



Alloy characterization of a 7th Century BC archeological bronze vase — Overcoming patina constraints using Monte Carlo simulations[☆]



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ABSTRACT

In this work we evaluate the composition of a bronze alloy using X-ray fluorescence spectrometry (XRF) and Monte Carlo (MC) simulations. For this purpose, a 7th Century BC archeological vase from the SW Iberian Peninsula, displaying a well formed corrosion patina was analyzed by means of a portable X-ray fluorescence spectrometer. Realistic MC simulations of the experimental setup were performed with the XRMC code package which is based on an intensive use of variance-reduction techniques and uses XRAYLIB a constantly updated X-ray library of atomic data. A single layer model was applied for simulating XRF of polished/pristine bronze whereas a two-or-three-layer model was developed for bronze covered respectively by a corrosion patina alone or coupled with a superficial soil derived crust. These simulations took into account corrosion (cerussite (PbCO₃), cuprite (Cu₂O), malachite (Cu₂CO₃(OH)₂), litharge (PbO)) and soil derived products (goethite (FeO(OH)) and quartz (SiO₂)) identified by means of X-ray diffraction and Raman micro analytical techniques. Results confirm previous research indicating that the XRF/Monte Carlo protocol is well suited when a two-layered model is considered, whereas in areas where the patina + soil derived products' crust is too thick, X-rays from the alloy substrate are not able to exit the sample. Quantitative results based on MC simulations indicate that the vase is made of a lead-bronze alloy: Mn (0.2%), Fe (1.0%), Cu (81.8%), As (0.5%), Ag (0.6%), Sn (8.0%) and Pb (8.0%).

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1. Introduction

The identification of ancient technologies of production from the determination of chemical composition and morphology of archeological objects often allows a clear attribution to a specific geographical region, as well as, an elucidation of the date of its manufacture, leading to a better understanding of ancient cultures. Furthermore, chemical characterization of archeological objects may also contribute to a better evaluation of their surface deterioration and to determine the authenticity of the artifacts [1].

X-ray fluorescence spectrometry (XRF) is a simple and fast technique that allows elemental determination and quantification ($Z > 12$) of metal alloys in a non-destructive way. This technique has been widely

used providing new insights concerning the evolution of the use of metals [2–5] and their provenance [6,7] through the identification of the component materials.

Advances in technology contributed to the miniaturization of X-ray detectors and tubes enabling the implementation of portable and handheld spectrometers able to provide in-situ measurements [8]. Realistic Monte Carlo (MC) simulations of experimental setups have been developed, in recent years, as a quantification tool able to take into account the roughness of the surface [7,9].

Alloyed objects suffer from atmospheric corrosion which leads to the formation of various chemical compounds such as oxides, carbonates, sulfides, or sulfates on the metal surface. This corrosion layer is known as a patina. The presence of patinas in alloyed objects is often a constraint for its accurate characterization, as the analysis by XRF determines the composition of this superficial corrosion layer formed on the object's surface and not the composition of the underlying material. For an accurate chemical characterization the patina must be removed, which is frequently not allowed. The difficulties in analyzing

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archeological bronzes because of the alloy's heterogeneity and surface patina, as well as the possibilities and limits of this analytical method have been emphasized in the literature as a topical issue in archaeometry and conservation science [6,10].

In this work we combine XRF spectrometry and MC simulations to evaluate the composition of a bronze alloy with no need for patina removal. An archeological bronze vase from a site near an important 7th Century BC Phoenician settlement (Alcácer do Sal) in the SW part of the Iberian Peninsula was analyzed using a portable XRF spectrometer. MC simulations of the experimental setup were performed with the XRMC code package [11,12].

2. Materials and methods

The archeological bronze vase selected for the analytical protocol was found in the 7th Century BC necropolis of Senhor dos Mártires near Alcácer do Sal in Southern Portugal, a well known Phoenician site in the protohistory of the Iberian Peninsula [13,14]. It was casted using the lost-wax technique and has been interpreted as having a ritual funerary use related to libation and purification practices.

Determination of corrosion and soil derived products was carried out by X-ray diffraction (XRD). Raman microscopy was used to study these alterations when they were found to be amorphous [15]. For this purpose samples were collected from the surface, and for the alloy determination, XRF analyses were carried out on the vase.

2.1. XRD

The mineralogical composition of the bronze corrosion patina was determined by XRD, using a commercial D2-Phaser, Bruker/AXS diffractometer with a Cu K α source operating at 30 kV and 10 mA and a lynxeye detector. Powdered samples scraped from the vase were deposited onto a flat glassy support and irradiated through a 0.6 mm slit. The angular range (2θ) was scanned from 6° to 80° at a step size of 0.02° with a counting time of 0.5 s/step. Evaluation of X-ray diffractograms was made by using the routines of the Diffrac software package (BRUKER/AXS GmbH, Germany) and the attached specific PDF database files.

2.2. Raman

Raman analyses were undertaken using a Horiba-Jobin Yvon XploRA confocal spectrometer, operating at wavelength of 785 nm, with maximum incident power of 0.2 mW. Using a 100 magnification objective with a pinhole of 500 μ m and an entrance slit of 100 μ m, the scattered light collected by the objective was dispersed onto the air-cooled CCD array of an Andor iDus detector by a 1200 lines/mm grating. Raman microscopy performed in a range of 100–1200 cm^{-1} . Spectra deconvolution was performed using the LabSpec software (V5.78). The identification of corrosion and soil derived products was made in good agreement with literature [16] and Spectral ID™.

2.3. EDXRF and MC simulations

The portable spectrometer consists of an Amptek Mini X X-ray tube equipped with a silver anode (50 kV, 80 μ A) and an Amptek X-123SDD detector with a resolution of 140 eV FWHM at Mn K α (5.9 keV). Due to the morphology of the samples, the detector was placed orthogonally to the sample surface while the X-ray tube was at 20° with respect to the normal at the surface. The setup was calibrated using a certificated bronze sample [17]. All measurements were carried out under ambient air.

An innovative reverse MC simulation quantification approach was used. In this approach XRF spectra were simulated using XRMC code package, which is a fast X-ray interaction simulation code. The code uses a set of files describing both the experimental setup and the composition and structure of the sample. The initial estimate is based on the expected composition of the sample. In the case of the bronze

layer only its composition is required whereas in the presence of a patina layer both composition and thickness are needed. Both compositions as well as the patina thickness are adjusted until the best fit is obtained for the peak part and background. MC simulations took into account the patina and soil derived products composition.

The XRMC code is based on an intensive use of variance-reduction techniques and uses XRAYLIB a constantly updated X-ray library of atomic data [18]. Besides the speed it has, to the best of our knowledge, another unique capability: simulation of arbitrarily rough surfaces [19]. Currently the fitting is user driven but an automatic procedure is being developed, despite the fact that, due to the large number of parameters involved, a full automatization will be hard to achieve.

The surface of the artifact shows strong effects of corrosion as well as the presence of a crust layer of soil derived products. The first step was to observe at the microscope the state of the surface in order to choose the optimal experimental setup. A model of the roughness was implemented in the MC code and the optimal X-ray beam size was defined to be 3 mm wide. The detector and the X-ray tube were modeled following the same geometry of the experimental layout. The distance of X-ray tube from the surface of the object was about 2 cm, while the detector was placed 3 cm from the surface. The X-ray tube worked at 40 kV and the current was chosen according to the flux reaching the detector. Several points have been analyzed. Here we report the results obtained taking the X-ray spectra at patinated points as well as at cleaned ones. The goal is to demonstrate that the MC simulation is able to give a correct estimate of the bronze composition even in the presence of a corrosion + crust layers. The geometry adopted for the structure of the artifacts depends on the surface state: a single layer geometry for the cleaned surface, a two-layer geometry for surfaces with patina (corrosion) and finally a three-layer geometry when a crust layer (soil derived products) is also observed. The layers' thicknesses as well as the elemental weight percentage have been changed in the simulations to fit the measured spectra.

3. Results and discussion

XRD analysis revealed that the main crystalline phases of the patina are cerussite (PbCO_3), cuprite (Cu_2O), malachite ($\text{Cu}_2\text{CO}_3(\text{OH})_2$) and quartz (SiO_2). Additional peaks can be attributed to the bronze support (Fig. 1). Raman analysis confirmed the presence of cerussite through the identification of the 1055 cm^{-1} band. The large band spreading from 490 to 660 cm^{-1} with two maxima located at 520 and 620 cm^{-1} has been reported as a characteristic of defective cuprite [20]. The band at 143 cm^{-1} is ascribed to litharge (PbO) (Fig. 2). Furthermore, goethite was also identified through the presence of the characteristic bands at

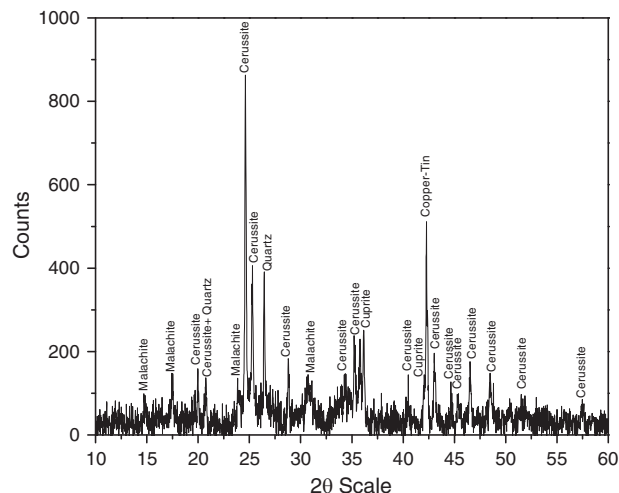


Fig. 1. XRD diffractogram of patina and soil derived products identified on the vase.

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