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Defining new technological traditions of Late Islamic Arabia: a view on Bahlā Ware from al-Ain (UAE) and the lead-barium glaze production

Jelena Živković 1 o · Timothy Power 2 · Myrto Georgakopoulou 1 · José Cristobal Carvajal López 3

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Abstract

In this paper, the monochrome glazed Bahlā Ware from al-Ain dated between the seventeenth and twentieth centuries (Late Islamic Arabian Period) has been analysed aiming to reconstruct the production technology of the ceramic fabrics and glazes. The results of the petrographic and chemical analyses suggest a unique technological tradition embedded in the culture of Late Islamic Arabia. This tradition incorporates the production of a lead-barium glaze coated over a single type of ceramic fabric that spans for nearly three centuries. Since this is the first evidence for the production of a lead-barium glaze in the Islamic World, the origins of this technology remain uncertain, but the results of the ceramic petrography identify the Omani Peninsula as the most likely source for the ceramic fabric. During the economic peak of al-Ain in the eighteenth century, this tradition shows signs of technological diversity visible in the appearance of new fabrics and glazes. Considering the wide distribution of Bahlā Ware in the Western Indian Ocean, understanding of the production technology and provenance of al-Ain's ceramics has important implications for archaeological interpretation.

Keywords Bahlā Ware · Lead-barium glaze · Ceramic technology · Islamic ceramics · South East Arabia

Introduction

Scientific data presented in this paper are obtained in a pilot project designed to set the stage for the extensive research on ceramic production and technology in Late Islamic Arabia. The material under study is the monochrome class of Bahlā Ware, found in consumption contexts at al-Ain dated between the middle of the seventeenth- and the early twentieth centuries. The focus here is particularly on the lead-barium glaze that has been identified for the first time in the context of Islamic ceramics. Considering the lack of production debris

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- UCL Qatar, Georgetown Building 2nd floor, P.O. Box 25256, Doha, Qatar
- College of Humanities and Social Sciences, Zayed University, P.O. Box 144534, Abu Dhabi, UAE
- University of Leicester, University Road, Leicester LE1 7RH, UK

that can be linked with this ceramic class, the reconstruction of production practices starts with an in-depth analysis of consumed pottery.

Bahlā Ware, the Oases of al-Ain and the regional context

Bahlā Ware constitutes a class of monochrome glazed ceramics dominated by open bowl forms. It was first identified by Andrew Williamson as part of his survey of the Mīnāb Plain, the land behind the great Late Islamic Iranian emporium of Hormuz, published posthumously by Seth Priestman (2005), pp. 269–70, 2013, pp. 631–32). Williamson thought it was produced in the town of Khunj, in the hinterland of Bandar Lengeh, in southern Iran (Fig. 1). However, Priestman notes that no wasters were found among the sherds collected by Williamson, implying that Khunj is unlikely to have been the production centre. Conversely, this class is widely found in South East Arabia and it has alternatively been suggested that it was a product of the well-known kilns of Bahlā, a large oasis town in central Oman, which remained in use until the 1970s (Whitcomb 1975, p. 129; Priestman 2008, pp. 277–78, Plate 12). This identification seems



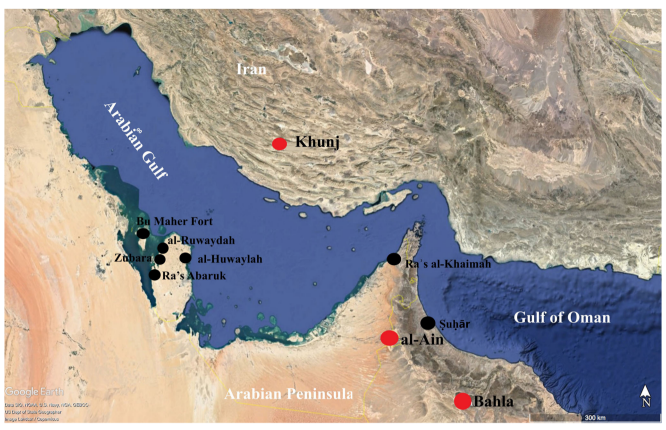


Fig. 1 The map showing locations of major sites of al-Ain, Bahlā, Khunj and other archaeological sites discussed in the paper

supported by a quantified study of the Late Islamic assemblage in the oases of al-Ain, further north along the same plain wherein Bahlā is situated, where this class represents as much as 48% of the glazed assemblage (Power 2015, p. 29, Table 26).

If the production centre in South East Arabia can be established, Bahlā Ware becomes a key marker of Omani trade in the early modern period. It is widely found at Late Islamic sites in the Arabian Gulf and western Indian Ocean between the sixteenth and twentieth centuries. Well-published instances in the UAE and Oman include Ra s al-Khaimah (de Cardi et al. 1994, p. 63; Kennet 2004, pp. 54-55), al-Ain (Power and Sheehan 2011, 2012; Power 2015), and Şuḥār (Costa and Wilkinson 1987). It is commonly found in Qatar, at Zubara and al-Ruwaydah (Petersen et al. 2010, p. 48), Ra's Abaruk (Garlake 1978a, p. 167) and al-Huwaylah (Garlake 1978b, p. 174), and also in Bahrain, at the Bu Maher Fort (Kennet 2004, p. 54; Carter et al. 2011, pp. 90-91). Importantly, it has also been found at sites in East Africa, including Gedi (de Cardi and Doe 1971, pp. 266-67) and Fort Jesus (Kirkman 1974).

The present study focuses on a sample of Bahlā Ware from excavations at the Bin Ātī House in Qaṭṭāra Oasis, one of seven palm plantations which make up the historic Buraimi Oasis (Power and Sheehan 2012), today split by an international border into the towns of al-Ain (UAE) and Buraimi

(Oman). The oases of al-Ain occupy a strategic crossroads in the northern Omani Peninsula. They lie on a north-south axis between Ra s al-Khaimah and the Iranian Plateau beyond, on the one hand, and the Dhāhira Plain extending south to central Oman and the great oasis town of Bahlā on the other. To the east, the Wādī al-Jizī cuts through the Ḥajar Mountains to gain access to the Indian Ocean networks via the port of Ṣuḥār, whilst the road west crosses the tip of the Rub al-Khālī to reach the pearling towns of Abu Dhabi and Dubai.

Excavations at the Bin Ātī House revealed three archaeological horizons (HRZ) associated with the Late Islamic occupation (Table 1). HRZ 9.1 (c.1650–1720) witnessed the construction of a large mudbrick tower house, probably contemporary with the creation of the adjacent palm plantation (Power and Sheehan 2012, p. 299, Fig. 6), which further included a date-press (madbasa). HRZ 9.2 (c. 1720-1790) marks an expansion of settlement and construction of a second house near the original tower; the quantity and quality of ceramic and coin finds suggest that this phase represents the peak of occupation. HRZ 10 (c. 1870-1920) constitutes a partial and perhaps seasonal reoccupation following a period of abandonment. Whilst the presence of other ceramic imports fluctuated over time, sometimes quite dramatically, it is striking that Bahlā Ware remains a consistent and slowly growing component of the assemblage.



Table 1 The stratigraphic and chronological context of the Bahla samples from al-Ain (see Power 2015)

Code	Archaeological horizon	Chronology	Number of selected samples
B1-22	HRZ 9.1	c. 1650–1720	10
B23-272	HRZ 9.2	c. 1720–1790	28
B273-288	HRZ 10	c. 1870–1920	6

Materials and methodology

Sampling

The sampling strategy followed in this paper has been designed to explore long-term patterns in the technology. For this purpose, 44 samples, out of 288 Bahlā Ware potsherds documented at Bin 'Ātī House, were selected for petrographic examination (Table 1). Drawing on the results of ceramic petrography, a number of sub-samples were subjected to chemical analyses both of the ceramic fabrics and the glazes as well as lead isotope analysis of the glazes. The aim was to investigate the compositional patterns, detecting potential variations between the glazes applied over fabrics with the same or different petrographic characteristics.

Samples belong to open bowls (40) and jars (4), reflecting the relative abundance of these types in the original assemblage. All vessels are wheel-made and coated with glaze on the external (jars) or both (bowls) sides. The colour of the glazes ranges between brown, yellow and green—all three were included in the sampling (Fig. 2).

Thin section ceramic petrography

Ceramic petrography was used for the petrological and textural characterisation of 44 samples. Thin sections of ceramics were observed with polarising microscopes LEICA DM750P and LEICA DM2500P in transmitted plain polarised (PPL) and cross polarised light (XPL). Fabric groups were defined by the petrology of their main inclusions, their distribution and texture (Whitbread 1989, 1995, pp. 365–96).

Elemental analysis of fabrics and analysis of glazes

Further research on the relations between fabrics was carried out by means of wavelength-dispersive X-ray fluorescence analysis (WDXRF), on a BRUKER S8 TIGER with a 4-kW Rh X-ray tube at the Fitch Laboratory of the British School at Athens. A calibration dedicated to the analysis of ancient ceramics was used for quantification, measuring in total 26 elements (Georgakopoulou et al. 2017). In total, 16 samples of the Bahlā Ware were analysed, all prepared as glass beads.

For the glaze analysis of 17 representative samples, a scanning electron microscope (SEM: JEOL JSM 6610LV) with an attached energy dispersive spectrometer (EDS: Oxford Instruments X-max^N 50 operated with the Aztec software) was used. Cross-sections of the samples were mounted in resin and polished to ¼ µm. The Bahlā glazes are often heterogeneous, and thus the analysis included bulk measurements and separate measurements of the glassy matrix as well as individual inclusions. The bulk composition measurements included the glaze with randomly spread inclusions, excluding bubbles and pores to the extent that this was possible. The reported results represent the average values for 5 areas, standardised at approximately 100 × 120 μ m using \times 800 magnification. For comparison, the analysis of the glaze matrix, avoiding the inclusions, is included. An average of 3-5 scans of an arbitrary size is reported, depending on the glaze texture. Furthermore, the composition of the various inclusions was measured with EDS spot analysis. The analyses were run in high vacuum conditions, at an accelerated voltage of 20 kV, working distance 10 mm, process time 5, and acquisition time 60 s livetime. A cobalt standard was measured periodically to monitor the beam current and the spot size was adjusted around 59 to achieve 40% deadtime on the cobalt metal. The performance of the instrument was monitored using the Corning Glass Standard C (Brill 1999, p. 542). Precision, estimated as relative standard deviation, was found to be within 3% for major elements and deteriorated as concentration approaches the detection limits of the EDS at around 0.1%. The relative difference of the mean to the certified value ($\delta\%$) is within 5% for most elements, with the exception of barium and lead where it is within 10% and cobalt whose composition approached the detection limits of the instrument and the difference is 13%.

Lead isotope analysis of glazes

Lead isotope analysis was performed on 5 samples of glazes to investigate the provenance of the lead-rich component. For this purpose, a small amount of glaze (c. 150 mg) was scraped off as a powder. The isotopic measurements were carried out at Frankfurt University, under the supervision of Dr. S. Klein, using a multicollector-inductively coupled plasma-mass spectrometer (MC-ICP-MS) Neptun TM Finnigan MAT (for details of the methodology see Klein et al. 2009).



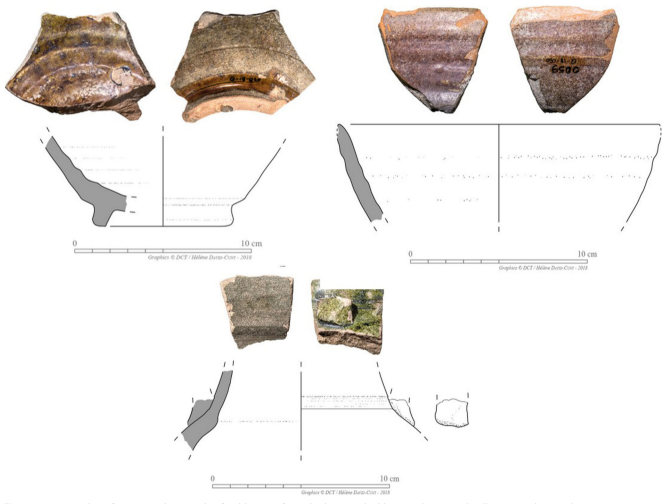


Fig. 2 Reconstruction of representative vessels of Bahla Ware from al-Ain, coated with green, brown and yellow monochrome glazes

Results

The results of ceramic petrography

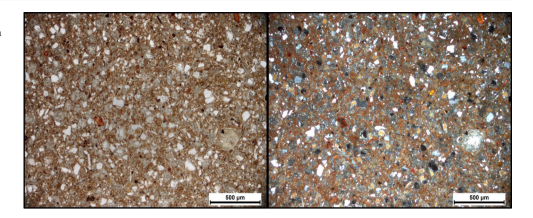
The results of ceramic petrography show that Bahlā Ware from al-Ain can be classified into a single fabric group, with two related samples and two petrographic loners (Appendix A). The ceramics are fine, with more than 90% of the inclusions occurring in the fine fraction.

Out of 44 analysed samples, 40 can be classified into the Limestone and Serpentinite fabric group (LS), defined by the presence of these two rocks set in a calcareous matrix (Appendix A; Fig. 3). The fabric is also characterised by detritic minerals consistent with an igneous geology of mafic or ultramafic rocks, such as clinopyroxene, plagioclase and olivine. This indicates that the raw clay was collected from a sedimentary environment where those minerals are deposited. The internal variability in the abundance of limestone, serpentinite and minerals associated with igneous rocks makes this fabric to some extent heterogenous.

The remaining four samples are petrographically diverse (Appendix A). All four samples have a bimodal distribution of inclusions, which indicates a different preparation of paste recipes compared to the LS fabric group. Apart from this common technological trait, the samples show a range of petrographic differences. Samples B132 and B252 contain rare inclusions of igneous rock/s, serpentinized in B252 (Fig. 4(a, b). Although these two samples have common inclusions in the coarse fraction, which makes them to some extent related, their relative quantity varies, making the degree of their relation uncertain. They also show a petrological association with the LS fabric group, considering the igneous nature of the inclusions in coarse fraction, but the different distribution of those inclusions (unimodal vs bimodal) is evident. B60 and B106 are described as petrographic loners because they contain even less inclusions that can indicate petrographic relations between them and the rest of the assemblage. The petrology of B106 is consistent with the LS fabric group, but the scarcity of inclusions in the coarse fraction prevent further



Fig. 3 Limestone Serpentinite fabric group (LS). Sample B32 in PPL (left) and XPL (right) with inclusions of quartz, limestone and serpentinite set in a calcareous matrix



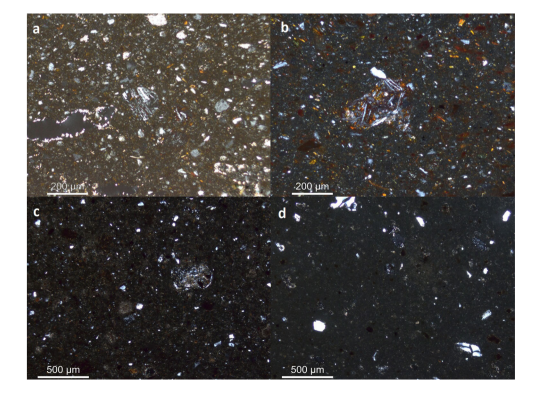
interpretation (Fig. 4(c). The same applies to B60, which is poor in inclusions in both the coarse and the fine fractions (Fig. 4(d).

Overall, the petrology of all four samples is in the same geological range with the LS fabric group, indicating exploitation of similar secondary clays. All analysed samples show well-controlled firing conditions as no colour differences can be observed in individual sherds. The equivalent firing temperature, based on the optical activity of the clay matrix, can be estimated to c. $800-850\,^{\circ}\text{C}$.

The chemical composition of ceramics

The results of WDXRF analysis are given in Table 2. Prior to pulverising the samples, the glaze layer was removed using a slow-speed saw with a thin diamond-coated blade and the external surfaces of the sherds were subsequently further cleaned with a tungsten-carbide drill. Still, the presence of significant contents of lead (Table 2) in the results suggests that part of the glaze layer was present in the pulverised sample. This will have affected a number of other elements in unpredictable ways as the glaze contamination is likely to be variable and the glaze compositions were also shown to be relatively heterogeneous. These elements are Ba, as will be shown below the glazes are Pb/Ba rich, as well as Cu present as a colourant in some cases. These three elements (Pb, Ba and Cu) are thus disregarded in the elemental grouping of ceramics. FeO is also present as a colourant in the glaze, but as this is a major element in the ceramic body, it is assumed that the traces of glaze contamination would not affect the overall iron content very much. The Th L α measured by the WDXRF partially overlaps the Pb L\u03b3. Although this is taken into

Fig. 4 Samples B132 (a), B252 (b), B60 (c) and B106 (d) that differ from the LS fabric group. Inclusions of igneous origin are visible on images a and b whilst images c and d show only inclusions of fine quartz. All images are given in XPL





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standard deviation of the samples belonging to the one chaste analysis are given for comparison. Les stands for the samples belonging to the one chaste analysis and interpretation of data as they are affected by the presence of lead-rich glaze, excluded because its relation to the burial contamination (see text for discussion)	excluded because its relation to the burial contamination (see text for discussion					2000	T IOL UE		6																	
	Na ₂ O) MgO	Al_2O_3	SiO_2	P ₂ O ₅) ₅ K ₂ O) CaO) TiO ₂	2 Fe ₂ O ₃)3 V	Cr	Mn	Co	Ni	Cu	Zn	Rb	Sr	7	Zr	Ba	La	Ce	pN	Pb	Th
rs	1.2	9:9	12.5	57.9	0.2	2.3	8.7	0.7	8.9	104	999 1	595	37	405	47	75	29	320	22	163	331	22	46	29	2322	18
rs	1.3	6.5	12.6	58.2	0.2	2.3	8.5	0.7	6.9	100	199 (577	41	406	43	73	69	327	21	163	320	23	53	16	2163	17
rs	1.4	6.5	12.8	59.2	0.2	2.1	8.6	0.8	6.5	113	644	473	33	327	55	75	51	224	24	168	311	24	50	25	4053	31
FS	1.1	0.9	11.4	51.9	0.2	2.3	13.4	1 0.7	0.9	66	503	477	31	300	65	74	64	588	20	44	1795	27	71	22	1895	15
rs	1.2	9.9	11.2	52.5	0.1	2.0	15.3	3 0.7	5.5	100	523	481	27	284	. 35	89	19	331	24	142	324	22	4	26	7420	57
rs	1.4	0.9	11.8	56.7	0.3	2.3	9.1	0.7	5.9	76	674	453	30	299	48	71	54	328	21	160	289	24	45	23	3231	25
rs	1.4	6.4	12.4	55.8	0.2	2.4	10.4	1 0.7	6.3	111	529	430	32	332	57	74	69	405	23	158	380	27	53	26	1452	12
rs	1.5	6.5	12.6	58.3	0.2	2.3	8.3	0.7	8.9	121	627	436	34	356	52	73	48	288	24	158	377	23	52	30	6210	48
rs	1.1	7.1	10.4	49.0	0.1	2.0	16.3	9.0 8	5.2	68	260	456	28	272	81	65	09	561	19	136	349	21	4	19	1010	∞
rs	1.2	7.3	11.0	49.9	0.1	2.1	16.5	9.0 5	5.8	103	549	491	30	320	92	09	72	570	19	139	435	23	48	23	74	7
loner	r 1.0	6.7	11.3	46.7	0.1	2.1	17.4	9.0 †	5.8	98	354	1 597	31	315	51	74	72	550	19	118	367	17	50	25	1074	6
LS	1.4	6.3	14.3	62.8	0.1	2.6	2.7	0.8	7.5	118	3 724	364	. 36	410	32	92	52	129	22	164	327	23	59	22	6889	48
rs	1.2	6.1	13.9	59.2	0.1	2.4	4.3	0.8	7.5	125	618	548	38	407	415	671	16	203	29	147	432	28	29	31	13,619	103
	1.3	6.5	12.2	55.2	0.2	2.2	10.8	3 0.7	6.3	105	587	488	33	341	82	119	55	371	22	151	457	23	52	24	3909	31
	11.1	0.9	9.3	8.7	28.2	2 8.2	43.1	10.1	11.6	11	17	14	12	15	124	140	34	41	13	10	88	12	16	17	95	06
loner	т 1.8	9.1	11.7	46.1	0.2	0.7	18.6	5 0.7	5.8	105	5 270	797	25	164	. 42	80	49	649	20	139	226	24	48	25	15	4
RLS	3 1.6	4.8	14.4	59.3	0.2	2.7	6.7	0.7	6.1	106	172	738	24	117	47	83	93	259	25	155	211	20	45	29	2990	23
RLS	3 1.4	12.5	10.9	51.3	0.2	1.4	10.6	9.0 9	7.6	105	516	1129	9 51	630	218	78	16	377	24	82	227	17	27	19	9841	75



account with an overlap correction in the Th calibration (Georgakopoulou et al. 2017), the lead contents in the Bahlā ceramics far exceed the standards used in the WDXRF calibration, and the correlation of Pb to Th values noted in the elemental results (Table 2), suggests the thorium contents are not reliable. They were thus disregarded as well and so was P_2O_5 that is associated with the burial contamination of archaeological ceramics (Freestone et al. 1985).

The cluster analysis performed on the raw chemical data additionally supports the heterogeneous character of Bahlā Ware, as the results of ceramic petrography suggest. Relations between the analysed samples are presented on the dendrogram (Fig. 5). One cluster emerges from the dendrogram (C1), consisting of samples of the LS fabric group and the petrographic loner B106. Slight compositional differences between the samples of C1, primarily in the CaO content (Table 2), are illustrated on the dendrogram. Excluding B114 and B195, the samples of C1 are calcareous, with the CaO content ranging between 8.3 and 17.4 wt%. However, similarities of other components suggest they should be seen as one group with varying CaO values. B114 and B195 are low-calcareous samples, with correspondingly higher contents of SiO₂, Al₂O₃ and Fe₂O₃. Sr, which is geochemically associated with Ca, is also lower, whilst Cr and Ni, associated with the serpentinitic component are increased. Furthermore, the value of Zn in B195 is affected by the glaze contamination being exceptionally high here and should thus be disregarded. It is thus clear that the two samples are rightly considered part of the same cluster, with the chemical distinctions noted here being a result of relative proportions of the different components.

The remaining three samples (B60, B132 and B252) can be described as outliers (Table 2; Fig. 5). Differences in major, minor and trace elements illuminate compositional differences compared to the samples of C1.

Fig. 5 Cluster dendrogram resulting from the cluster analysis performed on all samples included in the WDXRF analysis, using raw chemical compositions. Excluded oxides and elements are P₂O₅, Cu, Ba, Pb ad Th

The chemical composition of glazes

The chemical characterisation of the glazes determined through SEM-EDS shows that most samples of the LS fabric group are coated with a lead-barium glaze (Table 3). This is a glaze of heterogenous texture, as illustrated on the SEM photomicrographs (Fig. 6). The thickness of the glaze layer ranges between 80 and 200 µm. Quartz inclusions of various sizes are randomly scattered through the glaze, sometimes causing cracks in the texture (A on Fig. 6). Non-dissolved fragments of barium sulphate or baryte (BaSO₄) could be detected in the form of bright inclusions, ranging in size from small (20x20μm) rounded particles to large (100x100μm) subangular lumps (B on Fig. 6). The thickness of the ceramicglaze interface varies significantly, and in the case of BaO-rich glazes (B106) cannot be defined because the entire glaze layer is largely crystalline. The glaze is characterised by several crystalline phases of various compositions. The most abundant are Ca-rich pyroxenes, averaging 19.5at% Si, 8.8at% Ca, 6.8% at Mg and 4.0% at Fe, with minor amounts of Na, Al, K, Ba and Pb; some of these most likely also measured from the surrounding glaze. The composition of these crystals ranges between that of diopside (CaMgSi₂O₆) and hedenbergite (CaFeSi₂O₆) and they form at the ceramicglaze interface and/or float freely in the glaze (C on Fig. 6). Another crystalline phase, appearing brighter than the matrix in backscatter mode (D on Fig. 6) shows higher contents of Ba (4.2 at%), as well as Si (18.8at%), Fe (7.3at%), Al (4.4at%), and Mg, K and Ca at around 1at%. Their extremely small size precludes their individual analysis, as the surrounding matrix is also incorporated, so a direct identification of the mineralogy of these crystals is not attempted here. Furthermore, iron-rich clusters are occasionally present in the glaze.

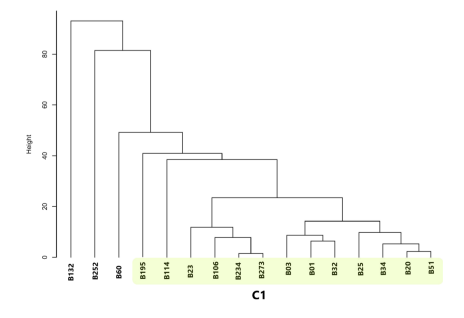
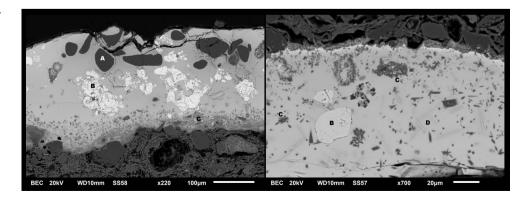




Table 3 The chemical composition of glazes determined through the SEM-EDS analysis. All results are normalised to 100 wt%. '-'indicates below detection limits. LS stands for the Limestone-Serpentine fabric group and RLS for related to Limestone-Serpentine fabric group

Sample	FG	Colour	Na ₂ O	MgO	Al_2O_3	SiO ₂	P_2O_5	K ₂ O	CaO	TiO ₂	MnO	FeO	CuO	ZnO	SnO_2	BaO	PbO
B1 bulk			1.1	3.0	4.7	44.7	0.3	2.2	7.2	-	-	12.0			-	15.4	9.0
B1 matrix	LS	Yellow	1.1	1.2	2.7	45.2		1.9	3.6			12.1	0.2			14.4	17.7
B3 bulk			1.4	1.3	3.0	45.3	_	3.3	3.9		_	9.8	_	_	-	10.2	21.8
B3 matrix	LS	Yellow	1.4	1.0	2.9	47.5		3.7	3.1			10.3				9.0	21.1
B8 bulk			1.3	2.3	3.0	43.5	_	1.4	6.3	_	_	9.8	0.2	_	_	11.1	20.8
B8 matrix	LS	Brown	1.5	0.5	3.2	40.0		1.5	3.7			10.1				14.0	25.6
B16 bulk			1.1	2.1	1.8	36.2	0.2	1.5	4.0	_	-	9.3	0.8	_	-	8.2	34.8
B16 matrix	LS	Brown	0.9	1.4	1.6	35.7		1.4	2.5			9.5	0.5			7.7	38.9
B20 bulk			1.3	1.3	1.8	33.7	_	1.7	4.4		_	10.4	_	_	-	6.4	39.0
B20 matrix	LS	Brown	0.9	1.2	1.5	34.1		1.4	2.7			9.0				5.7	43.5
B25 bulk			1.5	2.0	2.4	36.3	_	1.9	6.6	_	_	8.0	_	_	-	6.7	34.5
B25 matrix	LS	Brown	1.6	0.5	2.3	34.1		2.1	4.3			7.7				7.4	40.2
B32 bulk			1.1	1.0	2.4	46.4	_	2.3	3.4	_	-	9.5	0.2	_	-	9.2	24.5
B32 matrix	LS	Yellow	1.4	0.9	1.4	43.4		2.9	2.9			11.7				9.8	25.7
B34 bulk			1.3	1.3	1.3	46.2	-	3.1	3.4	_	-	11.6	_	-	_	10.5	21.4
B34 matrix	LS	Yellow	1.5	1.0	1.2	44.2		3.2	2.7			12.0				11.2	23.0
B106 bulk	LS	Yellow	0.8	2.9	5.7	48.7	_	2.9	8.3	_	_	13.5	_	_	_	12.4	4.5
B114 bulk			1.1	2.3	4.8	41.8	_	1.5	3.1		-	5.3	_	_	-	4.3	35.9
B114 matrix	LS	Yellow	1.1	1.3	4.3	39.9		1.4	1.5			5.6				4.4	40.6
B150 bulk			1.4	2.3	2.6	40.0	_	1.4	6.3	_	-	9.2	_	_	-	12.0	24.7
B150 matrix	LS	Brown	1.4	0.6	2.7	36.5		1.5	3.7			8.8				14.0	30.8
B279 bulk			1.2	1.1	2.9	39.2	_	2.2	2.5		_	11.4	_	_	_	5.0	34.5
B279 matrix	LS	Yellow	1.2	0.7	2.5	38.2		2.2	1.7			8.8				4.9	39.7
B281 bulk			0.4	2.1	3.6	44.1	0.3	0.6	4.8	_	_	13.4	_	_	_	29.5	1.0
B281 matrix	LS	Yellow	0.4	1.4	3.8	44.8		0.6	3.6			12.1				32.2	1.1
B287 bulk			0.7	2.0	1.9	40.0		1.0	4.3		-	8.1	_	-	_	10.1	31.9
B287 matrix	LS	Yellow	0.6	1.3	1.6	39.9		0.8	3.5			8.1				12.2	32.0
B195 bulk			0.6	1.2	2.2	33.7	-	0.4	1.8	-	-	1.3	0.8	3.4	_	0.6	54.0
B195 matrix	LS	Green	0.5	0.8	1.2	32.6		0.4	1.3			1.3	1.5	3.7		0.5	56.2
B252 bulk			0.6	1.2	0.9	33.5	_	0.8	1.4	0.3	0.4	5.5	_	-	0.8	-	54.8
B252 matrix	RLS	Yellow	0.6	1.2	0.8	33.1		0.7	1.4	0.3	0.4	5.2			0.5		55.9
B132 bulk			0.6	0.9	4.1	32.9	_	0.9	4.0	0.2	_	4.7	0.3	_	_	_	51.5
B132 matrix	RLS	Yellow	0.6	0.6	4.6	33.9		1.2	3.0	0.2		3.7	0.2				52.3

Fig. 6 SEM photomicrographs of lead-barium glazes detected on samples B3 (left) and B22 (right), taken in the BEC mode. Visible inclusions are quartz (A), baryte (B), Ca-rich pyroxenes (C), and probably Ba-rich pyroxenes (D)





The chemical composition of the glaze shows a negative correlation between PbO (1–39%) and BaO (4.3–29.5%), implying that the two are mineralogically associated (Table 3). Furthermore, the glaze contains 33.7–57.4% SiO₂, 2.5–8.3% CaO, 1.8–5.7% Al₂O₃, 1–3% MgO, 0.4–1.5% Na₂O and 0.6–3.3% K₂O. The content of FeO is high (5.3–13.5%), suggesting it was deliberately added as a colourant. The CuO value is <1%, which indicates it is probably an impurity associated with the lead minerals rather than a deliberately added colourant in the yellow and brown glazes.

An exception in the LS fabric group is B195 that contains a glaze of lead-zinc-barium type, whose chemical composition is characterised by 54% PbO, 3.4% ZnO, 0.6% BaO, 33.7% SiO₂, 2.2% Al₂O₃, 1.8% CaO, 1.2% MgO and 1% Na₂O + K₂O (Table 3). Compared to the group of lead-barium glazes, B195 has significantly less BaO, which does not fit the correlation with PbO noted in the lead-barium glazes. This result might indicate the use of different raw materials. The texture is, on the other hand, similar to the lead-barium group, containing non-dissolved fragments of quartz as well as Ca-rich pyroxenes in the interface and glaze (Fig. 7).

Interestingly, the samples B132 and B252 that do not belong to the LS fabric group are coated with a high-lead glaze that contains no traces of BaO (Table 3). Both samples contain over 50% of PbO and around 33% of SiO₂, but differ in the contents of Al_2O_3 and CaO. Furthermore, SnO_2 (0.8%) is detected in B252, visible as bright crystallites in the glaze (Fig. 7(c).

Lead isotope analyses of the glazes

Five glaze samples from the LS fabric group were analysed for their lead isotope ratios; one was B195, the lead-zinc-barium glaze, whilst the other four were all typical lead-barium glaze samples (Table 4). The results show relative standard deviations between 0.22% for the ²⁰⁸Pb/²⁰⁶Pb ratio to 0.45% for ²⁰⁶Pb/²⁰⁴Pb. Ore deposits of uniform geological and geochemical history are reported to have a

spread of lead isotope ratios up to a maximum of 0.3% (Gale and Stos-Gale 1992) or even 0.6% (Pernicka et al. 1990, p. 283). In principle thus, the lead for all five samples could have been sourced from the same deposit. Sample B195 is, however, clearly separated from the other four (Fig. 8) which together with its different chemistry, probably indicates a different source.

Discussion

Ceramic traditions of Late Islamic Arabia

The scientific assessment of Bahlā Ware consumed in al-Ain offers evidence for continuity of a technological tradition that spans between the mid-seventeenth and the early twentieth centuries. This tradition is defined by the clear correlation between the LS fabric group and the leadbarium glaze, showing standardisation in all segments of the *chaînes opératoires*. The transfer of knowledge and skills embedded in this tradition is telling of the cultural importance of Bahlā Ware for the communities settled in South East Arabia.

The technological diversity of Bahlā Ware is documented only in the eighteenth century (HRZ 9.2) at the Bin 'Ātī site. Although modest in number, the samples different to the LS fabric group indicate the presence of diverse technological practices, for both ceramics and glazes, that were used for manufacturing morphologically similar pots. The exploitation of similar secondary clays for the preparation of technologically different pastes coated with equally different glazes indicate the emergence of new workshops with distinctive practices.

Lead-barium glaze in the Islamic world

In the broad spectra of Islamic glazes and glasses in the Near East (e.g. Brill 2001; Freestone 2006; Tite 2011; Henderson 2013), the lead-barium glaze of Bahlā Ware has no parallels.

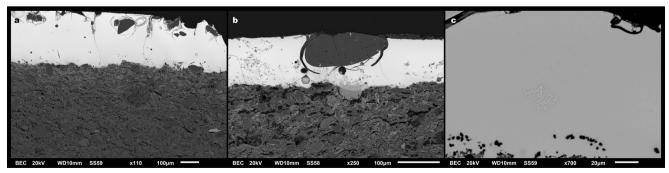


Fig. 7 SEM photomicrographs of lead-barium-zinc glaze of sample B195 (a) and high-lead glazes of samples B132 (b) and B252 (c) taken in BEC. Visible inclusions on images a and b are identified as quartz whilst image c contains inclusions of SnO₂



Table 4 Lead isotope ratios of Bahla Ware from al-Ain

Samples	²⁰⁶ Pb/ ²⁰⁴ Pb	²⁰⁷ Pb/ ²⁰⁶ Pb	²⁰⁸ Pb/ ²⁰⁶ Pb
B1	18.11690	0.86542	2.11393
B25	18.11119	0.86502	2.11283
B32	18.11895	0.86525	2.11371
B195	18.30110	0.85831	2.10357
B281	18.13577	0.86529	2.11465
RSD (%)	0.45	0.36	0.22

In fact, this type of glass is considered to be a Chinese innovation (Fuxi 2009, p. 20; Henderson 2013, p. 123; Henderson et al. 2018), and unknown outside of East Asia (Cui et al. 2011, p. 1671; Rehren and Freestone 2015, p. 234). Therefore, Bahlā Ware offers the first evidence for the production of lead-barium glaze in the Islamic World. In the absence of relevant analogies, the emergence and development of this glaze technology remain unclear. In the geographically close Iran, the seventeenth-century tiles of the Safavid period have glazes of the alkaline and lead types without traces of barium (Holakooei et al. 2014). The seventeenth-century Mughal tiles in India are coated with alkali glazes (Gill and Rehren 2011). Therefore, the lead-barium glaze of Bahlā Ware stands out from other contemporary traditions in the Islamic world.

The presented results enable a preliminary reconstruction of the lead-barium glaze technology and methods of application. The clear correlation between PbO and BaO determined through SEM-EDS supports their mineral association in nature. Baryte commonly occurs with lead sulphides such as galena. For the assessment of the original glaze composition and methods of application, the contents of PbO, BaO and CuO, oxides clearly coming from the glaze, were excluded from the analyses of the ceramic fabrics and glazes, and the

remaining compositions were recalculated and normalised to 100% (see Tite et al. 1998, pp. 249-250). The data presented in Table 5 show consistent differences in the compositions of bodies and glazes, for example, a significantly increased Si/Al ratio in the glaze, suggesting that the glaze was applied as a lead/barium-silica mixture. Since the glaze contains large pieces of non-dissolved baryte and quartz, it is more likely that they were applied as a suspension directly on the body. Experiments conducted on high-lead glazes suggest a high firing temperature (950 °C and above) and slow cooling rate (20 °C/h) for glazes showing an extended crystalline layer (Molera et al. 2001, pp. 1121–1122), which is the case with the Bahlā glazes. However, the reaction occurring between lead and baryte in glazes remains understudied, which precludes firm conclusions on the firing temperature. The sparkling look of the glaze, noticed by several scholars (Hansman 1985, p. 52; Kennet 2004, p. 42; Carter 2011, p. 37), could be explained by the presence of crystalline phases formed during devitrification. Finally, iron oxide was used as a colourant, which means that the colour gradient depends on the firing conditions.

It is challenging to explain the reasons for the use of sulphidic minerals in glaze preparation, as these are known for their poor fluxing properties (Brill et al. 1991b, p. 34). The role of deliberately added barium-bearing material in the early Chinese glasses was to achieve a jade-colour opacity (Brill et al. 1991b, p. 34; Cui et al. 2011, p. 1675), which was not needed in the case of the monochrome glazed Bahlā Ware. Also, it appears that PbO and BaO were part of the same naturally mixed batch and their amounts could not be controlled. A deliberate choice of the lead-barium compound as a raw material is indisputable, though. The lead isotope ratios dated to all three archaeological horizons in al-Ain suggest the exploitation of a single source for the lead-barium material. This continuity would not exist without a strong cultural association of craftsmen with this technological choice.

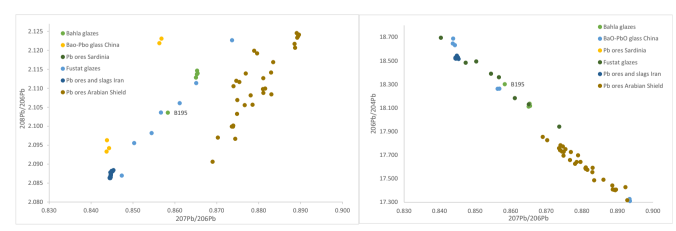


Fig. 8 The binary plot shows lead isotopes measured in five glazes of Bahla Ware (Table 4), compared with lead-barium glass from China (Cui et al. 2011), Pb ores from Sardinia (Stos-Gale et al. 1996; Gale 2011),

glazes from Fustat (Wolf et al. 2003), Pb ores and glazes from Iran (Pernicka et al. 2012) and Pb ores from the Arabian Shield (Stacey et al. 1980; Stacey and Stoeser 1983)



Table 5 Chemical compositions (SEM-EDS) of bodies and glazes excluding CuO, BaO and PbO in the latter and, normalised to 100 wt%. '-'indicates below detection limits

Samples	Na ₂ O	MgO	Al_2O_3	SiO_2	P_2O_5	K_2O	CaO	TiO_2	FeO
B1 glaze	1.5	3.9	6.3	59.5	0.4	2.9	9.6	_	16.0
B1 body	1.2	5.9	12.5	59.5	_	2.3	10.5	0.7	7.0
B8 glaze	2.0	3.4	4.5	64.2	_	2.1	9.3	_	14.5
B8 body	2.3	6.0	13.9	57.0	_	1.9	12.4	0.4	4.8
B16 glaze	1.9	3.7	3.2	64.5	0.4	2.7	7.0	_	16.5
B16 body	1.3	5.8	12.4	59.2	_	2.1	11.4	0.7	6.5
B20 glaze	2.4	2.5	3.4	61.8	_	3.0	8.0	_	19.0
B20 body	1.3	6.5	12.8	59.6	_	2.2	10.3	0.7	6.1
B25 glaze	2.6	3.4	4.1	61.9	_	3.2	11.2	_	13.6
B25 body	1.1	6.3	11.1	56.3	_	2.0	15.9	0.8	6.1
B34 glaze	1.9	1.9	1.9	67.8	_	4.6	4.9	_	17.0
B34 body	1.3	6.3	12.1	55.5	_	2.3	14.4	0.7	6.2
B150 glaze	2.2	3.6	4.2	63.2	_	2.2	10.0	_	14.6
B150 body	1.0	7.0	11.8	54.1	_	2.0	16.0	0.6	6.5
B281 glaze	0.6	3.0	5.2	63.5	0.5	0.9	6.9	_	19.3
B281 body	1.2	8.0	12.1	54.7	0.3	2.2	14.3	0.7	6.5
B287 glaze	1.2	3.4	3.2	69.0	_	1.8	7.5	_	13.9
B287 body	1.2	6.3	12.8	61.8	_	2.2	8.5	0.6	6.5

Ceramic provenance of the LS fabric group

An efficient provenance study of archaeological pottery requires a holistic approach, including testing potential clay sources and their comparison with archaeological pottery (Tite 2001). In this case, designed to be a pilot study, large-scale testing of clays was not feasible. Therefore, this preliminary provenance study relies on a comparison between the petrographic data and geological maps of Oman (Bechennec et al. 1986a, 1986b) and Iran (Spaargaren 1991).

The petrographic results point to an environment rich in limestone and serpentinite used for clay exploitation. The presence of limestone and rounded and sub-rounded quartz in the calcareous matrix suggest the formation of rocks by accumulation in a sedimentary basin, where detritic minerals associated with igneous rocks were deposited. Both petrographic and chemical compositional variations, such as the CaO variability, documented in the LS fabric group could be explained with this geological setting.

The petrology of the inclusions is consistent with the geological composition of the ophiolitic mountains in Oman (Lippard et al. 1986, p. 62), and more specifically with that of wadis that intersect rock formations (Hanna 1995). In semi-arid environments, alluvial fans are formed during a movement of materials from upper to lower elevations, especially during humid seasons (Rapp and Hill 2006, pp. 63–64). The town of Bahlā, one of the potential production centres, lies on one of these wadis surrounded by the ophiolitic mountains (Bechennec et al. 1986a). Alluvial terraces, located just 5–7 km north of the modern town, contain abundant olistoliths

of dolomite and biolithoclastic limestone, breccia and limestone with chert nodules, sandstone with calcareous matrix, serpentinized harzburgite and basalts (Bechennec et al. 1986b), and therefore could be a potential source of raw material. The historical and ethnographic importance of Bahlā for the regional ceramic production would support this possibility (Whitcomb 1975, p. 129). However, the absence of archaeologically documented production debris in the town obscures a more accurate identification.

With more certainty, it is possible to discard Khunj in Iran as a production centre for the Bahlā Ware consumed in al-Ain. Khunj is located in the anticline Fars domain of the Zagros Fold Belt, a region with sedimentary geology (Spaargaren 1991) and lies far beyond the ophiolitic zone (Momenzadehs 2004; Ghorbani 2013, pp. 47–50). Some other areas of Central Iran (High Zagros) are rich in ophiolites and geologically match the composition observed in the Bahlā Ware, but there is no archaeological evidence to suggest production of it there.

Glaze provenance of the LS fabric group

Although lead isotope analysis has been used for provenance determination of lead glasses for several decades (Brill and Wampler 1967), the method only recently started being used for the study of Islamic glazes (Wolf et al. 2003; Mason et al. 2011). The scarcity of comparable datasets for lead ores in the Middle East sets a limitation for absolute provenance attribution. This is especially relevant for this research; where the main subject of interest is the lead-barium mineralisation.



Oman is rich in copper ores (Begemann et al. 2010), but lead-barium sources are not known. Extensive lead-barium mineralisation occurs in Central Iran and Alborz, always in association with zinc-lead ores (Ghorbani 2013, p. 169). However, the available isotope ratios for representative minerals in the zinc-lead deposits from Sanadaj-Sirjan, Urumeih-Dokhtar and the Zagros zones (Ehya et al. 2010; Mirnejad et al. 2011) do not match the Bahlā glazes (Fig. 8). Even more distant are lead ores, lead slags and litharge from the Central Iranian Plateau (Pernicka et al. 2012, p. 670). The same is true for lead ores in the Saudi Arabia shield (Stacey et al. 1980; Stacey and Stoeser 1983).

An overview of lead isotope ratios from distant China, which is usually associated with lead-barium glass and glaze production (Brill et al. 1991a; Cui et al. 2011), does not offer a match with the glazes from al-Ain (Fig. 8). The closest association can be made with the ores of galena, pyrite and baryte from Sardinia (Stos-Gale et al. 1996; Gale 2011). Similarly, close ratios are reported for three glazed samples from Mamluk's Fustat dated to the fourteenth century (Wolf et al. 2003, p. 411). However, this proximity in lead isotope ratios does not provide an adequate framework for an archaeological interpretation of the provenance of Bahlā Ware, and may well be a consequence of random isotopic overlap (Henderson et al. 2005). In short, the absolute provenance of the lead-rich components of these glazes remains unknown.

Conclusion

This pilot project examines technological patterns of Bahlā Ware consumed in al-Ain between the mid-seventeenth and the early twentieth centuries in order to shed more light on the production and provenance of glazed ceramics in Late Islamic Arabia. The petrographic data presented here, together with those deriving from the archaeological investigations, indicate that Bahlā Ware is more likely to be the product of workshops located in Oman, possibly Bahlā itself, than Iran. This production was standardised and consisted of technological choices that were carefully transmitted from one generation of potters to the next over the course of three centuries. The peculiar aspect of this production refers to the glaze recipe, made of a lead/barium and silica mixture. Whilst the source of the lead-rich raw materials remains unclear due to the lack of comparable data, it is certain that the use of a mixed lead-barium mineral was a deliberate choice. The continuity in the exploitation of a single source, as the lead isotopes demonstrate, supports this conclusion. The chronological concentration of technological variability in the eighteenth century suggests the introduction of different workshops and technological solutions for the production of this ware, potentially related to increased regional demand. Further work will clarify whether this diversity is restricted to the eighteenth century or al-Ain only got access to multiple production networks in this period of economic peak.

A better grasp on the origins and technology of the leadbarium glaze in the Islamic World has to be sought within the regional context of Arabia and the Gulf. Further work is required into other assemblages dated to the Late Islamic period, including other classes besides Bahlā, as well as a holistic approach towards the question of production centres.

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