

PIN WITHOUT HEAD HR-18152 - TIN BRONZE - LATE BRONZE AGE - SWITZERLAND

Artefact name

Pin without head HR-18152

Authors

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Url

/artefacts/376/

➢ The object

Fig. 1: Tin bronze pin without head (after Rychner-Faraggi 1993, plate 74.29),



✤ Description and visual observation

Description of the artefact	Pin without head, light-brown patina typical of lake context (Fig. 1). Dimensions: L = 7,5cm; Ø = 3.3mm; WT = 4g.					
Type of artefact	Pin					
Origin	Hauterive - Champréveyres, Neuchâtel, Neuchâtel, Switzerland					
Recovering date	Excavation 1983-1985, object from layer 3 to 5					
Chronology category	Late Bronze Age					
chronology tpq	1054 B.C. 🗸					
chronology taq	1000 B.C. 🗸					
Chronology comment	Hallstatt B1 (1054/1037BC _ 1000BC)					
Burial conditions / environment	Lake					

Artefact location	Laténium, Neuchâtel, Neuchâtel							
Owner	Laténium, Neuchâtel, Neuchâtel							
Inv. number	Hr 18152							
Recorded conservation data	Not conserved							
Complementary information								
Nothing to report.								
	Fig. 2: Location of sampling area,							
➢ Binocular observation and repres	sentation of the corrosion structure							
Stratigraphic representation: none.								
✓ MiCorr stratigraphy(ies) – Bi								
✓ Sample(s)								
	Fig. 3: Micrograph of the cross-section showing the location of Figs. 5 to 12,							

Fig. 5, 6 Fig. 8, 9 Fig. 7 Fig. 7 Fig. 10-13	
	1mm

Credit HE-Arc CR.

Description of sample	The cross-section is circular and is a complete section through the pin (Fig. 2). It is covered with a rather thin and regular (in thickness) corrosion crust. One third of the corrosion crust is missing (Fig. 3).					
Alloy	Tin Bronze					
Technology	Annealed after cold working					
Lab number of sample	MAH 87-194					
Sample location	Musées d'art et d'histoire, Genève, Geneva					
Responsible institution	Musées d'art et d'histoire, Genève, Geneva					
Date and aim of sampling	1987, metallography and corrosion characterisation					

Complementary information

Nothing to report.

✓ Analyses and results

Analyses performed: Metallography (etched with ferric chloride reagent), Vickers hardness testing, XRF, ICP-OES, SEM/EDS, XRD, Raman spectroscopy.

➢ Non invasive analysis

✓ Metal

The remaining metal is a tin bronze and contains copper sulphide as well as heavy metal (Pb-rich) inclusions (Table 1, Figs. 5 and 6). Close to the surface of the remaining metal, copper sulphide inclusions are elongated and form rows (Fig. 5). The etched structure of the tin bronze shows polygonal grains; some of them are twinned (Fig. 7). In the centre of the sample and on the edges, the grains are smaller. The copper sulphide inclusions are located at the grain boundaries and in the grains. The average hardness of the metal is about HV1 110.

Elements	Cu	Sn	Pb	Sb	As	Ag	Fe	Ni	Со	Zn
mass%	89.22	9.57	0.34	0.26	0.19	0.15	0.09	0.05	0.06	0.05

Table 1: Chemical composition of the metal. Method of analysis: ICP-OES, Laboratory of Analytical Chemistry, Empa.



Credit HE-Arc CR..



Credit HE-Arc CR.



Credit HE-Arc CR.

Fig. 5: Micrograph of the metal sample from Fig. 3 (detail), unetched, bright field. Rows of elongated copper sulphide inclusions can be observed,

Fig. 6: SEM image of the metal sample from Fig. 3 (detail), SE-mode. Elongated copper sulphide inclusions (dark-grey) are visible as well as heavy metal (Pb-rich) inclusions (white),

Fig. 7: Micrograph of the metal sample from Fig. 3 (detail), etched, bright field. The metal shows a structure of polygonal and twinned grains. Copper sulphide inclusions are visible as blue spots,

Microstructure	Polygonal and twinned grains
First metal element	Cu
Other metal elements	Sn
Complementary information	
✓ Corrosion lavers	

The corrosion crust is regular in thickness (100µm). One third is missing (Fig. 3). At the metal - corrosion crust interface, there is a crack showing that the latter has separated from the metal core along its whole length (Figs. 8 and 9). The corrosion crust can be divided into three distinct layers (CP1-3). Directly above the crack is a first dense but cracked and irregular inner layer (CP3, Figs. 8 and 9). In bright field it appears brown (Fig. 10), in polarised light dark brown (Fig. 11). It is separated from the adjacent layer by a clear line (Figs. 9 and 10). The second layer (CP2) is dense with little porosity (Figs. 8 and 9). In bright field it appears light brown (fig. 10), in polarised dark grey (Fig. 11). The third and outermost layer (CP1) appears light brown in bright field (Fig. 10), contains particles (Fig. 9) and is very porous (visible as golden reflections under polarized light, Fig. 11). The elemental chemical distribution of the SEM image selected reveals that the inner layer (CP3) is depleted in Cu, but rich in Sn,O and Si (Fig. 12 and Table 2) and its interface with the intermediate layer (CP2) could represent the limit of the original surface (Figs. 8 and 12). The second and third layer (CP2 and CP1) are Fe,Cu and S-rich (Fig. 12) and have a composition similar to chalcopyrite/CuFeS₂ (Table 2). This was confirmed by XRD. The particles (inclusions) have a composition similar to covelline or covellite/CuS (Table 2). Both chalcopyrite and covelline have been identified by Raman spectroscopy (Figs. 13 and 14).

Elements		Fe	Cu	0	Si	Sn	Total
CP1 and CP2	35	30	34	<	<	<	99
Particles in CP1	26	4.1	68	<	<	<	98
CP3	5.8	5.0	13	32	2	41	99

Table 2: Chemical composition (mass %) of the corrosion layer from Fig. 10. Method of analysis: SEM/EDS, Laboratory of Analytical Chemistry, Empa.



Credit HE-Arc CR.

Fig. 8: SEM image of the metal sample from Fig. 3 (detail), SE -mode. The crack between the remaining metal and the corrosion layers is clearly visible,

Fig. 9: SEM image similar to Fig. 8, BSE-mode. A thin line is visible (white arrow) differentiating the inner Sn-rich layer from the intermediate and outer chalcopyrite layers. The latter contains bright covelline inclusions (black arrow) and pores,



Credit HE-Arc CR.



Fig. 10: Micrograph of the metal sample from Fig. 3 (detail) and corresponding to the stratigraphy of Fig. 4, unetched, bright field,

Credit HE-Arc CR.



Credit HE-Arc CR.

Fig. 11: Micrograph similar to Fig. 10, polarised light,

Fig. 12: SEM image and elemental chemical distribution of a selected area of Fig. 10 (rotated by 90°). Method of examination: SEM/EDS, Laboratory of Analytical Chemistry, Empa,



70µm Si P Credit Empa.



Credit SNM.



Credit SNM.

Corrosion form

Corrosion type

Uniform - pitting

Type I (Robbiola)

Complementary information

Nothing to report.

➢ MiCorr stratigraphy(ies) − CS

Fig. 13: Raman spectrum of the outermost layer (S23) of Fig. 9 compared to a chalcopyrite standard spectrum. Settings: laser wavelength 532nm, acquisition time 50s, 4 accumulations, filter D2 (0.75-0.8mW), hole 1000, slit 100, grating 600. Method of analysis: Raman spectroscopy, Lab of Swiss National Museum, Affoltern a. Albis ZH,

Fig. 14: Raman spectrum of the inclusions of the outermost layer (S42) of Fig. 9 compared to a covelline / covellite standard spectrum. Settings: laser wavelength 532nm, acquisition time 10s, 5 accumulations, D2 (0.75-0.8mW), hole 500, slit 80, grating 600. Method of analysis: Raman spectroscopy, Lab of Swiss National Museum, Affoltern a. Albis ZH,



Fig. 4: Stratigraphic representation of the object in cross-section using the MiCorr application. This representation can be compared to Fig. 10, Credit HE-Arc CR.

✤ Synthesis of the binocular / cross-section examination of the corrosion structure

Corrected stratigraphic representation: none.

imes Conclusion

The pin is made from tin bronze and has been annealed after cold working. It is covered with a regular, light-brown patina typical of lake context (Schweizer 1994). The inner, thin Sn-rich corrosion layer contains soil elements such as Si. The light-brown, thick intermediate and outer corrosion layers have the composition of chalcopyrite. This object was certainly abandoned rather quickly in an anaerobic, humid and S and Fe-rich environment, favouring the formation of the above mentioned compound. The limit of the original surface can be located between the chalcopyrite and the Cu depleted but Sn-rich inner corrosion layer. Thus, the corrosion is a type 1 according to Robbiola et al. 1998.

References on object and sample

References object

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Hochuli, S. et al. (1988) SPM III Bronzezeit, Verlag Schweizerische Gesellschaft für Ur- und Frühgschichte Basel, 76-77, 379.

References sample

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References on analytic methods and interpretation

6. Robbiola, L., Blengino, J-M., Fiaud, C. (1998) Morphology and mechanisms of formation of natural patinas on archaeological Cu-Sn alloys, Corrosion Science, 40, 12, 2083-2111.