

Sorosite $(\eta$ -Cu₆Sn₅)-bearing native tin and lead assemblage from the Mir zone (Mid-Atlantic Ridge, 26°N)

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Abstract – A number of small, irregular-shaped and spherical shiny metallic particles have been found in the sediments from the Mir zone, Trans-Atlantic Geotraverse hydrothermal field (Mid-Atlantic Ridge, 26°N). The phase variety of the particles examined is represented by metallic tin, tin-rich lead, and tin-copper phases. A detailed mineralogical study of these particles was carried out using optical microscopy, nuclear microscopy, scanning electron microscopy, electron microprobe, proton microprobe and X-ray diffraction analysis. Tin-lead grains have the typical eutectic microtexture of the metal components. Tin-copper grains are formed from single crystals of sorosite, η -Cu₆Sn₅. The Sn-Pb-Cu complex grains are composed of fine stannoan lead ($Pb_{0.74}Sn_{0.26}$) and tin crystals as well as fine (or occasionally larger) sorosite $(Cu_{6,1}Sn_{4,9})$ idiomorphic crystals, in a tin-lead matrix forming eutectic microtexture. On the basis of data obtained, a natural origin is proposed for the examined Sn-Pb-Cu association, and its parent relations with the tectono-magmatic events in the rift zone. This association has probably been formed (1) during the hydrothermal seepings through the Trans-Atlantic Geotraverse sediment cover, or (2) during the evolution of ridge crest magnatic systems. Crystallisation sequence from an initial melt with falling temperature is: firstly sorosite (Cu_6Sn_5) ($T \le 380^{\circ}C$) \rightarrow crystallisation of tin $(T \le 227^{\circ}\text{C}) \rightarrow \text{crystallisation}$ of Sn–Pb eutectic mixture, composed of tin + stannoan crystals lead $(T \le 183^{\circ}\text{C}) \rightarrow$ limited tin exsolutions in stannoan lead (T << 183^{\circ}\text{C}). Sn–Pb–Cu grains might be present as accessory minerals in the basic rocks of the east rift wall, which have undergone degradation, permitting their deposition into rift valley sediments. © 2001 Ifremer/CNRS/IRD/Éditions scientifiques et médicales Elsevier SAS

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Résumé – Association de Sn-Pb-Cu dans les dépôts de la zone hydrothermale de Mir (dorsale médio-Atlantique, 26°N). De petites particules métalliques brillantes de forme irrégulière et sphériques ont été trouvées dans les dépôts de la zone hydrothermale de Mir (dorsale, médio-Atlantique, 26 °N). Les particules examinées représentent l'étain métallique, l'étain riche en plomb et l'étain riche en cuivre. Une étude minéralogique détaillée de ces particules a été effectuée en utilisant la microscopie optique, la microscopie nucléaire, le microscope électronique à balayage, la micro-sonde d'électrons, la micro-sonde de protons et la diffraction de rayons X. Les grains étain-plomb présentent une microtexture typique des composants métalliques. Des grains d'étain-cuivre sont formés de cristaux simples de sorosite η -Cu₆Sn₅. Les grains complexes de Sn–Pb–Cu se composent de cristaux fins d'étain–plomb (Pb_{0.74}Sn_{0.26}) et d'étain ; les cristaux idiomorphes du sorosite ($Cu_{6,1}Sn_{4,9}$) fin dans une matrice d'étain-plomb formant la microtexture eutectique. Une origine « normale » est proposée pour l'association examinée de Sn-Pb-Cu, et ses relations de parenté avec les événements tectono-magmatiques dans la zone de rift. Cette association a été probablement formée (1) pendant une infiltration hydrothermale par la couverture de dépôt géotraverse trans-Atlantique, ou (2) pendant l'évolution des systèmes magmatiques des dorsales. La succession de cristallisations à partir d'une fusion initiale avec la température est la suivante : le $(T \leq 380^{\circ}\text{C}) \rightarrow \text{cristallisation}$ des en baisse sorosite (Cu_6Sn_5) cristaux d'étain $(T \le 227^{\circ}\text{C}) \rightarrow \text{cristallisation}$ du mélange eutectique de Sn-Pb, composé d'étain + étain-plomb $(T \le 183^{\circ}\text{C}) \rightarrow$ exsolutions limités d'étain dans étain-plomb ($T \le 183^{\circ}\text{C}$). Les grains de Sn-Pb-Cu pourraient être présents comme des minéraux accessoires des roches basiques de la paroi est du rift qui ont subi la dégradation, permettant leur dépôt dans les sédiments de la vallée de rift. © 2001 Ifremer/CNRS/IRD/Éditions scientifiques et médicales Elsevier SAS

hydrothermal field / Mid-Atlantic Ridge / native Sn and Pb / sorosite

zone hydrothermale / dorsale médio-Atlantique / étain et plomb natifs / sorosite

1. INTRODUCTION

Findings of native tin, established mainly in placer materials, have been known for a long time and data for them may be found in the basic references (Hosking, 1974; Ramdohr, 1975). Undoubted native tin in bedrocks has been first found in a calcite–quartz–sulfide association at Nesbitt La Bine Mines (Silman, 1954). More recently, native tin and tin alloys have been described in various types of terrestrial (Ramdohr, 1975; Kovalskii, 1981; Novgorodova, 1994) and lunar (Gay et al., 1970) rocks, and impact materials (Gurov and Kudinova, 1987).

In oceanic environments a wide spectrum of native minerals has been established in the mafic and ultramafic lithologies of the crust (Leinen et al., 1986; Shterenberg et al., 1993; Prichard et al., 1996), metalliferous sediments overlying the basalt basement (Jenkyns, 1976; Marchig et al., 1986), seafloor massive sulfides (Hannington et al., 1986; Zierenberg et al., 1993) and submarine hydrothermal plumes (Jedwab and Boulègue, 1984). Native tin and tin alloys were found in pelagic sediments (Arsamakov et al., 1988), and metalliferous sediments of the Galapagos Rift (Lazur et al., 1989) and East Pacific Rise (Dekov et al., 1996). We report here the first finding of a ternary sorositetin-lead (Sn-Pb-Cu) association in the near sulfide mound sediments of the Trans-Atlantic Geotraverse (TAG) hydrothermal field, Mid-Atlantic Ridge, 26°N.

1.1. Geologic setting

Since the discovery of high-temperature hydrothermal activity in the Mid-Atlantic Ridge (MAR) (Rona et al., 1986), the TAG hydrothermal field (MAR 26°N) has been the classic area for studies of recent hydrothermal discharge and mineralisation on slow-spreading centres. The TAG field, which comprises three sulfide mounds (one active and two dead), has been the focus of a number of studies over the past fifteen years (Campbell et al., 1988; Lisitsyn et al., 1989; Rona et al., 1993; Humphris et al., 1998). In an effort to collect novel data about the tectono-magmatic activity and related events at the TAG field, the 15th cruise of the RV *Akademik Mstislav Keldysh*, with the DSV *Mir-1* and *Mir-2* aboard, took place in the spring of 1988. The results of geophysical, tectonic, and hydrochemical observations, as well as

studies of basement rocks, hydrothermal deposits and rift valley sediment cover have been published (Lisitsyn, 1992).

The TAG hydrothermal field lies on the rift valley floor of the slow-spreading MAR (half rate of accretion 1.1-1.3 cm year⁻¹; McGregor et al., 1977) between the Atlantis (30°40'N) and Kane (24°N) fracture zones (Sempéré et al., 1990) (figure 1A). The rift valley is characterised by an asymmetric structure. The east wall is higher and steeper than the west wall (Rona et al., 1986; 1993) and exposes the sheet-dike complex and gabbro layer on its lower part (Lisitsyn, 1992). The active high-temperature sulfide mound is located east of the spreading axis on the rift valley floor. It is a large and steep-sided edifice. The central part consists of a black smoker complex surrounded by a wide platform with a group of white smokers and an apron of metalliferous sediments (Rona et al., 1986; 1993). Two inactive sulfide deposits (named Mir and Alvin) occur on the lower east wall of the rift valley and are undergoing extensive mass-wasting (Lisitsyn et al., 1989; Lisitsyn, 1992; Rona et al., 1993). The Mir relict hydrothermal zone showing various stages of weathering was found in 1988 (Lisitsyn et al., 1989). It is situated between the low-temperature zone, some 300 m higher on the wall (Rona et al., 1984), and the active high-temperature sulfide mound (Rona et al., 1993).

2. MATERIALS AND METHODS

The materials we used in this study were subsamples of a gravity core taken during the 15th cruise of the RV Akademik Mstislav Keldysh. The core was collected close to the Mir zone (figure 1B). The sediment cover surrounding the Mir mound is composed of nanofossil-foraminiferal and foraminiferal-nanofossil oozes with intercalations of metalliferous sediments and layers rich in basaltic clasts (figure 1C). The samples have been hermetically stored at room temperature in polythene boxes (100 mL) since the time of collection. Before analyses all samples were washed with distilled water to remove interstitial salts. The samples were separated into size fractions (> 2.0,2.0-1.0, 1.0-0.1,0.1 - 0.01, 0.01-0.001, < 0.001 mm) by wet sieving in a controlled lab environment (20°C).

The extremely low contents of metallic particles in oceanic sediments (Dekov et al., 1996) prevented the use

of routine bulk mineralogical techniques for particle diagnostics. Due to the difficulty to investigate the possible metallic particles from the fine (< 0.10 mm) sediment fractions (Dekov et al., 1996), we examined exclusively the coarse fractions (> 0.10 mm) under a stereomicroscope. Single metallic grains were handpicked with a steel needle for further mineralogical studies. Preliminary diagnostics of the separated grains was done by means of semi-quantitative energy-dispersion analyses (EDS).

The micromorphology, size and chemical composition (point analyses and area scanning onto polished sections) of the Sn–Pb–Cu specimens were investigated by scanning electron microscope Jeol T-300 with a Link 860-500 EDS and ZAF/PB program, and a Philips SEM-515 with an Edax PV 9100 EDS system (with an operating voltage of 20–25 kV and electron beam diameter of 1 μ m). The chemical composition (point analyses) was determined by employing a Cameca microprobe unit (accelerating voltage of 15 kV; beam current of 15 nA; electron beam diameter of 1 μ m). The following standards and X-ray lines were used: pure synthetic tin (Sn L_{α}), pure synthetic copper (Cu K_{α}) and PbTe (Pb L_{α}). The detection limits were 0.5% (weight).

Together with the conventional electron microscopy we applied a nuclear microscopy to the analysis of native $Sn-(\pm Pb) - (\pm Cu)$ grains. The chemical composition (point analyses) of representative specimens was analysed using the Oxford scanning proton microprobe facility (Grime et al., 1991; beam of 3 MeV protons focused to 1 µm diameter with a current of 100 pA) with a proton induced X-ray emission (PIXE) system and GUPIX program (Maxwell et al., 1995). The highly quantitative PIXE technique (Johansson and Campbell, 1988; an accuracy of 5-10% and a sensitivity in the order of 1–10 ppm) can detect all elements heavier than Na simultaneously. X-rays were detected using a lithiumdrifted silicon detector with 70 mm² active area positioned at 45° to the beam at a distance of 70 mm. The spatial distribution of different phases within the samples was investigated by scanning the beam over different sized areas (between $750 \times 750 \ \mu m^2$ and $25 \times 25 \ \mu m^2)$ of the sample and forming PIXE maps of the major elements. The X-ray diffraction patterns of the metallic particles were obtained with a 57.3 mm Gandolfi camera employing Ni-filtered Cu K_{α} radiation without internal standard (with an operating voltage of 40 kV and a beam current of 19 mA).

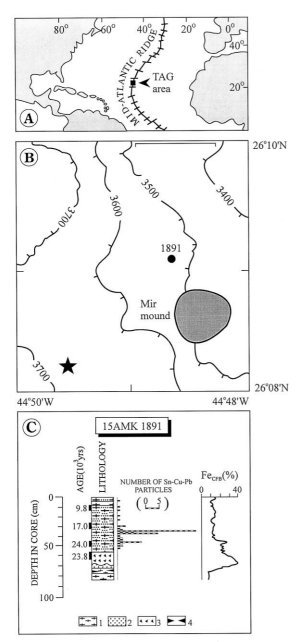


Figure 1. A. Index map showing the location of the Trans-Atlantic Geotraverse area at the Mid-Atlantic Ridge. **B.** Bathymetric map (based on Rona et al., 1993) of the investigated area showing the position of sediment sample station (solid dot), the active high-temperature sulfide mound (star) and the inactive Mir mound (bar in upper right is 1 km long). **C.** Stratigraphic section showing the lithology, number of Sn–Pb–Cu particles found and vertical distribution of the main chemical tracer of hydrothermal activity - Fe (on a carbonate-free basis) in the studied core (based on Lisitsyn et al., 1989). 1 = nanno–foram and foram–nanno oozes; 2 = disseminated hydrothermal component; 3 = clastic sulfide–oxyhydroxide flow; 4 = nontronitic lenses.

3. RESULTS

The EDS analyses of the separated metallic grains (total of 88) have established metallic tin (14% of total grains), tin–lead (48%), tin–copper (12%) and tin–lead–copper (26%) phases. About 80% of total particles found have an irregular shape while the rest 20% are spherical and rounded. Microprobe studies onto polished sections revealed that most of the tin and tin–lead particles contained traces of Cu. This is why we concentrated our investigations on the Sn–Pb–Cu occurrences.

Metallic particles were found to occur along with oxides and hydroxides (chromite, cuprite, tenorite, magnetite, maghemite, hematite, ilmenite, rutile, goethite), sulfides (chalcocite, pyrite, marcasite, chalcopyrite), halogenides (atacamite), sulfates (barite), and silicates (quartz, olivine and plagioclase phenocrysts).

The Sn–Pb–Cu particles are soft, lamella- to irregularshaped (*figure 2A*), spherical (*figure 2B*) or drop-like, silver-white to grey–white in colour and with metallic lustre. A typical eutectic microtexture (*figure 2C,D*) is observed on the smoothed grain surfaces. Findings of tin (traces of Pb and Cu; *figure 2E*) and needle-like tin–copper (*figure 2F*) crystals at the periphery of the Sn–Pb–Cu particles are fairly rare. Occasionally, the metallic particles are partly covered by a mass of red–brown Fe-oxyhydroxides and coccoliths (*figure 2E,F*). Coatings of secondary oxides, hydroxides and oxychlorides are present on some particles. They are thin, whitish grey, soft and fragile. The bulk (90%) of the Sn–Pb–Cu grains are concentrated in 0.10–0.25 mm fraction.

In reflected light (polished section examinations) the metallic particles are observed to consist almost entirely of a soft, highly reflective, white intergrowth, corresponding to a eutectic texture (figure 3C) of platy white pure tin (EDS study) with greyish-white Sn-rich lead. Within this eutectic are observed fine micro-inclusions as well as single brownish-rose 'porphyritic' crystals of a tin-copper phase. This rose mineral is non-bireflectant in air and oil, and exhibits pronounced anisotropism in oil. Its reflectance is lower than in tin and lead-tin phases. Despite their obvious softness, the tin-lead-copper particles take a good polish, with only a few scratches. The relative hardness of tin-copper crystals is greater than those of tin and lead-tin. The tin-copper crystals much resemble sorosite in their optical properties, appearance as well as associated minerals (native tin and lead).

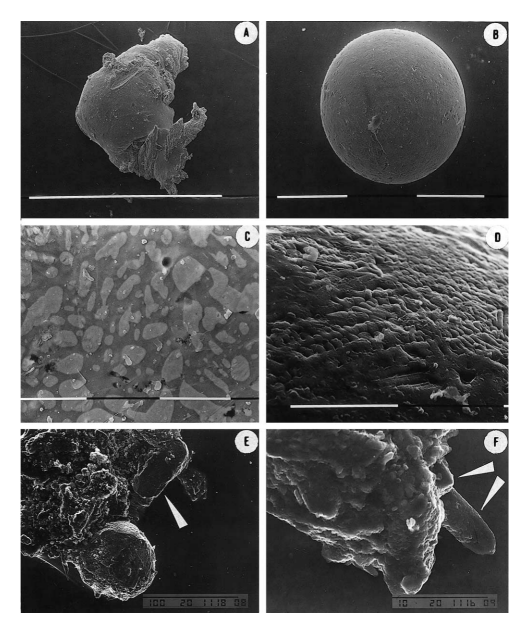


Figure 2. Scanning electron microscope photographs secondary electron image of: A. irregular-shaped Sn-Pb-Cu grain (H105I.S48); B. spheri-Sn-Pb-Cu cal grain (H105I.S60); C. eutectic microtexture of the Sn-Pb-Cu grain shown at (A) (enlargement of the central grain surface); lead-tin inclusions (light grey) in a tin matrix (grey); D. eutectic microtexture of the Sn-Pb-Cu grain shown at (B) (enlargement of the upper grain surface); E. tin crystal (arrow) at the periphery of a Sn-Pb-Cu grain (H105II.S40); F. tin-copper crystals (arrows) at the periphery of a Sn-Pb-Cu grain (H105II.S12). Scale bars equal to 1000 μm (A), 100 μm (B, E) and 10 µm (C-F).

PIXE mapping of the polished sections (*figure 3*) supports our microscopic observations: the particles studied are composed of fine lead-tin and tin-copper inclusions even distributed within a tin matrix. The tin-copper phases occur as tiny, probably elongated grains (*figure 2F*) and very seldom as bigger idiomorphic crystals of 'porphyritic' type (*figure 3A,B*). Tin and lead-tin phases are uniformly distributed within the eutectic mixture and the rare tin-copper crystals are notable on that background (*figure 3*). The lead-tin inclusions in the tin matrix have an uniform composition close to $Pb_{83.5}$, $Sn_{16.5}$ (in percent of the weight, i.e. $Pb_{74.3}Sn_{25.7}$ in percent of atoms) (*table I*) and fairly constant average atomic ratio of $Pb/Sn = 2.9 \pm 0.05$. They have to be defined as stannoan lead, i.e. solid solution of lead with high Sn content in the crystal lattice. Phases with similar compositions have only been found in alluvial placers from the Central Ural Mountains (Aleksandrov, 1955), and basic rocks from the Siberian Platform (Okrugin et al., 1981; see also *table I*).

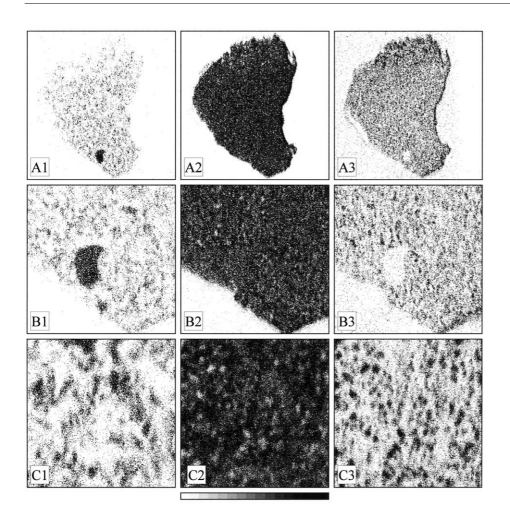


Figure 3. Internal microtextural peculiarities of Sn-Pb-Cu association (H105I.S48). A. Cross section of the Sn-Pb-Cu grain. B. Tin-copper crystal (η-Cu₆Sn₅-type) in the tin-lead eutectic matrix (enlargement of the lower part of the section shown in (A)). C. Eutectic texture of the tin-lead matrix with fine tin-copper crystals disseminated (enlargement of the upperright part of the scan area shown in (B)). Proton-induced X-ray emission scans in CuKa (A1-C1), SnL α (A2–C2) and PbM α (A3–C3). Scale bars (lower-right) in µm.

Tin-lead occurrences from the East Pacific Rise metalliferous sediments differ in composition from the TAG Sn-Pb-Cu particles (*table I*).

The composition of tin–copper crystals from the particles studied is also uniform and fairly constant (*table I*). Their average crystallochemical formula is $Cu_{6.1}Sn_{4.9}$, i.e. very close to sorosite (η -Cu₆Sn₅) (Barkov et al., 1998) defined at Baimka placer, despite the relative excess in Cu and the absence of Sb in the composition of the studied crystals. Similar tin–copper phases were also found in gold–platinoid concentrates from southeast Borneo (Stumpfl and Clark, 1965), tin ores from Panasqueira mine, Portugal (Clark, 1972), alluvial heavy mineral concentrates from Rio Tamaná, Colombia (Rose, 1981), some gold–silver deposits in Russia (Gamyanin et al., 1981), and soil and stream sediments from the US Virgin Islands (Alminas et

al., 1994; *table I*). We could not perform correct quantitative microprobe analyses on the thin, fragile secondary rind of the particles because of its bad polishing.

Area scanning applied to sections $(200 \times 200 \,\mu\text{m})$ of the eutectic tin–lead–copper mixture gave the following mean composition (in percent of weight): Sn 76.4 ± 0.9, Pb 16.6 ± 0.05, and Cu 7 ± 0.9 (respectively in percent of atoms: Sn 77.3 ± 1.5, Pb 9.6 ± 0.05, and Cu 13.1 ± 1.5). These data are indicative of the probable composition of the initial melt.

During the semi-quantitative EDS microprobe investigations, some trace elements were detected: Zn (in all types of particles – tin–lead–copper, lead–tin and tin–copper), Fe (in tin–copper particles) and Mn (in one lead–tin grain). Table I. Chemical composition of native Sn° and $Sn-(\pm Pb)-(\pm Cu)$ phases from different natural objects.

Refer- ence	Rudashe al. (198		Kovalski (1985)	ii and O	leynikov								Okrugir (1981)	n et al.			Glavatsk (1990)	ih		
Object and area of investi-	Ultrabas rocks, K skoe mo Russia	Koryak-	Kimberli pipes, ki lite breco Yakutiya	mber- cia,	Kimberl Russia	ite pipes	, serpent	inized g	arnet-spi	nel perido	otites, Y	akutiya,	Basic ro Russia	ocks, Sib	erian Pla	tform,	Basalts,	Kamchat	tka, Russ	ia
gation Sample No.			Russia OL-55, 44-77		OL- 210, 44-75		OL- 214, 42-28		OL- 214, 45-14		OL- 149, 44-46		OB- 242-3	OB- 54-4D	OL- 54-2					
Phase	tin	lead	tin	lead	tin	lead	Cu ₆ Sn ₅ -	Cu ₃ Sn-		Cu ₆ Sn ₅ -		Cu ₆ Sn ₅ -	tin	tin	lead	tin	lead	tin	lead	tin
Sn Pb Cu Zn Sb Total	99.00 1.27 - - 100.27	96.30 	2.39 - - 0.51	2.04 95.77 - 97.81	100.58 0.21 0.06 - 100.85	2.46 98.41 0.08 - 100.95	type 61.09 - 37.68 - - 98.77	type 42.16 1.46 54.51 - 98.13	98.30 0.25 0.30 - 98.85	1.64	99.19 0.27 0.24 - 99.70	type 66.10 0.16 33.87 - 100.13	97.11 0.48 	0.13	85.53 - 0.13	92.89 4.22 2.31 99.42	99.42	94.27 - - 4.73 99.00	0.30 99.96 - - 100.26	96.75 - - 3.26 100.01

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Reference	Gamyanin et al (1981)		Ippatyeva (1981)	Lazur et al. (1988)			Gurov and Ku (1987)	dinova	Rose (1981)	Clark (1972)		Kucha (1981)	Lawrence (1951)
Object and area of in- vestigation	Granitoids, Sou Verkhoyanskii rium, Russia	synclino-		Sandstones, bases, Ural,	clay shales and Russia		Impact materia oritic crater El Russia		Alluvial heavy min- eral con- centrates, Rio Tamaná, Colombia	Tin ores, Pana Mine, Portugal	-	Zechstein copper de- posits, Lu- bin Mine, Poland	Alluvial placers, Emmaville, NSW, Aus- tralia
Phase	Cu ₆ Sn ₅ -		tin	tin			Sn-Pb-Cu		Sb-bearing	η '-Cu ₆ Sn ₅		lead	tin
Sn	type 67.70	66.60	99.00) 99.60	63.76	63.34	63.00	67.00	η-Cu ₆ Sn ₅ 56.12	61.10	61.20	-	- 95–92
Pb	-				-		35.00	30.00				100.00	
Cu	32.30	33.40	-		36.14	36.98	0.30	0.30	35.99	38.70	38.50	-	
Zn	_	_	1.60) –	_	_	_	-			_	-	
Sb	_	_	-	- –	_	-	_	-	7.89	_	_	-	
Total	100.00	100.00	100.60) 99.60	99.90	100.32	98.30	97.30	100.00	99.80	99.70	100.00) 95-92

Table I. Continued.

Cu - - - - 38.4 38.2 38.2 36.12 35.33 35.96 36.45 36.88 36.9 Ni - - - - 0.0 0.3 0.6 -	area Franklin		stream_s								Barkov e	et al. (1998)				
Sample NMNH # SC514A SJ601 A SC514 SJ601 29 10C 10D SJ738-1 2 3 4 5 6 No. C6061-1 C6061-1 2-16<			i stream-so	ediments,	US Virg	gin Islan	ds, Grea	ter Antilles	s Island ar	rc	Gold pla	cer deposit	, Baimka	River, west	tern Chuko	tka, Russian	Far East
Phase lead tin lead lead lead Cu_6Sn_5 Cu_6Sn_5 sorosite sorosite <t< th=""><th>ple NMNH</th><th></th><th>SJ601A</th><th>SC514</th><th>SJ6</th><th>01 29</th><th>)</th><th>10C</th><th>10D</th><th></th><th>1</th><th>2</th><th>3</th><th>4</th><th>5</th><th>6 9</th><th></th></t<>	ple NMNH		SJ601A	SC514	SJ6	01 29)	10C	10D		1	2	3	4	5	6 9	
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Pb 99.5 0.3 100.6 94.6 99.7 101.0 -																	n 98.17
Cu - - - - 38.4 38.2 38.2 36.12 35.33 35.96 36.45 36.88 36.9 Ni - - - - 0.0 0.3 0.6 -																	90.17
Ni - - - - 0.0 0.3 0.6 -<																	-
Sb - - - - - 4.93 4.77 4.89 0.91 0.87 1.1 Fotal 99.5 99.9 100.6 100.6 99.9 101.0 99.9 99.6 99.9 99.46 99.53 99.95 100.73 100.5 Reference Dekov et al. (1996) Present study - - - Hydrothermal sediments, Mir mound, TAG field, MAR 26°N area of in-		_	_	_	_	_	_										-
Total 99.5 99.9 100.6 100.6 99.9 101.0 99.9 99.6 99.9 99.96 99.46 99.53 99.95 100.73 100.5 Reference Dekov et al. (1996) Present study Dbject and Metalliferous sediments, EPR 21°S area of in- Hydrothermal sediments, Mir mound, TAG field, MAR 26°N		_	_	_	_	_	_	-	-		- 0.92	2 1.18	3 1.2	- 6			-
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vestiga-	rence Dekov	et al. (199	6)						Pres	sent study							
iion Sample H2.S6 H22.S10 H105I.S48 No.	ect and Metallif of in-			PR 21°S							sediments	, Mir mou	nd, TAG f	ìeld, MAR	26°N		
Phase tin copper tin lead tin lead tin Cu ₆ Sn ₅ - Lu	ect and Metallif of in- iga-		ments, EF						Hyd	lrothermal	sediments	, Mir mou	nd, TAG f	ield, MAR	26°N		
	ect and Metallif of in- iga- ple H2.S6	ferous sed	ments, EF	510	1 1	in	lead	tin	Hyd H10: Cu ₆ S	Irothermal 5I.S48 Sn ₅ - Cu _o	₅Sn₅- Cι	ı₀Sn₅- C	u ₆ Sn ₅ - (pe t	Cu ₆ Sn ₅ - 1 type		lead-tin tir	1
	ect and Metallif of in- iga- ple H2.S6 se tin	ferous sed	H22.S	510 lead 99.21	5.15				Hyd H10: Cu ₆ type 99.27	Irothermal 5I.S48 Sn_5 - Cu_6 yp	₅ Sn ₅ - Cu e tyj	ı ₆ Sn₅- C pe ty	u ₆ Sn ₅ - (pe t	Cu ₆ Sn ₅ - 1 type	ead-tin	16.47	
Cu 0.33 79.16 - 0.03 0.10 0.09 0.22 39.68 39.96 39.82 40.46 40.13	ect and Metallif of in- iga- ple H2.S6 se tin 97.	copper .61	H22.9 tin 5.55	510 lead 99.21	5.15 94.27	97.4 2.4	9 ⁻ 1 9	7.51 9 1.61	Hyd H10: Cu ₆ type 99.27 0.51	Irothermal 5I.S48 Sn₅- Cu ₆ y typ 59.98 -	₅ Sn ₅ - Cu e tyj 59.65 –	1 ₆ Sn₅- C pe ty 59.29 −	u ₆ Sn₅- (pe t 59.13 -	Cu ₆ Sn ₅ - 1 ype 59.82 -	ead-tin 16.73		
	ect and Metallif of in- iga- ple H2.S6 se tin 97.	copper .61	H22.9 tin 5.55	510 lead 09.21 0.79	5.15	97.4 2.4	9 1 - 9 0	7.51 9 1.61 0.09	Hyd H10: Cu ₆ type 99.27	Irothermal 5I.S48 Sn_5 - Cu, typ 59.98	₅ Sn ₅ - Cu e tyj 59.65	1 ₆ Sn ₅ - C pe ty 59.29	u ₆ Sn ₅ - (pe t 59.13	Cu ₆ Sn ₅ - 1 type 59.82	ead-tin 16.73 83.27	16.47	1 98.43 -
	ect and Metallif of in- iga- ple H2.S6 se tin 97.	copper .61	H22.5 H22.5 tin 5.55 0.16 -	\$10 lead 99.21 0.79 –	5.15 94.27 0.03	97.4 2.4 0.1	9 1 - 9 0	7.51 9 1.61 0.09	Hyd H102 Cu ₆ 3 type 99.27 0.51 0.22	51.S48 51.S48 Sn ₅ - Cu ₆ 59.98 - 39.68	58n₅- Cu e tyj 59.65 _ 39.96	l_6Sn_5 - C pe ty 59.29 - 39.82	$u_6 Sn_5 - 0$ pe t 59.13 - 40.46	Cu ₆ Sn ₅ - 1 type 59.82 - 40.13	ead-tin 16.73 83.27	16.47 84.55	
Fe – 0.13 – – – – – – – – – – – – – – –	ect and Metallif of in- iga- ple H2.S6 se tin 97. 0.	copper .61	H22.8 H22.8 tin 5.55 0.16 0.13	510 lead 09.21 0.79 - - -	5.15 94.27 0.03 –	97.4 2.4 0.1	9 · 1 9 0 ·	7.51 9 1.61 0.09 0.19	Hyd H102 Cu ₆ 3 type 99.27 0.51 0.22	51.S48 51.S48 Sn ₅ - Cu ₆ 59.98 - 39.68	58n₅- Cu e tyj 59.65 _ 39.96	l_6Sn_5 - C pe ty 59.29 - 39.82	$u_6 Sn_5 - 0$ pe t 59.13 - 40.46	Cu ₆ Sn ₅ - 1 type 59.82 - 40.13	ead-tin 16.73 83.27	16.47 84.55	
	ect and Metallif of in- iga- ple H2.S6 se tin 97. 0.	copper .61 : .33 79 	H22.8 H22.8 tin 5.55 0.16 0.13 0.31	510 lead 09.21 0.79 - - - - -	5.15 94.27 0.03 - -	97.4 2.4 0.1	9 / 1 9 0 (-)	7.51 9 1.61 0.09 0.19 -	Hyd H102 Cu ₆ 3 type 99.27 0.51 0.22 –	51.S48 51.S48 Sn ₅ - Cu ₆ 59.98 - 39.68 - - -	58n₅- Cu e tyj 59.65 _ 39.96	l_6Sn_5 - C pe ty 59.29 - 39.82	u ₆ Sn ₅ - (pe t 59.13 - 40.46 - -	Cu ₆ Sn ₅ - 1 type 59.82 - 40.13	ead-tin 16.73 83.27	16.47 84.55	

Values are in percent of weight.

All reflections typical for Sn_{syn} and Pb_{syn} were established on the X-ray diffraction (XRD) pattern of the Sn-Pb-Cu phase (table II). Basic reflections of sorosite or its synthetic equivalent η -Cu_{6.26}Sn₅ (PDF # 47-1575) were absent on the XRD pattern, although such a phase was undoubtedly identified by microprobe analyses in the particle studied. This could be due to relatively low content of the tin-copper crystals in the specimen examined and the close proximity of n-Cu_{6.26}Sn₅ standard basic lines (2.9640 Å (101), 2.1030 Å (110) and 2.0886 Å (102)) with those of the obtained pattern (2.92 Å (I = 6) and 2.06 Å (I = 3); table II). There is, however, a coincidence of some minor XRD peaks of the studied phase (indexed as Sn° and Pb° peaks) with those of sorosite (table II). The XRD patterns of previously studied tin-lead particles from the metalliferous sediments from the EPR 21°S suggest that these phases probably contains sorosite too (table II).

Estimated cell parameters of the tin phase are practically identical with the standard ones (*table II*). In contrast, the measured $a_{\rm Pb} = 4.947$ Å is less than the pure standard one ($a_{\rm Pb} = 4.9506$ Å; PDF # 4-686) (*table II*). This is obviously a result of the inserting through solid solution of Sn atoms (with atomic radius (1.58 Å) smaller than Pb (1.75 Å)) into the Pb cell.

4. DISCUSSION

4.1. Genesis of the Sn-Pb-Cu association

The occurrence of a large proportion of unorthodox native metals (other than noble Au°, Ag° and platinum group elements) and alloys had long been regarded as sampling artefacts. Indeed, it is difficult to understand how elements generally combined with oxygen or sulfur could exist in a metallic state in Nature, although a number of papers on this matter have been published during the past years (summarised by Ramdohr, 1975; Kovalskii, 1981; Novgorodova, 1994; etc.). We are well aware of the possibility that metallic occurrences found in the TAG sediments may be anthropogenic. In our previous works on native metals in oceanic hydrothermal sediments (Dekov et al., 1996; Dekov et al., 1999) we discussed the possibility of the technogenic nature of these occurrences. Here we will add some additional facts that lend support to the idea of the natural origin of studied particles:

- The probable initial melt composition of the investigated particles (Sn₇₆Pb₁₇Cu₇; weight percentage) differs from that of anthropogenic Sn-Pb systems. The most typical tin-lead solders are predominantly Sb-bearing. Some special types of solders contain up to 1.5% of Cu (weight) (for example: LSn60Cu1.5 - TGL 14908 standard; Neumann and Richter, 1979), while in the present case, the Cu content is considerably greater (table I). The material from tin-lead artefacts typically contains 100 ppm or more silver together with alumina spheres, while the natural material does not (Alminas et al., 1994). It is also indicative the fact that alloys with similar compositions have been less favourable objects of investigation during the study of the ternary Sn–Pb–Cu system (see data summarised by Chang et al., 1979; Marcotte and Schroder, 1983);

– The close association of a great number of Sn–Pb–Cu particles with native nickel, native copper and copper–zinc alloys which are characteristic for some oceanic sediments (Shterenberg, 1993) and continental ultrabasic and basic rocks (Kovalskii, 1981; Novgorodova, 1994). The association of the Sn–Pb–Cu particles with typical hydrothermal and basaltic minerals;

- The bulk of particles were found in core samples from sediment layers formed in pretechnogenic times (older than 9.10^3 years) (*figure 1C*). There are no data (radio-carbon, geochemical, lithologic, biostratigraphic) supporting resedimentation of the materials studied.

The facts discussed above and our previous investigations (Dekov et al., 1996; Dekov et al., 1999) favour a natural origin for the Sn–Pb–Cu particles from the TAG sediments. Different hypotheses may explain the natural origin of the studied particles:

- Fluvial origin. The remoteness of the studied area from the landmasses, and relatively high specific gravity of the Sn-Pb-Cu particles suggest that a fluvial origin of these metallic grains is unlikely.

– Aeolian origin. Eastern winds from the Sahara desert carry a lot of terrestrial dust to the Central Atlantic (Lisitsyn, 1972). Extensive alteration of the shape of the aerosol particles during transport in dust clouds forms typical surface microtextures that we have not observed. We think that the aeolian origin of studied particles is also not possible.

- Meteoritic origin. As a rule, volatile metals such as tin and lead are depleted in extraterrestrial matter (Mason and Melson, 1970). Meteorites burning during their passage through the atmosphere could not generate mi Table II. X-ray diffraction data for native Sn° and Sn-(± Pb)-(± Cu) phases from different natural objects.

Reference	Pres	ent study	Dek	ov et al.	(199	6)	Butuz al. (19		t Okrugin (1981)	ı et al.	Gurov Kudir (1987	nova	Toms (1989		Bark (1998	ov et al 8)	. PDF							
Object and area of investigation	sedin mou	rothermal ments, Mir nd, TAG , MAR N		alliferous 2 21°S	s sedi	ments,	Metal sedim Atlant Deep, Sea	ents, is-II	Basic ro Siberiar Platforr Russia	1	rial, n itic cr	ytgyn,	ilmen metas tites,	ite oma- Kavale- listrict, orie,	depor Baim River ern Chuk	nka								
Metallic phases	Sn–l	Pb–Cu	Nati	ve Sn°	Sn-	-Pb	Pb-Sr	I	Sn–Pb		Sn–Pl	b–Cu	Sn-Pt	,	soros	site	Sn _{syn}			Pb _{svn}			η-Cu	_{6.26} Sn ₅
Sample No.		5I.S48	H23			2.S10	_		OB-54-	4b	_		_		_		# 4-6			# 4-6			# 47-	
Analytical condition	is																							
Radiation	Cu I	Ka	Cu l	Ka	Cu	Ka	Co K	,	Fe K _{α,β}		Cu K	a	_		Fe K	a	Cu K	a		Cu K	a		Cu K	
Filter	Ni		Ni	u	Ni		-	~	-		_		_		_		Ni			Ni	u.		Ni	u
U (kV)	40		40		40		_		_		_		_		_		_			-			_	
I (mA)	19		19		19		_		_		_		_		_		_			-			_	
Exposition time (h)	25.0	5	35		40		5–6		_		-		-		_		_			-			-	
Camera	57.3 Gan	mm dolfi	57.3	mm Ga	ndolfi		57.3 n Debye Scher	÷-	57.3 mr Debye- Scherre		57.3 r Debye Scher	e-	-		114.6 Deby Sche	/e-	-			-			Diffra	actometer
Results	<i>I</i> *	d (Å)	Ι	d (Å)		d (Å) 2.95(s)	Ι	d (Å)		d (Å)		d (Å)	Ι	d (Å)		d (Å) 2.970	Ι	$d(\text{\AA})$	hkl	Ι	d (Å)	hkl	I 100	d (Å) hkl 2.9640 101
	6	2.92	10	2.92			10	2.89	2	2.914	3	2.902	10	2.920			100	2.915	200					
	9	2.85			10				3	2.857	6	2.849	10	2.860						100	2.855	111		
	9	2.78	10	2.77	10				4	2.789	3	2.780	10	2.792			90	2.793	101					
					1	2.62									1	2.560							5	2.5489 002
	5	2.44			8	2.48	4	2.49	4	2.476	5	2.452	7	2.484	1	2.560				50	2.475	200	3	2.5489 002
	3	2.44			0	2.40	4	2.49	4 .	2.470	3	2.432	/	2.464	8	2.112				50	2.473	200	82	2.1030 110
															9	2.094							64	2.0886 102
	3	2.06	5	2.05	4	2.06			2	2.060	2	2.058			,	2.094	34	2.062	220				04	2.0000 102
	8	2.00	10	2.00	10	2.00	2	2.02		2.017		2.010	9	2.000			74	2.002						
	7	1.746	10	2.00	5	1.746	3	1.75		1.752		1.746		1.752				2.017	211	31	1.750	220		
															3	1.720							18	1.7153 201
	3	1.658	4	1.652	3	1.663			2	1.661	3	1.661					17	1.659	301					
															2	1.627							8	1.6222 112
															2	1.546							10	1.5399 103
	10	1.492			8	1.488	5	1.49	10	1.492	10	1.490	10	1.493						32	1.493	311		
	10	1.484(s)	4	1.479(s	.)										5	1.487	23	1.484	112				20	1.4819 202
	1	1.457	2	1.450	1	1.454					1	1.451					13	1.458	400					
	2	1.441	4	1.434	3	1.442				1.442	1	1.443					20	1.442	321					
	1	1.427					2	1.43	2	1.426	1	1.431								9	1.429	222		
					1	1.357(s)			1	1.328					4	1.333							4	1.3291 211

rosite (Å) rosite (Å)															5.120 (· ·						74 (1)
, (Å) rosite (Å)	4.94	7 (5)			4.95	5	4.95		4.95		4.951		4.95		4.217 (4)			4.950)6	4 204	62 (1)
(Å) (Å)		3 (4) 3 (9)	5.82 3.17		5.84 3.19				5.84 3.20		5.831 3.182		5.83 3.18				5.831 3.182					
(Å)	5 07	3 (4)	5.07	,	5.84				5.84		5.831		5.92				5.831				5	0.0027 40.
	1	0.806	1	0.806												-	3	0.8058 701			3	0.8027 403
	1	0.809	2	0.808														0.8086 640			0	5.0111 22-
																					3 8	0.8193 21: 0.8111 22
	2	0.825(s)																	4	0.8251 600	4 3	0.8247 32
	-	0.000																		0.0051 -005	3	0.8274 10
	5	0.836																	9	0.8369 531		
	1	0.839	3	0.838												4	4	0.8386 631				
			5	0.846												1	10	0.8466 532				
	5	0.848														4	4	0.8485 413			-	
																					3	0.857640
	5	0.070(8)	5	0.075(8	·/											4	-	0.0755 541	1	0.0752 440	3	0.8684 31
	1 3	0.887(s) 0.876(s)		0.886(s 0.875(s													4 2	0.8868 323 0.8755 541	1	0.8752 440	1 6	0.8896 20 0.8791 30
	1	0.007()	2	0.006(0.0060.202			2	0.8965 40
	1	0.917	1	0.916												4	5	0.9178 611				
	1	0.922																0.9219 620				
	3	0.929	2	0.927	2	0.929										1	13	0.9286 512				
											2	0.931				3	3	0.9310 303				
	5	0.955				0.945					5	0.755							5	0.9520 511	8	0.9392 31
	3	0.972(8)			1	0.945					3	0.955	3	0.972			2	0.9718 000	5	0.9526 511	2	0.9721 22
	1 1	0.983(s) 0.972(s)	1	0.980(s	5) 1	0.985(s)		5	0.984			3	0.972			5 2	0.9824 213			2	0.9818 10 0.9721 22
	1	0.082(a)	1	0.980(s	A 1	0.095(2)	``		5	0.984							-	0.9824 213			4 2	0.9910 31
	3	1.011			1	1.010			6	1.011	1	1.011	10	1.010					6	1.0105 422		
	1	1.025	1	1.024	1	1.027			4	1.028		1.027				4	5	1.0252 521				
									1	1.034							2	1.0309 440				
	2	1.040	2	1.040	1	1.040										4	5	1.0401 332				
									4	1.043	1	1.044					3	1.0434 103			-	
							2	1.06	1	1.046						1.054					7	1.0515 22
															2	1.072					10 5	1.0898 11 1.0697 21
	3	1.096(s)	3	1.092(s	s) 2	1.095(s)		3	1.096	1	1.100	3	1.095	4	1.095	13	1.095 431			2	1.0961 30
	3	1.106			2	1.105			4	1.107		1.107		1.105					7	1.1069 420		
	3	1.135			2	1.132	1	1.13		1.137		1.137		1.134					10	1.1359 331		
	4	1.204(s)	8	1.201(s	5)				7	1.207	4	1.207	5	1.204	5	1.212	20	1.205 312			16	1.2112 21
					5	1.217										1.218					10	1.2142 30
	1	1.238(s)							2	1.237	1	1.238				1.248			2	1.236 400	4	1.2423 20
	2	1.293	3	1.289	2	1.296			4	1.295	1	1.296			2	1.279	15	1.292 411			3	1.2742 00
	2	1 202	3	1 200	2	1 200			4	1 205	1	1 200					15	1.292 411				

* Intensities estimated visually. (s): XRD peaks close to sorosite peaks

crospherules with Sn–Pb composition. Cosmic metallic droplets found in the deep-sea sediments consist mainly of Fe and Ni (Blanchard et al., 1980; Brownlee et al., 1997). Particles quite similar in composition (*table I*), shape and size have been found in impact materials of the El'gygytgyn meteoritic crater (Gurov and Kudinova, 1987). The origin of these microspherules has been attributed to the impact process: condensation of a high-temperature gas phase mobilised from the land target under reducing conditions realised during the impact. In the case of open ocean, the target for meteoritic falls could be the oceanic surface and similar processes of Sn–Pb mobilisation and condensation are hardly possible. That is why we believe that an extraterrestrial or impact origin of examined particles has also to be rejected.

– Probable hydrothermal or magmatic origin. We suggest the major factors controlling the Sn–Pb–Cu phase occurrences appear to be the hydrothermal activity and magmatic processes in the TAG rift valley. Most probably these metallic particles have been formed in the hydrothermal sediments themselves and/or have been presented as accessory minerals disseminated in the basic rocks exposed at the foot of high-elevated east rift wall. During bedrock degradation the metallic grains could have been liberated and, in turn, deposited into adjacent sediments.

· Formation related to hydrothermal processes. We consider that the high-temperature hydrothermal discharge at the TAG field have not played a role in the formation of the studied particles. The high activity of reduced sulfur in the TAG hydrothermal vents (Campbell et al., 1988) would provide a sink for chalcophiles (Pb, Cu, Sn) deposited as sulfides rather than their deposition in zero-valent state. Nevertheless, it is known that laboratory synthesis studies (Alminas et al., 1994) in which Pb- and Sn-rich saline solutions were introduced into a CaCO₃-rich environment at 25°C, 1 atm and low pH, produced native tin. Metallic particles very similar in composition, shape and size, and in the same mineral associations have been found in the soils and stream sediments of the US Virgin Islands (Alminas et al., 1994). Their origin was attributed to the interaction between metal-rich chloride hydrothermal solutions and calcareous oozes into which the solutions had been injected (Alminas et al., 1994). Seepage of hot hydrothermal fluids through the TAG sediment cover could cause boiling and phase separation (vapor and liquid) of the fluid as in the sediment hosted submarine hydrothermal environments. The low temperature, saline, metal-rich and exhausted in H_2S residual liquid would interact with the calcareous matrix of the TAG oozes and might have precipitated native Sn–Pb–Cu association within the sediment cover.

• Formation in the oceanic crust during the magmatic crystallisation. There are two possible pathways: (1) formation from an immiscible metal liquid during crystal-liquid fractionation processes in magmatic melt; and (2) reduction and crystallisation from a silicate melt matrix. We have already discussed these two models (Dekov et al., 1996) and would like to give attention to some features of the second mode of formation. Experimental studies on the interaction of hydrogen with basic magmatic melts (Persikov et al., 1986) revealed that during these processes Pb, Fe, Ni and Co had been separated as metallic microspherules and irregular-shaped particles in the silicate matrix. We consider that beneath the rift zones with high intratelluric H₂ flux similar processes of metallisation of the peridotitic upper mantle, which partially melts to produce MORB, could occur.

Sorosite-bearing assemblage of native tin and lead in the Baimka placer (Barkov et al., 1998) occurs in an environment, which contains widespread ophiolitic rocks. This implies that metallic Sn–Pb–Cu phases could derive from the mafic and serpentinised ultramafic lithologies of the ancient and recent oceanic crust.

4.2. Formation conditions and sequence of crystallisation

The phase diagrams of the binary systems Cu–Sn and Sn–Pb (after Hansen and Anderko, 1958; Champion et al., 1981) are given in *figure 4*. The Cu–Pb system is monotectic with a eutectic reaction near the high Pb corner. It is not discussed hereafter since the crystallisation pathway of the studied particles does not pass through the Cu–Pb field. The Pb–Sn system shows a simple eutectic, while the Cu–Sn one is more complicated with binary phases and peritectic transformations. The ternary Cu–Pb–Sn system has been studied by Marcotte and Schroder (1983). These authors established the reduction of the decomposition temperature of η -Cu₆Sn₅ with addition of Pb in the Sn–Cu system and a ternary eutectic reaction near the binary Pb–Sn eutectic.

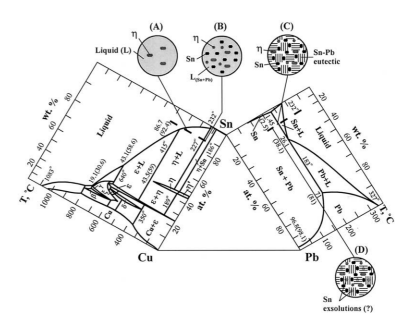


Figure 4. Diagrams of equilibrium of Cu–Sn, Sn–Pb and Cu–Pb binary systems (after Hansen and Anderko, 1958; Champion et al., 1981) and proposed phase variety and microtextural relationships at different solidification stages of the initial ternary Sn–Pb–Cu melt (see explanation in the text).

On the basis of the observed microtextural relationships, and phase compositions of the particles examined, the formation of the Sn–Pb–Cu particles from the TAG metalliferous sediments was probably realised with falling temperature in the following sequence (*figure 4*):

– Crystallisation of sorosite (η-Cu₆Sn₅) from the parent melt (L) at around 380°C (*figure 4A*). Metallic melt becomes practically exhausted in Cu. The temperature, experimentally found by Marcotte and Schroder (1983) as a η-Cu₆Sn₅ decomposition temperature of Sn₇₀Pb₂₀Cu₁₀ system, was 35°C lower than the one given in *figure 4* because of the Pb addition in the system. The single large ('porphyritic') η-Cu₆Sn₅ crystals on the background of the fine microcrystalline structure are probably due to gravity segregation.

– Formation of tin crystals from liquid with tentative composition $Sn_{80}Pb_{20}$ started at $T < 227^{\circ}C$ (*figure 4B*).

– Sn–Pb eutectic, composed of intimate intergrowths of Sn and (Pb,Sn), formed from the remainder liquid at T < 183°C and the whole of the melt transformed into solid state (*figure 4C*). It is well known that Pb at 183°C and under ambient pressure is able to insert up to 29% of Sn atoms in its crystal lattice without changing the crystal structure. This value at 0°C is respectively 3.2% of Sn atoms (*figure 4*). The lead–tin phase (Pb_{74.3}Sn_{25.7} in percent atoms) in the particles studied has Sn content

close to the uppermost limit, which may be explained by three different ways:

• very rapid melt cooling (quenching effect) to below the temperature of eutectic crystallisation of the Sn–Pb system (i.e., < 183°C); in this case, such a rapid cooling should prevent appearance of Sn exsolutions within the lead–tin inclusions and the system should remain under non-equilibrium state, in comparison with the low temperature of the parent rock;

• the compositions of the lead-tin inclusions (*table I*) have been obtained from non homogeneous sections including also Sn microexsolutions, this could lead to an overestimation of the actual Sn content during the microprobe investigations (analytical artefact); some non homogeneity of these inclusions can be observed in *figure 2C* and *figure 3C2*;

• the Sn–Pb phase diagram (*figure 4*), gives only a tentative notion of the probable development of the geological processes because it takes into consideration one basic physicochemical parameter, the temperature; the behaviour of a melt under high pressures of the magmatogenic mineral-forming processes is not known although the influence of low pressures on the phase equilibrium and crystallisation pathways of metallic systems in liquid and solid state is generally insignificant.

We suppose that in the present case some quantity of tin exsolutions has been separated from the lead-tin solid solution as the temperature falls to below the eutectic level. It is quite possible that this process had not been fully completed and at a certain moment sudden cooling provoked by various geological reasons (for example, opening of the system caused by tectonic movements, mixing with low-temperature igneous derivatives, etc.) took place. The eutectic microtexture and spherical shape of some of the Sn–Pb–Cu particles studied (*figure 2B,D*) support such a conclusion.

- Sn exsolutions were separated from the lead-tin phase (Pb_{74.3},Sn_{25.7}); shortly afterwards sudden cooling of the system stopped the exsolution process giving the association its end phase composition (*figure 4D*).

The influence of the cooling rate and related diffusion in the system on the microtextural and phase characteristics of the Cu–Sn association in solid state has been established by Champion et al. (1981). Using the explanations of these authors, we may conclude that, as a whole, the investigated Sn–Pb–Cu association might have been formed as a result of relatively rapid cooling of parent melt. This led to an appearance of numerous nuclei creating eutectic and dendrite microtextures of intimate intergrowths of the main solid phases (*figure 2C,D*). The presence of large 'porphyritic' inclusions of sorosite (η -Cu₆Sn₅), however, testifies to relatively low cooling rate during the initial stage of peritectic reaction (A) when gravity segregation takes place.

5. CONCLUSION

The examined metallic particles of the Sn–Pb–Cu system found in the sediments from the Mir zone, TAG hydrothermal field (Mid-Atlantic Ridge, 26°N) are thought to have a natural origin related to either: 1) the hydrothermal seeping processes in the sediment cover; or 2) formation of rift valley igneous rocks. On the one hand, metal-rich hydrothermal fluids could seep through the TAG sediment cover, react with the calcareous matrix and precipitate tin–lead based metallic particles. On the other hand, Sn–Pb–Cu metallic microsegregations could have been formed magmatically during the evolution of ridge crest magmatic systems: either from an immiscible metal liquid; or by reduction from the silicate melt. The pathway of formation of the Sn–Pb–Cu association and crystallisation kinetics of the process have probably been as follows: relatively slow crystallisation and gravity segregation of sorosite (η -Cu₆Sn₅) at the initial stage of solidification under moderate fall in temperature below $380^{\circ}C \rightarrow$ rapid crystallisation at the middle and final stage of η -Cu₆Sn₅ formation as well as during the Sn–Pb eutectic reaction creating a mixture of tin and stannoan lead (total range of $300^{\circ}?$ ~183°C) \rightarrow rapid cooling at T << 183°C (?) stopping the process of tin exsolution.

Magmatic intrusions in the ridge axial zone might have transported the Sn–Pb–Cu segregations within the silicate matrix to the upper earth's crust levels. During the processes of rift valley formation (Lisitsyn, 1992) the gabbro and sheet–dike layers have been exposed and have undergone active disintegration. The native Sn–Pb–Cu grains liberated together with other rock shards have been dispersed into the rift valley sediments.

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