorable. To verify this, and to investigate the extent to which the heme d_1 may alter the properties of the ferrous-dioxygen adduct and of its reduced congeners, we report here DFT geometry optimization results on models of the cd1NIR ferrous-dioxygen, reduced ferrous-dioxygen, and ferric-hydroperoxo adducts.

Also elaborating on our "thiolate obstruction" theory, we report DFT geometry optimization results for models consisting of a histidine-ligated *b* heme coordinated by nitric oxide, in three oxidation states: Fe(III)-NO, Fe(II)-NO, and Fe(II)-NO; for the latter model, the two protonated states, Fe(II)-N(H)-O and Fe(II)-N-O-H are also investigated. These heme *b* models mimic the nitric oxide adducts of myoglobin, and, in addition to corroborating the "thiolate obstruction" theory, they provide insight into myoglobin's ability to perform nitric oxide reductase chemistry.

METHODS

Geometries for all models were optimized without any constraints with the UBP86 functional, which uses the gradient-corrected exchange functional proposed by Becke (1988),[19] the correlation functional by Perdew (1986),[20] and the DN** numerical basis set (comparable in size to 6-31G**) as implemented in Spartan.[21] For the SCF calculations, a fine grid was used, and the convergence criteria were set to 10⁻⁶ (for the root-mean square of electron density) and 10⁻⁸ (energy), respectively. For geometry optimization, convergence criteria were set to 0.001 au (maximum gradient criterion) and 0.0003 (maximum displacement criterion). Partial atomic charges and spin densities were derived from Mulliken population analyses.

Low-spin states (S=0, S=1/2) were assumed for all models, in line with previous experimental and theoretical findings.[11, 16, 18, 22]

RESULTS AND DISCUSSION

Table 1 and Figure 1 show geometry optimization results for cd1NIR models. Overall, the optimized geometries are consistent with crystal structure of the ferrousdioxygen cd1NIR.

The models in Figure 1 feature a non-planar heme. This is consistent with our previous findings[11] on dioxygen/peroxo heme *b* models. Conversely, the heme was calculated to be essentially planar in the nitric oxide and nitrite adducts of cd1NIR.[18, 23] Data in Table 1 are very similar to those previously reported on equivalent heme *b* models,[11] thus suggesting that the unusual heme d_1 at the active site of cd1NIR does not alter the heme-iron-dioxygen/peroxo chemistry to any significant extent. Importantly, one-electron reduction of the formally ferrous-dioxygen species is still significantly "exothermic", confirming our "thiolate obstruction" theory. Consistent with previous findings[11] on related heme systems, $[FeO_2]^{2+}$ cd1NIR appears as a ferrous-dioxygen/ferric-superoxo hybrid. $[FeO_2]^+$ cd1NIR features one unpaired electron on the oxygen atoms and in this respect is best described as a superoxo adduct rather than peroxo. Assigning the iron in $[FeO_2]^+$ as ferrous may at this stage be tempting; however, a ferric-superoxo description (with one extra electron delocalized onto the porphyrin, and with the iron and porphyrin unpaired electrons strongly covalently coupled) is more likely.[11]

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Figure 1. Geometry-optimized structures for (from left to right): formally ferrous-dioxygen ([FeO₂]²⁺), ferric-peroxo ([FeO₂][†])), and ferric-hydroperoxo ([FeO₂H]²⁺) cd1NIR adducts.

Table 1.

Energies (a.u.), distances (Å), partial atomic charges and spin densities (the latter shown in parentheses) for formally ferrous-dioxygen ($[FeO_2]^{2^+}$), ferric-peroxo ($[FeO_2]^+$)), and ferric-hydroperoxo ($[FeO_2H]^{2^+}$) cd1NIR adducts.

	energy	Fe-O	0-0	Fe-N	Fe	01	O2	OOH
[FeO ₂] ²⁺	-2780.07420	1.78	1.28	2.11	0.48	-0.02	-0.15	-0.17
[FeO ₂] ⁺	-2780.16461	1.90	1.31	2.15	0.46	-0.13	-0.24	-0.37
					(-0.07)	(0.42)	(0.52)	(0.94)
[FeO ₂ H] ²⁺	-2780.69712	1.82	1.44	2.06	0.49	-0.19	-0.22	-0.07
					(0.56)	(0.34)	(0.13)	(0.47)

Table 2 and Figure 2 show geometry optimization results for the myoglobin iron-nitric oxide adducts. These results are consistent with our previous reports on similar models with thiolate, lysine, or with no axial ligand instead of the imidazole.[16] Importantly, one-electron reduction of the formally Fe(II)-NO species is significantly "exothermic", similar to ammonia-ligated models but unlike thiolate-ligated models – in line with the "thiolate obstruction" theory.[16]



Figure 2. Optimized geometries for heme-histidine [FeNO]⁶, [FeNO]⁷, [FeNO]⁸ (top row, from left to right), [FeNHO]⁸, and [FeNOH]⁸ adducts (bottom row, from left to right).

The [FeNO]⁶ model features a significant amount of positive charge on the NO ligand and is thus best described as Fe^{2+} -NO⁺, in line with previous assignments.[16] The [FeNO]⁷ model features more electron density on the NO as well as a bent Fe-N-O unit, which is consistent with a Fe^{2+} -NO/Fe³⁺-NO⁻ hybrid – again as previously

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assigned in related models.[16] The [FeNO]⁸, [FeNHO]⁸, and [FeNOH]⁸ models all clearly feature an NO⁻ ligand, implying a ferrous center as previously assumed.[16] However, we note that, as in all other heme complexes examined to date,[11, 16, 22, 24] the bulk of the two electrons upon going from [FeNO]⁶ to [FeNO]⁸ is in fact not found on the NO, even though formal changes in oxidation state of the NO ligand from +1 to -1 cannot be negated. There is thus a distinction between *formal oxidation state*, which changes for NO from +1 to -1 upon going from [FeNO]⁶ to [FeNO]⁸, and *electron density*, which in fact varies less on the NO ligand and more on the porphyrin – even though the latter cannot be claimed to undergo any formal change in oxidation state.[22]

Table 2.

	energy	Fe-	N-O	Fe-	Fe-	Fe	N	0	NO(H)
		NO		Ν	N-				
					0				
[FeNO] ⁶	-	1.65	1.15	2.05	179	0.48	0.28	0.01	0.29
	2608.96248								
[FeNO] ⁷	-	1.75	1.19	2.15	140	0.49	0.13	-0.13	0.00
	2609.16404					(0.57)	(0.26)	(0.15)	(0.41)
[FeNO] ⁸	-	1.81	1.21	2.41	123	0.51	0.02	-0.22	-0.20
	2609.21773								
[FeNHO] ⁸	-	1.79	1.24	2.13	131	0.52	0.07	-0.24	0.04
	2609.76451								
[FeNOH] ⁸	-	1.72	1.38	2.20	116	0.55	-0.07	-0.22	0.00
_	2609.73346								

Energies (a.u.), distances (Å), partial atomic charges and spin densities (the latter shown in parentheses) for heme-histidine [FeNO]⁶, [FeNO]⁷, [FeNO]⁸, [FeNHO]⁸, and [FeNOH]⁸ adducts.

Data on the protonated models, [FeNHO]⁸ and [FeNOH]⁸, is particularly important, since a protonated [FeNO]⁸ species has indeed been isolated at the histidine-ligated b heme of myoglobin.[25] Table 2 confirms that, as previously proposed,[25] the proton must lie on the nitrogen rather than on the oxygen atom. Second, examining partial atomic charges and bond lengths in [FeNHO]⁸, we find all parameters to be consistent with previously described thiolate-ligated heme [FeNHO]^{8 [16]} Based on knowledge on the reactivity of this latter thiolate system,[16] the myoglobin [FeNHO]⁸ adduct should also react with a second NO molecule, thereby generating N_2O and H_2O . Indeed, evidence for such a reaction in myoglobin was available experimentally and has to this date not been rationalized.[25] This N₂O production by myoglobin appears however to be slow and incomplete: this deficiency is likely due to three factors. Firstly, myoglobin lacks good proton donors at the active site; two protons are in fact needed to convert [FeNHO]⁸ + NO into N₂O and H₂O. Secondly, the myoglobin heme site is relatively small and designed to bind one, rather than two diatomics.[12] Thirdly, myoglobin is not designed to receive electrons from biological electron-transfer agents in a rapid and efficient manner.[12] By contrast, the P450nor active site is designed to allow unrestricted contact between the NADH hydride donor to the Fe-NO adduct, and is also large enough to accommodate entry of a second NO molecule - as required for N₂O formation.[16]

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CcNIR, cd1NIR and respiratory cytochrome oxidases feature additional heme groups, which are located very close to the catalytic active site and deliver electrons immediately after initial binding of the diatomic substrate (NO or O_2).[16, 17]

The energetics of the two processes examined here, dioxygen reduction by cd1NIR and nitric oxide reduction by myoglobin, correlate well with previous findings on other hemoproteins, and allow us to better elaborate on our newly proposed "thiolate obstruction" theory. This theory, orthogonal if not opposed to the classical "thiolate push" dogma, argues that efficient reduction of diatomics such as dioxygen and nitric oxide is more readily accomplished by histidine-ligated hemes than by thiolate– ligated hemes. Generalizing, for reductive processes involving a small diatomic as the sole substrate, neutral ligands (e.g., histidine in cytochrome oxidases and heme oxygenase, lysine in cytochrome c nitrite reductase) are found to always be preferable over anionic ligands. By contrast, in enzymes designed to deal with more than one substrate, anionic ligands (e.g., the cysteine or tyrosine in heme-containing monooxygenases) are preferable, since they allow the safety switches needed to avoid uncoupling in their race against entropy.

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Dedicated to professor Gh. Marcu at his 80th anniversary

CATALYTIC SYNTHESIS OF DIMETHYL CARBONATE FROM METHANOL AND CARBON DIOXIDE

JENŐ BÓDIS¹ AND GEORGE A. OLAH²

ABSTRACT. The direct synthesis of dimethyl carbonate (DMC) from carbon dioxide and methanol, or dimethyl ether, in presence of magnesium and aluminum methoxide as catalysts was studied. The reactions were performed under batch conditions with catalysts in homogeneous liquid phase of excess methanol (no other solvent was used) at different partial pressures of carbon dioxide and at different temperatures. Depending on the reaction conditions and using magnesium methoxide as catalyst, dimethyl carbonate in 2-7% yield was detected.

Keywords: Dimethyl carbonate; Carbon dioxide activation; Homogeneous catalysis; Magnesium methoxide

INTRODUCTION

The versatile chemical property of dimethyl carbonate (DMC) makes it useful as a phosgene equivalent as well as a methylating and methoxycarbonylating agent [1-3]. On the other hand, replacing hazardous compounds is one of the many goals of environmental chemistry [3].

DMC can also be used as an additive in gasoline to improve fuel combustion and reduce automotive emissions [4]. Formaldehyde emissions were also lower with DMC as octane enhancer than with MTBE.

The industrial synthesis of DMC was based on phosgene [3].

 $CICOCI + 2 CH_3OH \rightarrow (CH_3O)_2CO + 2 HCI$

In the Enichem synthesis the CuCl/CuCl₂ catalyzed oxidative carbonylation reaction of methanol to DMC is used [1,3,4].

 $CH_{3}OH \ + \ 1/2O_{2} \ + \ CO \ \rightarrow \ (CH_{3}O)_{2}CO \ + \ H_{2}O$

Despite the fact that the Enichem process looks very feasible, the reaction is still subject of intensive studies. Kricsfalussy *et. al.* [5] tested molten CuCl-KCl salts as catalysts, Cavinato *et. al.* [6] Pd complexes, and Delledonne *et. al.* [7] Schiff base Co

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complexes. A Japanese patent [8] reports a complex catalytic system (PdBr₂-CuBr-KBr) in the presence of quinones and KOOCCH₃, with modest yields at atmospheric pressure. Relatively good space-time yields were obtained by gas phase oxidative carbonylation of methanol over (PdCl₂-CuCl₂-CH₃COOK)/activated carbon catalytic system [9]. Transition metal (Cu, Pd, Co, Rh) based catalytic systems [10] have been found to catalyze the electrochemical oxidative carbonylation of methanol.

UBE Chemical Industries [1,3,4,11] developed a pilot plant process for DMC production that uses the carbonylation of methyl nitrite over metal halides (Pd+Fe, Cu, Co, Ni or Sn) supported on active carbon.

Texaco Chemical [12-14] realized the coproduction of DMC and ethylene glycol through the transesterification of ethylene carbonate with methanol.



Titanium silicate molecular sieve (TS-1), exchanged with an aqueous solution of K_2CO_3 is described to be an excellent catalyst for the above transesterification reaction [15], reaching 57% yield (based on ethylene carbonate) of DMC. Some reports [16-18] claimed that by carrying out the transesterification of alkylene carbonates with methanol in presence of carbon dioxide, part of the methanol was reacting directly with CO_2 to DMC. However, the selectivity was very low and involved significant problems during separation.

The present work focuses on direct synthesis of DMC from carbon dioxide, an environmentally friendly building block, and methanol or dimethyl ether, cheap and available feedstocks of the chemical industry. Both reactions are thermodynamically unfavorable having positive ideal gas phase free energy changes (Δ G) at 100 °C.

 $\begin{array}{rcl} 2 \ CH_3OH &+ \ CO_2 &\rightarrow & (CH_3O)_2CO &+ & H_2O & \Delta G_{100^\circ C} = +12.2 \ kcal/mol \\ CH_3OCH_3 &+ & CO_2 &\rightarrow & (CH_3O)_2CO & \Delta G_{100^\circ C} = +15.1 \ kcal/mol \end{array}$

There are only few publications dealing with the reaction of carbon dioxide and methanol [19-29], and no literature report was found on the dimethyl ether reaction route. In almost all cases, the reported DMC yields were based on the amount of catalyst used. Using Bu₂Sn(OCH₃)₂ as catalyst [19], a 50% DMC yield was reached at 100 °C and 5 bar CO₂ pressure after 24 hours reaction time. In the presence of thallium compounds (ethoxide, or oxide) [20], only low reaction performances were detected (e.g. 20 cm³ CH₃OH resulted only 67-68 mg DMC, corresponding to 0.3% DMC yield based on used methanol). Bu₂Sn(OBu)₂-NaBr, Sn(OBu)₄-NaBr, and Ti(OBu)₄-NaBr systems are described as good catalysts, especially in the presence of dicyclohexylcarbodiimide as water scavenger [21-23] reaching 0.5- 2.68 % DMC yields based on methanol after 20 hours reaction time at 176 150 °C and 25-60 bar. Basic catalysts (K_2CO_3 , Na_2CO_3 , Cs_2CO_3 , K_3PO_4 , and (CH₃)₄NOH) applied together with CH₃I as promoter [24] gave high DMC yield in reactions carried out at 100 °C and 50 bar CO₂ pressure for 2 hours. Part of the CH₃I promoter is also consumed, making it difficult to evaluate the DMC yield based on the amount of methanol introduced into the reaction mixture. Nevertheless, in the case of K_2CO_3 , DMC was obtained even in the absence of CO₂ as reactant.

Without any experimental data regarding the amount of methanol, carbon dioxide, or catalyst used, a short publication [25] claimed 30 % conversion and 99% selectivity after 12 hours reaction time by using *in situ* prepared magnesium methoxide at 180 °C and at ~15 bar CO_2 pressure.

Results on the use of zirconia (ZrO_2) and zirconia promoted with phosphoric acid as catalysts for direct synthesis of DMC from methanol and CO_2 were also reported [26,27]. The maximum DMC yield (based on methanol) was found to be ~ 0.6% in case of zirconia and around 0.9% in case of promoted ZrO_2 catalysts.

The aim of our present contribution was to find effective catalysts or catalytic systems, and to establish the optimum reaction parameters for the direct conversion of carbon dioxide and methanol to DMC, a high scale required product both as reactant for chemical synthesis, and as gasoline octane enhancing additive.

EXPERIMENTAL

The carboxylation of methanol with carbon dioxide to dimethyl carbonate was carried out in a Parr autoclave (120 cm³) operated at 10-150 bar and 25 - 180 °C. Magnetic stirring at a speed of 1100 rpm was used. Weighed amount of catalysts were introduced into the autoclave. After addition of the dry methanol (usually 0.5 - 1.0 mol, Mallinkrodt, AR), the autoclave was sealed, and flushed with nitrogen or helium to remove atmospheric oxygen. The inside temperature was measured by a thermocouple. Carbon dioxide (Air Gas, research grade) was introduced in two different ways:

1.- at room temperature until \sim 10-55 bar total pressure and the autoclave was then placed in a thermostat bath preset to the reaction temperature.

2. - by adding carbon dioxide only after the catalyst-methanol mixture reached the desired reaction temperature. In this case the reactions had a well-defined starting time.

In the case of *in-situ* prepared methoxide catalysts, both versions were applied, but most experiments were run by introducing carbon dioxide only after the methoxide was prior formed at a higher temperature (1-2 hours at 200-210 °C for $Mg(OCH_3)_2$ and 220-230 °C for $Al(OCH_3)_3$). The progress of reaction in time was evaluated from the carbon dioxide consumption (added from a well-known volume reservoir keeping the working pressure in the autoclave constant). After the reaction was terminated, the autoclave was cooled to 0 °C in a water-ice bath, and the carbon dioxide vented carefully before opening the reaction vessel.

The reactants and products were analyzed using a Hewlett Packard 5971 mass spectrometer and Varian 3300 gas chromatograph equipped with a DB 5, or HP 1 capillary columns and FID detector. When the product mixture appeared as a suspension, filtering was employed before analysis. In most cases of metal methoxide containing mixtures, distillation was applied for the separation of volatile liquid compounds (mostly DMC and methanol) from dissolved catalyst. The water content of the reaction mixture (H₂O formation is supporting the catalytic pathway of the reaction

between methanol and CO_2 in presence of methoxide) was analyzed by a gas chromatograph equipped with TCD detector.

RESULTS AND DISCUSSION

The reaction of dimethyl ether with carbon dioxide was checked in different experiments realized at 20 - 180 °C and 20 - 150 bar with catalysts such as MgO in DMF, Pd(PPh₃)₄ in toluene, $Cr(CO)_6$ in pyridine, and magnesium methoxide in methanol. None of these experiments provided positive results. In order to achieve the DMC synthesis from methanol and carbon dioxide, a great number of attempts were carried out at temperatures and pressures mentioned above by testing different catalysts (Et₃N, PPh₃, MgO, Ru₃(CO)₁₂, Pd(PPh₃)₄, CuPhtc, 5% Rh/C, 10% Pd/C, 5% Pd/BaCO₃) and catalytic systems (5% Rh/C + CuCl + KCl, 5% Rh/C + PPh₃ + CH₃I, 5% Rh/C + Et₃N, 5% Rh/C + MgO, 5% Rh/C + Et₃N + CuCl, 5% Pd/BaCO₃ + Et₃N, 10% Pd/C + Et₃N). However, the catalysts tested were not able to produce measurable amount of DMC. Tin compounds (SnO₂, (CH₃)₃SnCl) were also unsuccessful, as well.

The experimental conditions of our first successful experiments and the results obtained for the conversion of methanol to DMC upon treatment with carbon dioxide are compiled in Table 1. The magnesium methoxide catalysts were prepared *in-situ* from Mg (turnings, Mallinckrodt, AR) and absolute methanol under the depicted reaction conditions, carbon dioxide being present from the beginning. Analyses were done without any preliminary separation by injecting samples in GC, and MS respectively.

Table 1

Reaction conditions and results for the synthesis of DMC from methanol (or dimethyl ether) and carbon dioxide in the presence of magnesium methoxide catalyst prepared *in situ* from Mg and methanol at the reaction conditions.

n _{MeOH} (mol)	n _{MeOMe} (mol)	n _{Mg} (mol)	p _{CO2} (bar)	T _{react} (°C)	p _{react} (bar)	t _{react} (h)	Y _{DMC} /MeO H (%)	Y _{DMC} /cat (%)
0.50	0.00	0.04	30	170	80	3	1.9	9.7
0.50	0.00	0.04	30	180	130	4	3.2	16.0
0.10	0.10	0.03	35	180	150	5	0.0	0.0

Y_{DMC}/MeOH: yield, molar percentage of methanol converted to DMC.

Y_{DMC}/cat: yield, molar percentage of DMC reported to the amount of catalyst.

As is shown in Table 1, only very low yields of DMC could be detected. By adding dimethyl ether, even the transformation of methanol was blocked.

In the next series of experiments (see Table 2), the magnesium methoxide was prepared in advance by stirring metallic magnesium with methanol at 200 °C for at least 1 hour, cooling the mixture at the desired reaction temperature, than adding carbon dioxide and running the reaction at constant temperature. Analyses still were done on crude reaction mixtures.

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Table 2

Reaction conditions and results for the synthesis of DMC from methanol and carbon dioxide (added only at the desired reaction temperature) in the presence of magnesium methoxide catalyst prepared previously *in-situ* from Mg and methanol at 200 °C.

n _{MeOH}	n _{Mg}	t _{react}	Y _{DMC} /MeOH	Y _{DMC} /cat
(mol)	(mol)	(h)	(%)	(%)
0.50	0.05	5	1.9	7.5
0.50	0.04	12	4.3	22.6
0.50	0.04	12	4.6	24.0

 p_{CO2} = 25 bar; T_{react} = 177 °C; p_{react} = 55 bar

Y_{DMC}/MeOH: yield, molar percentage of methanol converted to DMC.

Y_{DMC}/cat: yield, molar percentage of DMC reported to the amount of catalyst.

As shown in Table 2, the preformed catalyst led to a small increase of DMC yields, about 1%. The increased amount of magnesium didn't have a benefic effect on the reaction as part of the magnesium was found untransformed in the crude product mixture. Of course, the lower reaction time can be also a reason for the lower yield, as well.

In order to eliminate the analyses difficulties (needle blocking, reactions induced by the methoxide in the GC injector), the resulted reaction mixtures of the next experiments were separated by distillation before analysis. The yields included in Table 3 are dependent on reaction temperatures and pressures.

Table 3

Reaction conditions and results for the synthesis of DMC from methanol and carbon dioxide (12 hrs reaction time) catalyzed by magnesium methoxide preformed *in-situ* from Mg and methanol at 200 C. Carbon dioxide (p_{CO2} = 50 bar) was added to the CH₃OH-Mg(OCH₃)₂ mixtures at 50 °C.

Entry	n _{MeOH}	n _{Mg}	T _{react}	p _{react}	Y _{DMC} /MeOH	Y _{DMC} /cat
	(mol)	(mol)	(°C)	(bar)	(%)	(%)
1	0.50	0.04	50	50	0.0	0.0
2	0.50	0.04	100	67	0.0	0.0
3	0.50	0.04	150	85	2.7	14.4
4	0.50	0.04	160	92	5.0	26.2
5	0.50	0.04	170	98	6.7	35.3
6	0.50	0.04	180	106	7.2	37.6
7*	0.50	0.04	180	140	5.5	28.2
8	1.00	0.08	170	103	6.8	35.8
9	0.50	0.02	170	103	3.7	42.9

Y_{DMC}/MeOH: yield, molar percentage of methanol converted to DMC.

 Y_{DMC} /cat: yield, molar percentage of DMC reported to the amount of catalyst. *added 50 bar CO₂ at 25 °C.

At low temperatures (50-100 °C), the reaction practically did not work. Increase in temperature had a positive effect on the DMC yield, the maximum yield being detected

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at 180 °C. However, the variation of yields had a higher slope in the range of 150 - 170 °C than in the temperature range of 170 - 180 °C. The problem encountered with these experiments was, that by going up with temperature, the partial pressures of methanol and especially CO₂, are rising up. Therefore, the positive effect on the DMC yield must be attributed to both parameters, temperature and pressure together. The diminished yields obtained at high partial pressures of carbon dioxide (140 bar total pressure, Entry 7), indicates that at a certain value of the CO₂ pressure, the catalyst can be carbonated, losing its activity. A decreased amount of catalyst is leading to lower DMC yield, but to a higher catalyst related yield (Entry 9).

The content of water was measured by GC-MS analysis in the product mixture (Entry 6) and it was found to be 056 mol/l water, lower than the concentration of DMC (0.88 mol/l). It appeared to us that part of water reacts with magnesium methoxide forming CH_3OH and $Mg(OH)_2$ and deactivating the catalyst. Nevertheless, the presence of water in the product mixture is supporting our initial assumption that the reaction should be catalytic and DMC was mainly formed by insertion of CO_2 in magnesium methoxide. The resulting adduct reacts with methanol leading DMC, water and the reformed magnesium methoxide (Scheme 1).



Scheme 1. Catalytic pathways during the formation of DMC from carbon dioxide and methanol in presence of magnesium methoxide.



Scheme 2. Possible non-catalytic route for the formation of DMC through nonreversible reaction of magnesium methoxide with excess carbon dioxide.

However, at higher CO_2 pressures and higher temperatures (Entry 7), can be supposed another reaction route involving the Mg(OCH₃)₂ catalyst and CO₂, yielding

DMC and $MgCO_3$ (Scheme 2). This reaction is also deactivating the catalyst and should be avoided during the DMC synthesis.

The upcoming experiments were done at different temperatures by using also preformed methoxide catalysts. CO_2 was introduced only when the methanolmagnesium methoxide mixture reached the desired reaction temperature. The reaction conditions and the obtained yields are given in Table 4.

Table 4

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Reaction conditions and results for the synthesis of DMC from methanol and carbon dioxide (12 hrs reaction time) catalyzed by magnesium methoxide preformed *in-situ* from Mg or Al and methanol at

200 °C and 220 °C respectively. Carbon dioxide (p_{CO2} = 20-25 bar) was added to the CH₃OH-methoxide mixtures only when the desired reaction temperature has been reached.

		•			•	
Entry	n _{MeOH}	n _M	T _{react}	p _{react}	Y _{DMC} /MeOH	Y _{DMC} /cat
	(mol)	(mol)	(°C)	(bar)	(%)	(%)
1	0.50	0.04	180	60	4.3	22.6
2	0.50	0.04	170	55	3.8	19.7
3	0.50	0.04	160	55	3.6	18.7
4	0.50	0.04	150	55	2.8	14.5
5*	0.50	0.04	170	55	0.4	1.7
6**	0.50	0.05	170	55	0.2	0.4

Y_{DMC}/MeOH: yield, molar percentage of methanol converted to DMC.

 Y_{DMC} /cat: yield, molar percentage of DMC reported to the amount of catalyst.

*4 Å molecular sieve (Mallinckrodt) as water scavenger.

** Al as catalyst precursor.

Aluminum methoxide has also shown some activity (Entry 6). Although seen as a promising option, was not further explored in this study.

The use of a molecular sieve as water scavenger (Entry 5) had a negative effect on the reaction. Our finding is different from the almost neutral influence found for a 3Å molecular sieve on DMC formation over ZrO_2 catalyst [26], or from the positive effect of a 4Å molecular sieve revealed in case of DMC synthesis from methanol, carbon dioxide and methyl iodide mixture in presence of K_2CO_3 as catalyst [29].

At almost constant carbon dioxide partial pressures, an increase of temperature slowly increased the yields in the temperature range of 150-180 °C (see Figure 1).

A few experiments of DMC synthesis were also realized by operating the batch pressure reactor in a continuous mode. Thus, with the help of a dip tube, a constant CO_2 flow was passed through the methoxide-methanol mixture keeping the pressure at 40–50 bar inside the autoclave, and continuously venting CO_2 , DMC, CH_3OH , and water. Having the amount of distillate/time and the DMC content of the flux, we were able to evaluate the rate of DMC formation (turnover frequency, TOF = 0.08 - 0.1 mol_{DMC}/mol_{Mq}.h).



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Figure 1. Effect of reaction temperature on the DMC yields under the conditions given in Table 4.

CONCLUSIONS

Magnesium methoxide looks to be a promising catalyst for the direct synthesis of dimethyl carbonate from carbon dioxide and methanol. Since the reaction can be performed in excess of methanol, there is no need for another organic solvent as reaction medium.

The catalyst is deactivated by water (a green byproduct of the reaction) and by high carbon dioxide partial pressures. Continuous operation of the process in both, homogeneous or heterogeneous reaction at optimum pressures and temperatures will be able to diminish this inconvenience in the near future.

Dimethyl ether could not be converted to dimethyl carbonate through direct reaction with carbon dioxide under the conditions investigated in this study.

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REMOVAL OF SOME HEAVY METAL IONS FROM SYNTHETIC WASTEWATERS USING NATURAL ZEOLITES. A COMPARATIVE STUDY

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ABSTRACT. This paper presents some results obtained for removal of lead (II), zinc(II) and iron (II) ions from wastewaters by ionic exchange on natural zeolites from Transilvania. We used zeolitic (clinoptilolitic) volcanic tuffs from Măcicaş (M), Pâglişa (P) and Cepari Vultureni (CV) areas (Cluj county). The experiments were conducted in a batch reactor (static regime) and in a fixed bed column (dynamic regime). The efficiency of removal is high for all the natural zeolites and ions considered. A pretreatment of the zeolite with Na⁺ solutions increased the efficiency of the ionic exchange process.

KEY WORDS: Natural zeolites, clinoptilolite, wastewaters, lead, zinc, iron, ionic exchange.

1. INTRODUCTION

It is well known that 70% of the Earth's surface is covered by water, which follows a cycle driven by the absorption of energy from the Sun (evaporation-condensation-precipitation-infiltration to ground water). During this cycle, in which human activities play an important role, water pollution occurs. Heavy metal ions pollution is considered to be of high risk, due to the toxic effect upon the living organisms. Cadmium, chromium, cobalt, copper, lead, mercury, nickel and zinc are considered to have high toxicity, while iron and manganese are considered non toxic but with aesthetic effect. Copper and zinc ions are linked to some processes that take place in human body, but if their concentration is too large can become toxic.¹ Other heavy metal ions are highly toxic al low concentrations and can accumulate in the human body leading to different diseases.

Heavy metal ions in water come from natural and anthropogenic sources. Natural sources include rock weathering, soil erosion or dissolution of water-soluble salts. Usually metal ions introduced in water by natural sources are in small quantities and they not have any adverse effects on humans or other living organisms. Anthropogenic sources include municipal wastewater treatment plants, mining, industry (chemical, metallurgic, leather and textile, petroleum refining) corrosion and electroplating. If these wastewaters are not properly treated, heavy metal ions can be transported as

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dissolved species, can be adsorbed on sediments, volatilize in atmosphere and finally can be taken up by organisms causing deleterious effects.^{1,2}

Pollutants removal from wastewaters and water disposal in nature or recycling in the economic circuit is one of the main strategies applied to avoid environmental pollution.

Heavy metal ions removal from wastewaters can be realized using methods like: extraction, reduction, precipitation, electrochemical processes, adsorption or ionic exchange,²⁻⁶ each of them with advantages and disadvantages. The main disadvantage in most of the cases is the high cost of the process.

Adsorption and ionic exchange processes are widely applied to remove heavy metal ions from wastewaters due to their high efficiency even for small concentration of metal. In recent years many researchers oriented their efforts to improve existing materials⁷⁻¹² and processes (e.g. use of fluidised bed to remove copper on ionic exchange resin)^{13,14} or to find inexpensive materials that can be used as adsorbents or ionic exchangers. In the second category are included natural materials and natural wastes. There were investigated many natural zeolites and clays: Mexican clinoptilolite,¹⁵ Croatian clinoptilolite,¹⁶ American clinoptilolite,¹⁷ montmorillonite,¹⁸⁻²² scolecite (Brazilian natural zeolite),^{23,24} phillipsite (Jordanian zeolite),²⁵ vermiculite,²⁶ erionite, chabazite, mordenite,¹⁷ kaolinite,^{27,28} dolomite,²⁹ smectite and hectorite (magnesian smectite),³⁰⁻³² gibbsite,³³ sepiolite,³⁴ micas, illites³⁵⁻³⁷ and bentonites.³⁴⁻⁴² Other studies were oriented on the investigation of adsorptive properties of some natural materials and wastes such as apple residues,⁴³ grape stalk wastes,⁴⁴ crab shell particles,^{45,46} rice bran,⁴⁷ coffee grounds,⁴⁸ tree leaves,⁴⁹ ash particles derived from palm oil wastes,⁵⁰ peanut shell,⁵¹ sheep manure wastes^{52,53} or algae and microalgae.⁵⁴⁻⁵⁶

In this work we used zeolitic (clinoptilolitic) volcanic tuffs from Măcicaş, Pâglişa and Cepari Vultureni areas (Cluj county) to remove lead, zinc and iron from synthetic wastewaters. The deposits of volcanic tuffs from these areas are very important, from the environmental point of view, due to their clinoptilollite content, which is between 70 and 90% of rock mass. Volcanic tuff samples were collected from two stratigraphical horizons, those from the higher level are marked with "1" and they have a microporous structure, while those from the lower level are marked "2" and they have a mediumporous structure. Physico-chemical and mineralogical characterisation of all the zeolitic volcanic tuffs used in this study will make the object of another paper.

2. EXPERIMENTAL

Adsorptive and ionic exchange properties of natural zeolites are determined by the structure of their crystalline network, which determine the channel system and also by the negative charge excess due to $[AIO_4]^{5-}$ tetrahedrons compensated by mono- or divalent cations (eg. Na⁺, K⁺, Ca²⁺, Mg²⁺). These counter ions are mobile and can be total or partial exchanged during ionic exchange processes. Heavy metal ions removal was realised on zeolitic volcanic tuff modified by treatment with 1M HCI (Z-H form) and 1M NaCl + NaOH pH = 10 (Z-Na form). There were used two granulations, 0.63-1.0 mm and 0.2-0.4 mm of the zeolitic volcanic tuff. The stages involved in the zeolite preparation are as follows: crushing, grinding, size

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separation, washing with distilled water, drying at 105°C for 6 hours, treatment with 1M HCl in a stirring reactor with a zeolite acid solution ratio of 1:10, washing with distilled water to pH = 7 and finally drying at 105°C for 6 hours. At the end of this sequence we obtained the zeolite in -H form (Z-H). Also, during the treatment with HCl, zeolite channels are cleaned and pores opened. To bring the zeolite in -Na form which proved to be more efficient in the ionic exchange process,¹⁷ the zeolite is subjected subsequently to an alkaline treatment with strong Na⁺ solution (1M NaCl solution brought to pH = 10 with a 1M NaOH solution) according to equation (1). After the alkaline treatment the zeolite samples are washed again with distilled water to pH = 7 and dried for 6 hours at 105°C.

$$Z-H + NaCl = Z-Na + HCl$$
(1)

There were used monocomponent synthetic solutions containing Zn^{2+} , Pb^{2+} or Fe^{2+} ions, prepared from $ZnSO_4 \cdot 7H_2O$, $(CH_3OO)_2Pb \cdot 3H_2O$ and $FeSO_4 \cdot 7H_2O$ respectively, all analytical purity reagents. Concentrations of the solutions used in this study were selected accordingly to their toxicity and the maximum admissible concentrations in surface and drinking water.^{57,58} Determination of heavy metal ions in solution was realised using a Jenway spectrophotometer. Iron was determined as total iron. Experiments were carried out without any modification of the pH for the synthetic solutions.

The ionic exchange process was realised in a batch reactor in static regime using 10 g zeolitic volcanic tuff and 100 ml metal ions solution (zeolite:solution = 1:10). Synthetic wastewater samples were taken every 24 hours until the equilibrium was reached. We also worked on a fixed bed column ($d_i = 15$ mm) containing 5 g zeolite, with a flow rate of 0,055 ml/s (for all experiments) when samples were collected every 100 ml until the zeolite was exhausted (dynamic regime). The ionic exchange reactions are as follows:

$$2Z-Na + Zn^{2+} = Z_2-Zn + 2Na^+$$
⁽²⁾

$$2Z-Na + Pb^{2+} = Z_2-Pb + 2Na^+$$
(3)

$$2Z-Na + Fe^{2+} = Z_2-Fe + 2Na^+$$
(4)

Zeolitic volcanic tuffs used in our study were labelled considering the stratigraphical horizon and their origin as follows: Măcicaş area – M1 and M2, Pâglişa area – P1 and P2 and Cepari Vultureni area – CV.

The ionic exchange efficiencies, η , in %, were calculated with equation (5), considering the final concentration as follows: (a) in static regime – after the equilibrium was reached and (b) in dynamic regime – after 100 ml of synthetic wastewater passed the zeolitic volcanic tuff column bed.

$$\eta = \frac{C_i - C_f}{C_i} \cdot 100 \tag{5}$$

where

 C_{i} is the initial concentration, in mg/L

 C_f is the final concentration, in mg/L.

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3. RESULTS AND DISCUSSIONS

3.1. Influence of the chemical treatment on the ionic exchange capacity in dynamic regime.

The influence of the chemical treatment over the ionic exchange capacity was studied on the P1 sample brought in P1-H and P1-Na forms. The experiments were carried out in a column filled with 5 g zeolite with granulation of 0.2-0.4 mm and a synthetic solution containing Zn^{2+} ions, concentration 136.64 mg/L. Our results indicated that the efficiency of the ionic exchange process is almost twice as large in case of the -Na form modified zeolite (figure 1) – 42.03% for P1-H and 76.13% for P1-Na. This result is in agreement with conclusions of some previous studies performed on clinoptilolitic type natural zeolites.^{17,59,60} In case of the treatment with NaCl solutions,^{17,60} experimental results indicate that natural zeolites improve significantly their ion exchange capacity. Also H⁺ radius is 2.08 Å, while Na⁺ radius is only 0.95 Å, which will increase diffusion rate through the zeolite channels. If a NaOH treatment was applied,⁵⁹ an increase of the crystallinity degree was observed (explained by the amorphous phase reduction), which will improve zeolite exchange properties.

3.2. Comparison between the ionic exchange efficiencies of the zeolitic volcanic tuffs in static regime.

We studied the ionic exchange for zinc (237.58 mg/L) and lead (5.38 mg/L) ions using M1-Na, M2-Na, P1-Na, P2-Na and CV-Na samples with a granulation of 0.63-1.0 mm. Variation of heavy metal ions concentration in time and the ionic exchange efficiency for all the zeolitic volcanic tuffs are presented in figures 2 and 3 for zinc and figures 4 and 5 for lead. The ionic exchange reactions took place for both ions and all zeolitic volcanic tuffs considered (Zn^{2+} and Pb²⁺ concentrations in water decrease), therefore the zeolites under study are considered to be active in the ionic exchange reactions. In case of zinc, the efficiency of the ionic exchange process was higher than 96% for all samples considered.

The equilibrium in case of lead ion was reached more rapidly by comparison to zinc ion, which indicates a favourable kinetic for Pb^{2+} ion. Experimental ionic exchange studies performed on natural zeolites containing as a main component clinoptilolite and multicomponent heavy metal ions solutions showed that Pb ion is in the top of the selectivity sequences (Pb>Cu>Cd>Zn>Cr>Co>Ni).⁶¹⁻⁶³ In case of lead, ionic exchange efficiencies were over 93% with the exception of M1-Na microporous zeolitic volcanic tuff sample, where the efficiency is only 71.18%. This result can be correlated with the Pb²⁺ ionic radius (1.2 Å), which we assume that is too large by comparison with the pore diameter (microporous sample). After a comparison of ion exchange efficiencies for al the zeolitic volcanic tuffs and each ion, we can conclude that for the CV-Na zeolitic sample were obtained the best results in both cases. Also P1-Na zeolitic sample, in case of lead, and M1-Na zeolitic sample, in case of zinc, showed similar efficiencies for the process under study.

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Figure 1. Ionic exchange efficiency for zinc removal on P1 zeolitic volcanic tuff subjected to acid (P1-H) and acid-alkaline treatment (P1-Na)

3.3. Comparison between the ionic exchange efficiencies of the zeolitic volcanic tuffs in dynamic regime.

In this part of the work we studied the ionic exchange process in dynamic regime, in a fixed bed column, for Zn^{2+} and Fe^{2+} on M1-Na, P1-Na and CV-Na zeolitic volcanic tuffs. We used each time a 5 g zeolitic sample with granulation of 0.2-0.4 mm and solutions with different concentrations. In case of zinc we studied the influence of concentration over the efficiency of the ionic exchange process using solutions containing 136.64 mg Zn^{2+}/L and 59 mg Zn^{2+}/L , while for iron we considered solutions with 22.93 mg Fe^{2+}/L and 10 mg Fe^{2+}/L . Preliminary tests performed with zeolitic volcanic tuff with granulation 0.63-1.0 mm and higher concentrations.





Figure 2. Comparison between variations of Zn²⁺ concentration (in the synthetic wastewater) in time for all zeolitic volcanic tuffs studied, in -Na form and static regime.

Figure 3. lonic exchange efficiency for zinc removal in static regime on all zeolitic volcanic tuffs studied.

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Figure 4. Comparison between variations of Pb²⁺ concentration (in the synthetic wastewater) in time for all zeolitic volcanic tuffs studied, in -Na form and static regime.

Figure 5. Ionic exchange efficiency for lead removal in static regime on all zeolitic volcanic tuffs studied.

For removal of zinc in dynamic regime, were considered M1-Na, P1-Na and CV-Na zeolitic volcanic tuffs, which proved to have higher ionic exchange efficiencies in the static regime study. Results are presented in figures 6 and 7. We found out that in this case, the zeolitic volcanic tuff from Pâglişa area (P1-Na) presents a higher ionic exchange capacity, at the beginning of the process. This might be due to the pores distribution in the P1 sample, which has a surface area smaller than M1 sample, therefore pores are more accessible for the ionic exchange process. In case of P1-Na sample, we also investigated the ionic exchange for a second solution, diluted approximately at half. The efficiency of the ionic exchange process was calculated to be 81.57%, while for the first solution the efficiency was 76.13% after first 100 ml of synthetic wastewater passed the column bed. Also, the wastewater volume that can be processed, before the zeolite exhaustion, increased from 400 ml for the concentrated solution to 1000 ml for the diluted solution.





Figure 6. Comparison between variations of Zn²⁺ concentration (in the synthetic wastewater) with the efluent volume for M1-Na, P1-Na and CV-Na zeolitic samples in dynamic regime.

Figure 7. Ionic exchange efficiency for zinc removal in dynamic regime on M1-Na, P1-Na and CV-Na zeolitic samples

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Figure 9. lonic exchange efficiency for iron removal in dynamic regime on M1-Na, P1-Na and CV-Na zeolitic samples

The same zeolitic volcanic tufts were used to study the removal of iron from synthetic wastewaters (initial solution 10 mg Fe^{2+}/L). Results are presented in figures 8 and 9. All zeolitic samples demonstrated high efficiencies for iron removal. Iron was completely removed after first 100 ml of synthetic wastewater passed the zeolitic bed for M1-Na and P1-Na zeolitic samples, while for CV-Na zeolitic sample an efficiency of 98.4% was calculated.

We also carried out tests on P1-Na zeolitic sample using a second, more concentrated solution (figure 10). We observed that for the concentrated solution the exhaustion of the zeolite sample took place after we processed 900 ml wastewater. In case of the diluted solution after 1200 ml, when we stopped the experiment, the residual concentration of iron in solution was only 3.43 ml Fe_{total}/L (the ionic exchange process takes place until the initial concentration of iron is reached after the solution passed the zeolitic volcanic tuff bed, as a consequence of the ionic exchanger exhaustion). During the experiment we observed that Fe(OH)₃ brown precipitate started to build up on top of the zeolite sample, leading to an increase of the pressure drop, which modifies the hydraulic regime across the column.



Figure 10. Comparison between variations of Fe_{total} concentration (in the synthetic wastewater) with the effluent volume for P1-Na zeolitic sample in dynamic regime for two concentrations of iron solutions.

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4. CONCLUSIONS

(1) All zeolite samples considered can be used in wastewaters treatment to remove lead, zinc and iron.

(2) Our results indicated that the efficiency of the ionic exchange process is almost twice as large in case of the -Na form modified zeolite by comparison with the -H form.

(3) In static regime the most efficient samples were CV-Na and M1-Na for zinc removal and CV-Na and P1-Na for lead removal.

(4) The equilibrium in case of lead ion (static regime) was reached more rapidly by comparison to zinc ion, which indicates a favourable kinetic for Pb^{2+} ion.

(5) In case of dynamic regime for zinc removal, the most efficient sample was P1-Na. Also the ionic exchange process was more efficient and column exhaustion took place after a larger volume of wastewater was processed when solution concentration is lower.

(6) The iron was removed completely after 100 ml wastewater passed on M1-Na and P1-Na samples and the life of the zeolite was extended when diluted solution was used.

(7) After the heavy metal ions removal from wastewaters, the zeolites samples in metal ion form (Z-Meⁿ⁺) can be regenerated and reused in the same process with metals recovery² and as catalysts in flue gas treatment to remove NO_x⁶⁴ or in Fenton type processes to remove organic pollutants from wastewater.⁶⁵

(8) Further studies will consider multicomponent synthetic solutions and wastewaters collected from industrial effluents.

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Dedicated to professor Gh. Marcu at his 80th anniversary

SPECTROSCOPIC AND MAGNETIC INVESTIGATIONS OF THE CHROMIUM(III) TUNGSTOARSENATE

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ABSTRACT. The sandwich-type tungstoarsenate complex Na₆[Cr₄(H₂O)₂)(AsW₉O₃₄)₂] . 18H₂O (**Cr^{III}AsW₉**) was synthesized and investigated by elemental analysis, thermogravimetry, FT-IR, UV-VIS, EPR spectroscopy and magnetic susceptibility methods. The results suggest a sandwich-type structure in which two of the [AsW₉O₃₄]⁹ trilacunary Keggin fragments are connected by a Cr₄O₁₆ group of four coplanar CrO₆ octahedra sharing edges.

The aim of this study consists in establishing the modifications that Cr^{III} ions make in the trivacant Keggin units, in determination of the local symmetry around the chromium ions and the identification of the nature of the chromium-chromium interactions.

Keywords: polyoxometalates, heteropolyoxometalates, chromium(III), tungstoarsenates

Introduction

Scientific interest in polyoxometalates is increasing worldwide, due to the enormous variety of structures of these compounds, often with unusual properties, rendering them useful candidates in medicinal applications and other fields such as nanomaterials, catalysis and magnetochemistry [1-4]. Although the mechanism of formation of polyoxometalates is commonly described as self-assembly, it is not well understood. In spite of this drawback, the synthesis of new compounds can be based on a rational synthetic approach and efficient control of the reaction path.

Lacunary polyoxometalates are usually synthesized from precursor ions by loss of one or more MO_6 octahedra [1, 5]. These lacunary polyoxometalates show an increased reactivity for metal ions and organometallic fragments, leading to the formation of a broad variety of complexes, in which the heteropolyanion framework remains unchanged. Taking into account the coordination requirements and the size of a given transition metal ion, the geometry of the reaction product can often be predicted.

Although polyoxometalate complexes of the trilacunary $[PW_9O_{34}]^9$ with metal cations and organometallic fragments have been extensively investigated, there are relatively few papers concerning $[AsW_9O_{34}]^9$ analogs [6-9].

Herein the synthesis and characterization of trilacunary $[AsW_9O_{34}]^9$ (**AsW**₉) and Na₆[Cr₄(H₂O)₂](AsW₉O₃₄)₂] 18H₂O (**CrAsW**₉) are reported. The complex was investigated using chemical and thermal analysis, UV-VIS, FT-IR and EPR spectra,

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as well as magnetic susceptibility measurements. The results suggest a sandwichtype structure similar to that reported by Knoth *et al.* [10], in which two of the AsW_9 units are connected by a belt of four chromium atoms (Fig. 1).



Fig. 1. Proposed structure of the Na₆[Cr₄(H₂O)₂](AsW₉O₃₄)₂]·18H₂O polyoxometalate complex.

Experimental

1. Methods

Tungsten, chromium, sodium and arsenicum were determined by atomic absorption. The water content was determined on the basis of thermal analysis performed using a METTLER-TGA/SDTA 851^eSTAR^eSoftware derivatograph. FT-IR spectra were recorded on a Jasco FT/IR 610 spectrophotometer in the 4000–400 cm⁻¹ range, using KBr pellets. Electronic spectra in the visible range were performed in aqueous solutions on an ATI Unicam-UV-Visible spectrophotometer with Vision Software V 3.20. EPR spectra on powdered solids have been recorded at room temperature and 80 K at *ca.* 9.6 GHz (X-band) using a Bruker ESP 380 spectrometer. The magnetic susceptibility measurements were performed using a Faraday type balance in the temperature range of 77–273 K.

2. Materials

Reagent grade chemicals were used and all syntheses and studies were carried out in distilled water.

The sodium salt of the polyoxometalate ligand, Na₈H[AsW₉O₃₄] (**AsW₉**) was prepared following the procedure described previously in the literature [11].

Synthesis of Na₆[Cr₄(OH₂)₂(AsW₉O₃₄)₂]·18H₂O CrCl₃·6H₂O (1.34 g, 3.094 mmol) was dissolved in water (10 ml) and added dropwise, while stirring, to a boiling aqueous solution (50 ml) of Na₈H[AsW₉O₃₄]·11H₂O (5.5 g, 2.070 mmol). The slightly turbid final solution was boiled for 2h, its pH adjusted between 4.5 and 5.0 with 6M HCl, and filtered under suction. The filtrate was heated at 80°C, while stirring, until half of the volume remained. It was then cooled to room temperature. A few days later a green precipitate appeared. It was recrystallized from hot water, at pH= 6.0-6.5 and dried under vacuum over P₂O₅. The yield was 2.3 g (60%). Anal. Calc. for H₄₀As₂Cr₄Na₆O₈₈W₁₈ (5776.95): As 2.80; Cr 4.00; Na 2.60; W 63.10. Found: As 2.85; Cr 3.96; Na 2.63; W 63.04. IR (polyoxometalate region, cm⁻¹): 955, 890, 839, 795, 734, 517, 472, 440.

SPECTROSCOPIC AND MAGNETIC INVESTIGATIONS OF THE CHROMIUM(III) TUNGSTOARSENATE

Results and Discussions

The reaction of Cr^{III} with AsW_9 in water can be described by equation. (1). Isolation of the products as sodium salts and recrystallization from hot water, resulted in relatively good yields of the $Cr^{III}AsW_9$ complexes.

$$4Cr^{3^{+}} + 2[AsW_{9}O_{34}]^{9^{-}} + 2H_{2}O \rightarrow [Cr_{4}(OH_{2})_{2}(AsW_{9}O_{34})_{2}]^{6^{-}}$$
(1)

Elemental analysis of the final products are consistent with the suggested formula for $Cr^{III}AsW_9$.

Thermal study behavior of the new chromium (III) tungstoarsenate reveals that complex contain 18 crystallization water molecules and two coordination water molecules. For **Cr^{III}AsW**₉, two endothermic processes were observed between 120 and 260 °C (Fig. 2). These processes were assigned to water loss. The TG curves showed that the last process, corresponding to coordination water molecule loss, was the slowest. The exothermal process, appearing around 480°C on the DTA curve of complex, was assigned to the decomposition of polyoxometalate [12]. On the basis of these thermogravimetric studies, it can be thus concluded that **Cr^{III}AsW**₉ are thermally stable up to 480°C.



Fig. 2. Thermogravimetric and thermodifferential curves of Cr^{III}AsW₉

Electronic Spectra

It have been registered the electronic UV and VIS spectra on aqueous solutions of the sodium salts of the polyoxometalates complex $Cr^{III}AsW_9$ and have been compared with this of the ligand AsW_9 .

The UV electronic spectra (Fig. 3) are characteristic to the polyoxometalates and similar to the ligand. [13,14] The higher energy band (v_1) at ~ 50000 cm⁻¹, due to the d π -p π proper transitions from the W=O_t (O_t -terminal oxygen atom) bonds, insignificantly shifted in the complex, compared to the ligand, which can been associated with the lack of involvement of the terminal oxigen atoms in the coordination of the cations of transitional metals.





Fig. 3. UV spectra of the sodium salts of $Cr^{III}AsW_9$ (solid) and AsW_9 (dotted) obtained in $5x10^{-5}$ mol·l⁻¹ aqueous solution.

The lower energy band (v₂) at ~ 39000 cm⁻¹, due to the d π -p π -d π electronic transitions from the tricentric W-O_{c,e}-W (O_{c,e}-bridging oxygens between two WO₆ octahedra sharing corner or edge respectively), bonds, with an expected absorbtion maximum in 40000-38000 cm⁻¹ range, in Cr^{III} complex spectrum, apearing as to shoulders in the spectrum, is due to the decrease of the symmetry as well as to the distorsion of the WO₆ octahedra through complexation, which influences the electronic transfer from these bonds. [15]

Information about the local environment of Cr(III) ions has been obtained by means of d-d transitions from the visible electronic spectrum performed in aqueous solution. Two bands of the chromium complexes, at 16770 cm⁻¹ and 21200 cm⁻¹ are assigned to ${}^{4}A_{2g}(F) \rightarrow {}^{4}T_{2g}(F)$ (v₁) and ${}^{4}A_{2g}(F) \rightarrow {}^{4}T_{2g}(F)$ (v₂) transitions. The third band (${}^{4}A_{2g}(F) \rightarrow {}^{4}T_{1g}(P)$) expected in the UV domain is obscured by the change-transfer and ligand-specific bands.

The first broad and unsplit band (v₁) yields the 10D_q parameter, while the second (v₂) is also broad and asymmetric. This can be explained by a small distortion from the ideal octahedral symmetry, because of the Jahn-Teller effect, spin-orbit coupling and the mixing of neighboring quartet-doublet states (⁴T_{1g} and ²T_{2q} or ²A_{1q}).

The position of the third band and the spectral parameters were calculated by the method proposed by Lever [16], using a value of 400 cm⁻¹ for the interelectronic repulsion constant. The obtained values are: $D_q/B = 4.2$, $v_3/B = 88.6$, $\beta = 0.388$, $v_3 = 35286$ cm⁻¹. The latter band (v_3), assigned to a forbidden transition of two electrons, should have low intensity [17].



Fig.4. The visible electronic spectrum of the Cr^{JII}AsW₃ complex in 5x10⁻³ mol·l⁻¹ aqueous solution. The Gaussian components are represented with dashed lines.

Vibrational Spectra

All antisymmetric frequencies bonds involving tungsten ions are shifted towards higher or lower frequencies (1–18 cm⁻¹) in the complex spectrum compared to the ligand spectrum (Table 1).

The relative small shift of the $v_{asym}W-O_t$ vibration band from 945 cm⁻¹ in the ligand spectrum to 950 cm⁻¹ in complex spectrum is due to the fact that the terminal O_t atoms at the lacunary surface are not involved in the coordination of Cr^{III} ions [18]. The vibration frequencies of the As- O_i (O_i -internal oxygen atom) bonds are not observed at ~820 cm⁻¹ overlaps that of W- O_b -W (O_b – bridging oxygen atom) bonds. [18] Two vibration bands of tricentric W- O_c -W bonds of the corner sharing WO₆ octahedra appear in spectrum of the ligand at 878 and 843 cm⁻¹, respectively, and in complex spectrum at 890 and 839 cm⁻¹, respectively, suggesting the nonequivalence of the W- O_c -W bonds when linking octahedra from the equatorial and polar regions of the trilacunary fragments [18].

Two vibration bands of tricentric $W-O_e-W$ bonds of the edge-sharing WO_6 octahedra appear in the ligand spectrum at 795 and 752 cm⁻¹, respectively, suggesting that two nonequivalent bonds of this type are present. The complex spectrum shows two bands at 796 and 734 cm⁻¹ too. [19]

FI-IR Data of the Cr^{-A} SVV ₉ complex and the ASVV ₉ ligand				
Vibration band	ν (cm ⁻¹)			
	AsW ₉	Cr [™] AsW ₉		
vasW-Ot	945	950		
vasAsOi+vasW-Oc-W	878	890		
vasW-Oc-W	843	839		
v _{as} W-O _e -W	795	795		
	752	734		

TABEL 1.

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Magnetic measurements

The temperature dependence of the reciprocal molar susceptibility $1/\chi_m$ for the $Cr^{III}-AsW_9$ complex in the 77–273 K temperature range is presented in Fig. 5.

The effective magnetic moment obtained from the magnetic susceptibility data in the 77 – 273 K temperature range is μ_{eff} = 10.66 μ_B . The value of Curie-Weiss temperature θ = -6.1 K, indicates the presence of antiferromagnetic coupling between the Cr^{III} ions.

These measurements, together with the informations obtained from EPR spectroscopy, leaded to the establishment of intra- and intermolecular metal-metal interactions .



Fig. 5. Variation of reciprocal molar susceptibility of the Cr^{III}AsW₉ complex with respect to temperature. Solid line represents the best fit of the experimental data with a Curie-Weiss behavior.



Fig. 6. EPR Spectrum of the Cr^{III}AsW₉

SPECTROSCOPIC AND MAGNETIC INVESTIGATIONS OF THE CHROMIUM(III) TUNGSTOARSENATE

EPR Spectrum

The polycrystalline EPR spectrum of the Cr complex obtained at 80 K contains a single large \approx 2495 G pseudo-isotropic signal centered at g=1.963. The line width of the signal indicates the presence of the coupling between the chromium ions; the almost isotropic shape of this is due to an O_h symmetry around the chromium (III) ions. The shape of this signal is not modified at 293 K. By raising the temperature, the line width of signal decreases, which indicates the presence of small Cr(III)-Cr(III) super-exchange interactions (Fig. 6). [20]

Conclusions

The results of the performed investigations revealed that the chromium complex of the AsW₉ trilacunary polyoxometalate was prepared. A sandwich-type structure, similar to the structure of the other complexes reported by Knoth *et. al.*, [10] (Fig. 1) is inferred from the experimental results. The coordination of the chromium (III) ions in the lacunary region of the Keggin units results from the shift of the $v_{asym}(W-O_{c,e}-W)$ stretching frequencies in the FT-IR spectrum of the complex compared to the ligand spectrum.

Every metal ion is surrounded by six oxygen atoms in a distorted O_h symmetry and has a $d_{x^2-y^2}$ orbital as ground state. The almost isotropic shape form of the signal

from EPR spectrum registered on the Cr^{III}AsW₉ is due to an O_h symmetry around the Cr^{III}.

The magnetic susceptibility measurements in the 77–273 K temperature range have indicated the presence of antiferromagnetically coupled Cr^{III} ions.

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Dedicated to professor Gh. Marcu at his 80th anniversary

THE INFLUENCE OF ADHESIVE ON COMPOSITE MATERIALS BONDED JOINTS ASSEMBLIES

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ABSTRACT. The paper presents a theoretical calculation model of plane assemblies joined with adhesive, based on an energy method. After the determination of the cinematically acceptable field of stresses, according to the applied load, a variational calculus on the expression of elastic potential energy leads to the complete expression of the stress field in the whole assembly. A first parametric analysis (geometrical and physical parameters) is carried out on a double-lap plane assembly and makes it possible to deduce the optimal length and thickness of the adhesive. For the assembly, the total force-displacement behavior is well defined. Thus the analytical model makes it possible to determine the rigidity of the assembly and to obtain a simple formulation very rapidly which gives the total behavior of the assembly.

Keywords: Stress Analysis, Bonded joints, Numerical modeling

INTRODUCTION

The increase in use of adhesive bonded joints is due to the many advantages of this method compared to the traditional methods. This assembling method distributes the stresses over the whole joining surface and removes the concentrations of stresses to the boundary of holes generated by bolting or riveting assemblies. The mechanical performance of an adhesive bonded joint is related to the distribution of the stresses in the adhesive layer. Consequently it is essential to know this distribution, which, because of its complexity, makes prediction of fractures difficult. From the first works of Volkersen (1938) when only a distribution of the shear stress in the adhesive joint was taken into account to the more recent studies by finite elements, many formulations have made it possible to define the field of stresses in such assemblies better and better.

Since the first work of Volkersen until the more recent studies by finite elements many formulations allowed to better define the stress field in such assemblies. Among those, we can mention in principal the works due to:

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 Goland and Reissner (1944) – dealing with the description of the peeling stress. They showed that the effects of flexion create additional stresses which are superimposed on shear stresses,

 Hart-Smith (1973) - studied the elastoplastic behavior of the joint of adhesive and took into account the effects of temperature,

• Renton and Vinson (1977) have checked certain results by experimental way.

One should mention also the theoretical work of Oljado and Eidinoff (1978), or those due to Bigwood (1989), and Allman (1977),

Following Goland and Reissner (1944), Volkersen (1965) introduced into his new analysis the normal stress "stress of shearing" (peeling stress) which is variable in the thickness of the adhesive layer. This assumption enabled him to build a stress field observing the boundary conditions of the assembly. However, due to the complexity and difficulty of its implementation, this analytical formulation is not easily applicable.

Gilibert and Rigolot (1979 – 1991) propose, based on the method of the asymptotic developments connected in the vicinity of the ends, an analytical formulation of the stress field over the entire covering length. If this formulation constitutes a clear improvement of the modeling of the field of the constraints on the level of the ends and represents experimental reality better, it is however not valid near the free edges.

Other more recent studies, due to Liyong (1994) present an analytical formulation making it possible to calculate the mechanical strength of the double-lap joints by taking into account the effects of temperature; however the author disregards the stress distribution and calculates only the joint shearing deformation energy.

Adams and Peppiatt (1974), in their finite element analysis, circumvented this difficulty by studying a joint modified by the addition of a regularizing part. However even this study is not satisfactory on the level of the ends.

Tsai and Oplinger (1998) develop the existing traditional solutions by the inclusion of shearing strains, neglected until there. The solutions obtained ensure a better forecast of the distribution and intensity of the shear stress.

Mortensen and Thomsen (2002) developed the approach for the analysis and the design of the joints adhesive bonded. They held into account the influence of the interface effects between the adherents and they modeled the adhesive layer by assimilating it to a spring.

Nemeş (2004) use a technique based on the minimization of the potential energy. The first stage consists in building a statically acceptable stress field, i.e. verifying the boundary conditions and the equilibrium equations. Then, the potential energy generated by such a stress field is calculated. In the third stage, the potential energy is minimized in order to determine the stress distributions. As we have just seen, the analytical formulations and the finite element analysis provide a stress field satisfying for the median part of the joints. On the other hand, these two approaches provide results that do not satisfy the boundary conditions imposed at the ends of covering. However it is in the vicinity of these ends that one observes the majority of the phenomena of degradation (non-linear behavior, damage, cracking, even fracture). The analytical study that follows gives a first solution of the field of the constraints respecting the whole of these conditions. ADHESIVE INFLUENCE ON COMPOSITE MATERIALS BONDED JOINTS ASSEMBLIES

RESULTS AND DISCUSSION THEORETICAL MODEL

All work has encountered difficulties in modeling the stress field in the vicinity of the ends of the joint. The method used to obtain the optimal field for this type of assembly consists of:

- Construction of a statically acceptable field,
- Calculation of the potential energy associated with the stress field,
- Minimization of this energy by a variational method,
- Resolution of the differential equation obtained.

Definitions and hypothesis. Let us consider a plane joining with double covering (Figure 1) whose supports are maintained stuck by a marked elastic adhesive of the index ©.

The whole joining is in balance under the action of a tensile load. The two adherents are subjected to the same load of F/2 traction following axis X, the median plate it being subjected to an opposite load of intensity F. Considering the geometrical symmetries of the problem, our analysis will be limited to the study of the higher half of the assembly represented Figure **2**.



Figure 1. Double-lap adhesive assembly



Figure 2. Geometrical and material definition of the double-lap joint

Where:

E_c, v_c, Young's modulus and Poisson's ratio of the adhesive ©,

• E_{tl} , E_{ll} , v_{tll} , longitudinal, transverse modulus and Poisson's ratio of the inner tube,

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E_{2t}, E_{2l}, v_{t12}, longitudinal, transverse modulus and Poisson's ratio of the external tube,

- e_c, adhesive © thickness,
- e₁, e₂, adherents ① and ② thickness,
- L, joining length,
- f and q, tensile stresses following x axis, on the adherents.

The constraints in various materials will be located by the index (i), where i = \mathbb{O} , \mathbb{O} or \mathbb{O} . We are in the case of plane constraints and we will adopt the following assumptions:

- the state of plane stresses: $\tau^{(i)}_{zx} = \tau^{(i)}_{zy} = \sigma^{(i)}_{zz} = 0$
- the $\sigma_{xx}^{(i)}$, $\tau_{xy}^{(i)}$ et $\sigma_{yy}^{(i)}$ stresses are independent of z variable
- the $\sigma_{\rm xx}^{(l)}$ et $\sigma_{\rm xx}^{(2)}$ stresses are function only of x variable
- the normal stress in the adhesive will be considered null: $\sigma_{\rm vv}^{(c)}=0$

The stress field is thus reduced to:

- adherent Φ : $\sigma_{xx}^{(1)}(x)$, $\tau_{xy}^{(1)}(x,y)$, $\sigma_{yy}^{(1)}(x,y)$,
- adhesive ©: $\tau_{xy}^{(c)}(x)$, $\sigma_{yy}^{(c)}(x,y)$,
- adherent **Q**: $\sigma_{xx}^{(2)}(x)$, $\tau_{xy}^{(2)}(x,y)$, $\sigma_{w}^{(2)}(x,y)$.

The stress field definition. We build a statically acceptable stress field by respecting the preceding assumptions. i.e. checking the local equilibrium equations, the boundary conditions as well as the conditions of continuity to the interfaces with the adhesive. We write the balance of the forces which act on the whole of joining by carrying out a fictitious cut of the assembly following y axis.

The stress components for each three component must satisfy the equilibrium equations, the conditions of continuity of the vector forced with the crossing of the interfaces as well as the boundary conditions in x = 0, X = L, y = 0 and $y = (e_1 + e_2 + e_c)$. Starting from equilibrium equations, in order to determine the various component the of the stress vectors, we must write the boundary conditions as well as the conditions of continuity of the stress vectors with the interfaces.

On the free edges, the normal stress on the surface and shear stresses are null (equal with zero value). With the crossing of the interfaces, these same stresses must be continuous. The nullity of the shear stress for y = 0 is due to the symmetry of the problem. All these conditions were gathered below:

- For x = 0: $\sigma_{xx}^{(1)} = q$, $\sigma_{xx}^{(2)} = 0$, $\tau_{xy}^{(1)} = 0$, $\tau_{xy}^{(2)} = 0$
- For x = L: $\sigma_{xx}^{(1)} = 0$, $\sigma_{xx}^{(2)} = f$, $\tau_{xy}^{(1)} = 0$, $\tau_{xy}^{(2)} = 0$
- For y = 0: $\tau_{xy}^{(2)} = 0$
- For $y = e_2$: $\tau_{xy}^{(2)} = \tau_{xy}^{(c)}$, $\sigma_{yy}^{(2)} = \sigma_{yy}^{(c)}$
- For $y = e_2 + e_c$: $\tau_{xy}^{(1)} = \tau_{xy}^{(c)}$, $\sigma_{yy}^{(1)} = \sigma_{yy}^{(c)}$
- For $y = e_{_1} + e_{_c} + e_{_2} \colon \, \tau^{(1)}_{_{xy}} = 0$, $\, \sigma^{(1)}_{_{yy}} = 0$

The stress field is thus reduced to the following components:

$$\begin{aligned} \tau_{xy}^{(1)}(x,y) &= \left[\left(e_{1} + e_{2} + e_{c} \right) - y \right] \frac{d\sigma_{xx}^{(1)}}{dx}, \\ \sigma_{yy}^{(1)}(x,y) &= \frac{1}{2} \left[y - \left(e_{1} + e_{2} + e_{c} \right) \right]^{2} \frac{d^{2}\sigma_{xx}^{(1)}}{dx^{2}} \\ \tau_{xy}^{(c)}(x) &= e_{1} \frac{d\sigma_{xx}^{(1)}}{dx}, \\ \sigma_{yy}^{(c)}(x,y) &= e_{1} \left[\left(\frac{e_{1}}{2} + e_{2} + e_{c} \right) - y \right] \frac{d^{2}\sigma_{xx}^{(1)}}{dx^{2}}, \end{aligned}$$
(1)
$$\sigma_{xx}^{(2)}(x,y) = f - \frac{e_{1}}{e_{2}} \sigma_{xx}^{(1)} \\ \tau_{xy}^{(2)}(x,y) &= \frac{e_{1}}{e_{2}} y \frac{d\sigma_{xx}^{(1)}}{dx}, \\ \sigma_{yy}^{(2)}(x,y) &= \frac{e_{1}}{2} \left[\left(e_{1} + e_{2} + 2e_{c} \right) - \frac{y^{2}}{e_{2}} \right] \frac{d^{2}\sigma_{xx}^{(1)}}{dx^{2}} \end{aligned}$$

Deformation energy calculation. In the defined stress field, the only unknown factor is the expression of the normal stress $\sigma_{xx}^{(1)}$, expression which we will determine using the principle of minimum of complementary energy. The potential energy associated with the statically acceptable field previously given is written for a joining length 1 and width unit on axis z:

After integration following y, for the potential energy was obtained the following form:

$$\xi_{\rm P} = \int_{0}^{L} \left[A\sigma_{\rm xx}^{(1)^2} + B\sigma_{\rm xx}^{(1)} \frac{d^2 \sigma_{\rm xx}^{(1)}}{dx^2} + C \left(\frac{d\sigma_{\rm xx}^{(1)}}{dx} \right)^2 + D\sigma_{\rm xx}^{(1)} + E \left(\frac{d^2 \sigma_{\rm xx}^{(1)}}{dx^2} \right)^2 + F \frac{d^2 \sigma_{\rm xx}^{(1)}}{dx^2} + K \right] dx \quad (2)$$

the constants A, B, C, D, E, F and K depend on the geometrical and material characteristics of the three components as well as the applied loading. The expressions of these various constants according to the geometrical and physical characteristics of the assembly are given by the following equations.

By applying to the functional ξ_p a variational calculus and by using the boundary conditions which one also writes in the form:

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$$\sigma_{xx}^{(1)}(x=0) = q = \frac{e_2}{e_1} f \cdot \frac{d\sigma_{xx}^{(1)}}{dx}(x=0) = 0 \cdot \sigma_{xx}^{(1)}(x=L) = 0 \cdot \frac{d\sigma_{xx}^{(1)}}{dx}(x=L) = 0$$
(3)

we obtain that the energy is minimal when the stress function $\sigma_{xx}^{(l)}(x)$ is solution of the following differential equation:

$$E\frac{d^{4}\sigma_{xx}^{(1)}(x)}{dx^{4}} + (B-C)\frac{d^{2}\sigma_{xx}^{(1)}(x)}{dx^{2}} + A\sigma_{xx}^{(1)}(x) + \frac{D}{2} = 0$$
(4)

Parametric study of the adhesive bonded joints

Stress distribution. Figure 3 shows the stress distributions in the adhesive for the analyzed configurations. These data show that the peeling stresses are more important than shear stresses. This observation joined the remarks of Volkersen (1965), Gilibert and Rigolot (1985) who also argument that the peeling stress are most important. We notice that: for σ_{yy} , the maximum values are obtained on the free edges (z = 0, z = L). These values are much localized at the edges; however, the maximum constraint σ_{yymax} is obtained in compression, for τ_{xy} , we raises two peaks of stresses located at equal distance of the two free edges. The peaks do not have the same intensity because of the difference of rigidities of the two stuck adherents.



Figure 3. Distribution of the stresses in the adhesive of a VE \pm 45°-AV 119-VE \pm 45° assembly for F = 1 N/mm. a) the peeling stress (σ_{vv}), b) the shear stress (τ_{vv}).

Based on previous analysis of distributions we can note that the peeling stress are more important than the shear stresses thus the use of a criterion of rupture of the adhesive bonded joint must take into account not only the stress shear τ_{xy} but also the peeling stress σ_{yy} .

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Parametric study

Covering length influence. Figure 4 shows the influence of covering length on the distribution and the intensity of shear stresses. The fact of increasing the length of joining beyond a certain value does not have any influence on the maximum stresses in the adhesive. Indeed, for the assembly with double covering there is an optimal length of covering beyond which the added length is not under load. By increasing gradually the covering length we observed:

- the reduction of the values of the shear stress in the middle of the joint,
- the displacement of the stress peaks towards the free edges.



Figure 4. Variation of τ_{xy} in the adhesive according to the covering length (L = 10÷100 mm) and F = 1 N/mm, for a VE ±45°-AV 119-VE ±45° assembly.

Influence of rigidities. Figure 5 represents the influence of the elastic module of the adhesive on the shear stress in the adhesive. The maximum peaks will increase slightly when the elastic module grows.

The influence of relative rigidity, between the two stuck substrates, is illustrated in Figure 6.

We can notice that the maximum peaks on the two edges are not equaled any more if ratio E_2/E_1 is different from 1.

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Figure 5. Variation of the shear stress $(-\tau_{xy})$ in the adhesive according to the elastic module of the adhesive.



Figure 6. Variation of the shear stress $(-\tau_{xy})$ according to relative rigidity.

Influence of adhesive thickness. Figures 7 - 8 show the influence of adhesive thickness. When the adhesive thickness increases the maximum stresses in the adhesive decrease and the distribution (shear and peeling stress) tends to be uniform over the entire covering length, except in the vicinity of the free edges.



Figure 7. Variation of the shear stress $(-\tau_{xy})$ according to the adhesive thickness.

At the same time, for the peeling stresses, we can also note a considerable reduction in the maximum values to the level of the free edges.



Figure 8. Variation of the peeling stress (σ_{yy}) according to the adhesive thickness.

CONCLUSIONS

The present data point the followings:

- the maximum values of σ_{yy} are obtained on the free edges and the maximum values are localized at the edges,

- concerning $\tau_{\text{xy}},$ we found two peaks of stresses located at equal distance of the two free edges,

the peeling stresses are more important than shear stresses,

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 there is an optimal length beyond which the maximum stress do not evolve any more,

• the intensities of the peaks are influenced by the rigidities difference of the two adherents, the maximum peaks increase slightly when the elastic module grows,

 the shear stress in the adhesive increases with the increase in the relative rigidity of the adherents,

 more the thickness of adhesive is increased, more the values of the stresses decrease on the level of the free edges and the distribution tends to being uniform.

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Dedicated to professor Gh. Marcu at his 80th anniversary

MODELING AND SIMULATION OF THE AMMONIA ABSORPTION PROCESS IN SODIUM CHLORIDE SOLUTION USING CHEMCAD

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ABSTRACT. In this paper the mathematical modeling and the simulation results for ammonia absorption in sodium chloride solution (brine) have been presented. The ammoniacal brine is used to obtain sodium bicarbonate and sodium carbonate in soda ash plants (according to the Solvay process).

The ammonia absorption process is performed in the absorption columns sequence. Because the absorption process is an exothermic process, the absorption columns are provided with cooling systems.

Modeling and simulation of the ammonia absorption in sodium chloride solution were done using ChemCAD software package. The evolutions of the process parameters were studied during the absorption process. The model and the simulation results proved to be a reliable tool for analyzing the absorption processes and can be used to improve the real plant operation.

1. INTRODUCTION

Sodium carbonate is a common inorganic industrial chemical, also known as soda ash (Na_2CO_3) . The synthesis process of soda ash (sodium carbonate) using Solvay process is done starting from sodium chloride, limestone, coke and ammonia as raw materials [1, 2].

The natural sodium chloride solution (brine) is extracted from soil and purified (removal of solid impurities by filtration and removal of calcium and magnesium ions by precipitation). Into the purified sodium chloride solution, ammonia is absorbed (the recovered ammonia from the residual liquid phase is used). After ammonia absorption, the solution is carbonated with gaseous carbon dioxide coming from two main sources: thermal decomposition of the limestone and sodium bicarbonate calcination process. After carbonation of ammoniacal brine, a suspension of sodium bicarbonate results. Sodium bicarbonate is filtered and the residual liquid phase is treated with calcium hydroxide solution (slaked lime) in order to recover the ammonia

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from ammonium salts (ammonium chloride, carbonate, bicarbonate etc.). Sodium bicarbonate resulted after filtration is washed, dried and calcined in order to obtain sodium carbonate (soda ash).

2. MODELING AND SIMULATION OF THE PROCESS

The ammonia absorption process is done using an absorption columns sequence. Because the absorption process is exothermic the columns are provided with cooling systems. The ammonia absorption process is presented in figure 1 [1,2].

The purified brine is distributed in the ammonia recovering column I 2 and in ammonia recovering column III 4. The sodium chlorine solution (brine) rich in ammonia passes in the ammonia recovering column II 3, and into the first absorber 6. In the absorbers 6 and 7, the brine flows in counter-current with cold gas from distillation (the gas from distillation are cooled at 55-60°C, in the refrigerator 8). During the ammonia absorption process, the temperature increases and reaches 60-63°C to the bottom of the absorption column 6. The ammonia concentration in the first absorber is 15-20 DN. In the second absorber 7, the temperature is kept constant in the interval of 60-65°C, by cooling. The ammoniacal brine from the second absorber 7 (with concentration 100-106 DN/20 ammonia and 30-35 DN/20 carbon dioxide) is cooled at 30-35°C, in the refrigerator 9 and collected in the ammoniacal brine tank 10 in order to be send to carbonation process. The condensed phase is collected in tank 11 and sent to ammonia distillation. The outlet gas from the first absorber 6 contains 9-17% ammonia and 55-60% carbon dioxide. The ammonia from the outlet gas of the absorber 6 is absorbed in the recovering column 3, and the rich carbon dioxide gas passes to the calciner gas collector.

The main ionic reactions of the ammonia absorption in sodium chloride solution (brine) are presented below [2]. Most of the authors agreed to the following description of the ionic reactions that take place during the absorption process:

$$H_2O \leftrightarrows H^+ + OH^-$$
(1)

$$CO_2 + H_2O \leftrightarrows HCO_3^- + H^+$$
(2)

- $HCO_3^{-} \leftrightarrows CO_3^{2-} + H^+$ (3)
- $NH_3 + H_2O \leftrightarrows NH_4^+ + OH^-$ (4)
- $NH_3 + HCO_3^{-} \leftrightarrows NH_2COO^{-} + H_2O$ (5)
- $NaCl \leftrightarrows Na^{+} + Cl^{-}$ (6)
- $NH_4OH \implies NH_4^+ + OH^-$ (7)



MODELING AND SIMULATION OF THE AMMONIA ABSORPTION PROCESS IN SODIUM ...

Figure 1. Ammonia absorption process [1]

purified brine tank, 2 – ammonia recovering column I (from the filtration gas),
 ammonia recovering column II (from absorption stage), 4- ammonia recovering column III (from the carbonation column gas), 5 – gas cleaner, 6 – ammonia absorber I, 7- ammonia absorber II, 8 – refrigerator of the distillation gas, 9 – refrigerator of the ammoniacal brine, 10- ammoniacal brine tank, 11- condensed tank

The parameters used for modeling and simulation of the ammonia absorption process in sodium chloride solution (brine) are presented in the tables 1, 2 and 3.

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Table 1.

Parameter	Measuring Unit	Gas coming from sodium bicarbonate filtration process	Gas coming from carbonation columns	Gas coming from ammonia distillation process	
Temperature	[°C]	25	50	60	
Pressure	[bar]	0.5	1	0.9	
CO ₂	[mole %]	3.20	5.36	14.33	
CO	[mole %]	0	1.00	0.03	
O ₂	[mole %]	18.42	1.50	0.05	
N ₂	[mole %]	73.68	68.52	0.33	
H ₂ O	[mole %]	3.50	5.33	14.42	
NH ₃	[mole %]	1.20	18.28	70.83	
Flow	[kg/h]	850	1016	1136.15	

The properties of the inlet gaseous streams

Table 2.

The properties of the inlet liquid stream (purified brine)

Parameter	Measuring Unit	Value
Temperature	[°C]	20
Pressure	[bar]	1
NaCl	[weight %]	24,63
H ₂ O	[weight %]	75.47
Flow	[kg/h]	10600

Table 3.

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Parameters of the absorption columns					
Parameter	No. of stage	Feed tray for liquid stream	Feed tray for gas stream	Cooling duty	
First absorber	6	1,4	6	0	
Second absorber	6	1	6	1* 100 ⁷ J/h	
Ammonia recovering column I	4	1	4	0	
Ammonia recovering column II	6	1	6	0	
Ammonia recovering column III	6	1	6	0	

The modeling and simulation of the ammonia absorption process were done using ChemCAD (version 5.1.3) software package. The electrolyte package was used as thermodynamic option for simulation of the ammonia absorption process [4,5].

The main window of the application developed for ammonia absorption process is presented in the figure 2.

MODELING AND SIMULATION OF THE AMMONIA ABSORPTION PROCESS IN SODIUM ...



Figure 2. Simulation of the ammonia absorption process using ChemCAD

3. RESULTS AND DISCUSSIONS

The simulation results of ammonia absorption process in brine (sodium chloride solution) using ChemCAD, are presented below.

The properties of output gaseous streams resulted from the simulation in case of the recovering columns and absorber columns are presented in table 4 and 5.

Tabel 4.

The properties of the gaseous streams leaving the ammonia recovering columns

Parameter	Measuring	Gas from	Gas from	Gas from
	Unit	ammonia	ammonia	ammonia
		recovering	recovering column	recovering column
		column I		=
Temperature	[°C]	20	20	26
Pressure	[bar]	1	1	1
CO ₂	[mole %]	2.2	0	49.45
CO	[mole %]	0	1.38	2.85
O ₂	[mole %]	19.17	2.07	6.67
N ₂	[mole %]	76.77	94.7	38.07
H ₂ O	[mole %]	1.86	1.85	2.9
NH ₃	[mole %]	0	0	0
Flow	[kg/h]	820	775	80

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Tabel 5.

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Parameter	Measuring	First absorber	Second absorber
	Unit		
Temperature	[°C]	39	36
Pressure	[bar]	1	1
CO ₂	[mole %]	0.16	0.12
CO	[mole %]	5.58	7.82
O ₂	[mole %]	8.11	14.83
N ₂	[mole %]	79.11	71.57
H ₂ O	[mole %]	5.57	4.70
NH ₃	[mole %]	1.47	0.95
Flow	[kg/h]	8.2	8.6

The ammoniacal brine resulted in the absorption process has the following properties: temperature 62°C, pressure 1 bar, flow 12000 kg/h, and composition (mass %): 64.88 % water, 15 % ammonia, 5.34 % NH₄OH, 3.11 % NH₂COO⁻, 8.52 % Na⁺, 1.39 % CO₃⁻², 0.56 % HCO₃⁻¹, 1.93 % NH₄⁺, 13.14 % Cl⁻.

In the sequence of absorption columns, the temperature variations are: for the ammonia recovering column I the change is of 20-26°C, for the ammonia recovering column III is of 20-39.5°C, for the ammonia recovering column II the temperature is constant, for the first ammonia absorber is of 36-39°C and for the second ammonia absorber is of 30-60 °C.

The variations of the temperature for the ammonia recovering columns are presented in the figures 3 and 4.



Figure 3. Variation of the temperature in the ammonia recovering column I

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Figure 4. Variation of the temperature in of the ammonia recovering column III

The simulation results presented above were compared with data collected from real plant operation. The operation data collected from a real plant are presented in the table 6 [1, 2, 3].

Tabel 6.

The properties of the gaseous streams leaving the ammonia recovering columns (data collected from a real plant operation)

Parameter	Measuring	Ammonia	Ammonia	Ammonia
	Unit	recovering	recovering	recovering
		column I	column III	column II
Temperature	[°C]	-	-	25-30
CO ₂	[mole %]	1.3	3.2	69.7
H ₂ O	[mole %]	1.8	3.2	5.1
NH ₃	[mole %]	0.06	0.19	0.06
Inert	[mole %]	96.9	93.4	25.2

From the comparison of the simulation and real plant data, one can observe a close similarity between simulation results and experimental data. This fact validates the application developed for simulation the process and proves the utility of the model in analyzing and optimization of the real plant operation.

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4. CONCLUSIONS

Modeling and simulation of the ammonia absorption process in sodium chloride solution (brine) was done using ChemCAD software package (version 5.1.3).

The evolutions of the process parameters (liquid and gaseous flows, composition of the streams, temperatures) were studied during the carbonation process. The simulation results were compared with real plant operation data in order to validate the application developed for the absorption process.

The mathematical model and the simulation results proved to be a reliable tool for analyzing and optimizing the real plant operation of the ammonia absorption in sodium chloride solution used in soda ash manufacturing process according to the Solvay technology.

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