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An archaeometric study of Hellenistic glass vessels: evidence for multiple sources

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Abstract In the present study, 53 glass fragments from core-formed vessels and 3 glass beads are investigated using SEM/EDX, EPMA and LA-ICP-MS. All samples were excavated in the Latin settlement of Satricum in central west Italy and apart from two, were found in the so-called fourth–third c. BC Hellenistic Votive deposit, also known as Votive Deposit III, discovered in front of the sanctuary of *Mater Matuta* on top of the acropolis. The analytical results indicate that the glass from Satricum is a typical soda-lime-silica type with natron used as a flux. Its chemical compositions display a relatively low compositional variation. Small differences in the concentrations of major and minor oxides (SiO_2 , Al_2O_3 , CaO and Fe_2O_3) and in trace elements (Sr, Zr and Nd) between individual samples suggest the use of different types of raw materials, especially sand. In turn, this suggests that the glass derived from more than one glass making centre. The combined investigation of colourants (Co, Cu and Mn) reinforces and confirms the idea that glass from Satricum was made using different manufacturing traditions during the Hellenistic period.

Keywords Natron glass · Core-formed vessels · Hellenistic period · Italy · Satricum · Trace elements · Chemical composition · SEM-EDX · EPMA · LA-ICP-MS

Introduction

Archeometric glass studies have focused on the analysis of material from various periods, from Bronze Age to the Middle Ages. The scientific community has been mainly interested in investigating glass dating from the Late Bronze Age (fourteenth to twelfth c. BC) and from late antiquity (first to ninth c. AD) while little interest has been given to glass from the intervening period (seventh to first c. BC) (Brill 1999; Triantafyllidis 2000a, b; Rehren et al. 2005; Zacharias et al. 2008a, b; Oikonomou et al. 2008; Sokaras et al. 2009; Arletti et al. 2011; Beltsios et al. 2012; Connolly et al. 2012; Oikonomou et al. 2012; Triantafyllidis et al. 2012; Oikonomou et al. 2014; Palamara et al. 2015; Rehren et al. 2015). The focus of this paper is an analytical study of early Hellenistic glass excavated in Satricum, Italy, in order to redress the balance of our knowledge for glass from this time period. Little is known about primary or secondary glass production during Hellenistic times. The main objectives of this work are to shed light to the glass technology used in this period and to try to answer suggest a provenance for the glasses.

The glass finds investigated in this paper come from the Latin settlement of Satricum situated on the banks of the river Astura, ca. 60 km south of Rome (Fig. 1). This settlement developed from a modest hamlet of huts in the ninth c. BC, perched on top of an ‘acropolis’ hill, into a prosperous urban centre in by the sixth c. BC covering an area of nearly 40 ha. The site was discovered in the late nineteenth c., after which Italian archaeologists excavated large parts of the settlement (1896–1898 and 1907–1910). Research in Satricum was taken

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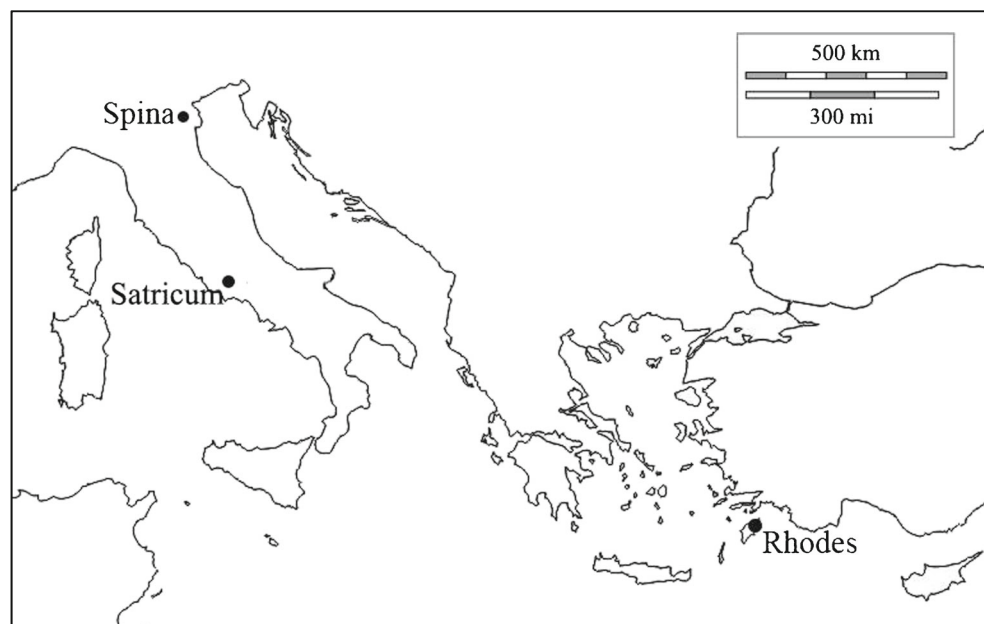
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Fig. 1 Map indicating the location of the sites discussed in this paper



up again in 1977 by Dutch archaeologists who are still active on the site.

Satricum's favourable location at the crossroads between the northern Etruscan and southern Greek areas is reflected in the remarkable diversity of the archaeological material. At the same time, it is one of the best preserved sites in central Italy revealing archaeological remains which cover a period of continuous occupation from nearly ten centuries (ninth c. BC–first c. AD). The site is probably best known for its sanctuary dedicated to *Mater Matuta*, goddess of dawn. Situated on top of the acropolis, it incorporates three successive temple buildings from the Archaic period onwards. The importance of the sanctuary is reflected in three substantial votive deposits which testify to offering practices over a long period of time. Among them is the so-called Hellenistic Votive deposit, also known as Votive Deposit III, discovered in front of the temple at the end of the nineteenth c. During the re-excavation of the deposit between 1986 and 1990, substantial numbers of ex-voto came to light that were either not excavated by the nineteenth c. excavators or left behind by them because of their fragmented state of preservation. Among these finds which consisted of a large variety of Hellenistic ceramics as well as numerous categories of terracotta votive offerings, some 140 fragments of core-formed glass flasks and several glass beads were found.¹ So far, this is the largest corpus of Hellenistic core glass vessels known in Latium. The fragments of the core-formed vessels belong to the Mediterranean Group II industry which dates between mid fourth to third c. BC. Core-formed vessels of the first millennium BC can be

distinguished in three major glass industries (Mediterranean groups I, II and III) that prevailed in Mediterranean area between mid sixth c. BC to early first c. AD. The typology of these glass vessels follows the typical shapes of pottery and metalware of Archaic, Classical and Hellenistic periods such as alabastra, amphoriskoi, oinochoai and aryballoi (Grose 1989; Stern and Schlick-Nolte 1994). The material constitutes an exotic category among the predominantly central Italian objects in the votive deposit. The current study aims to improve our knowledge on the composition of the glass and the provenance of the material, thus providing valuable information on the position of Satricum in the commercial networks of this period (Gnade 2002, 2007).

Materials and methods

Materials

In the present study, 53 fragments of core-formed glass vessels and 3 fragments of glass beads are investigated. The samples come from various vessels types. Specifically, the vessel assemblage consists of 25 alabastra, 8 oinochoai, 7 amphoriskoi, 4 hydriskoi, 1 aryballos and 8 unidentified glass vessels. The beads assemblage consists of 2 eye beads and 1 bead with white decorative trails (Table 1). The majority of glass vessels bear opaque white or/and yellow decorative zigzag or straight trails, while there are few examples of blue and turquoise decorative trails (Fig. 2).

Small pieces (2–3 mm) were cut from the vessel fragments using a diamond cutting disc. The samples were mounted in a resin block. The resin block was ground with silicon carbide

¹ The core-formed glass vessels are presently under study as part of a larger project concerning the publication of all votive offerings found in the Hellenistic votive deposit.

Table 1 General description of the glass artefacts analysed

Sample	Main body colour	Decoration colour/type	Vessel part	Typology	Analytical technique	Group
Sa.4	Dark blue	–	Unidentified	Unidentified	SEM/ICP	A
Sa.5	Turquoise	Yellow	Rim	Alabastron	SEM/ICP	A
Sa.16	Dark blue	White/zigzag	Body	Alabastron	SEM/ICP	A
Sa.21	Turquoise	Yellow and white/feather	Body	Alabastron	SEM/ICP	A
Sa.35	Dark blue	Yellow and white/festoon	Body	Aryballos	SEM/ICP	A
Sa.40	Brown	Yellow, white and Turquoise/feather	Body	Oinochoe	SEM/ICP	A
SAT.5	Blue	Yellow and white	Shoulder	Unidentified	EPMA	A
Sa.1	Dark blue	–	Unidentified	Alabastron	SEM/ICP	B
Sa.2	Dark blue	–	Rim	Alabastron	SEM/ICP	B
Sa.3	Dark blue	–	Body	Alabastron	SEM/ICP	B
Sa.6–14	Dark blue	Yellow and white/zig zag	Body	Alabastron	SEM/ICP	B
Sa.15, 17–20, 22	Dark blue	Yellow and white/feather	Body	Alabastron	SEM/ICP	B
Sa.23	Dark blue	White/feather	Body	Alabastron	SEM/ICP	B
Sa.24	Dark blue	Yellow and white/zig zag	Body	Alabastron	SEM/ICP	B
Sa.25–27	Dark blue	Yellow/straight	Body	Amphoriskos	SEM/ICP	B
Sa.28–29	Brown	Yellow and white/zig zag	Body	Amphoriskos	SEM/ICP	B
Sa.30	Brown	Yellow and white/straight	Body	Amphoriskos	SEM/ICP	B
Sa.31	Dark blue	-	Body	Hydriskos	SEM/ICP	B
Sa.32–33	Dark blue	Yellow and white/straight	Body	Hydriskos	SEM/ICP	B
Sa.34	Dark blue	–	Base	Hydriskos	SEM/ICP	B
Sa.36–37	Dark blue	Yellow and white/straight	Body	Oinochoe	SEM/ICP	B
Sa.38–39	Dark blue	White/zigzag	Body	Oinochoe	SEM/ICP	B
Sa.41	Dark blue	Yellow/straight	Body	Oinochoe	SEM/ICP	B
Sa.42	Dark blue	–	Base	Oinochoe	SEM/ICP	B
Sa.43	Dark blue	Yellow/zigzag	–	Bead	SEM/ICP	B
Sa.44–45	Dark blue	White/eyes	–	Eye bead	SEM/ICP	B
SAT.2	Dark blue	Yellow/straight	Base	Pinochoe	EPMA	B
SAT.1	Dark blue	Yellow and white/trails	Body	Amphoriskos	EPMA	C
SAT.3	Dark blue	Yellow	Neck and shoulder	Unidentified	EPMA	C
SAT.4	Dark blue	–	Neck	Alabastron	EPMA	C
SAT.6	Dark blue	Yellow/zigzag	Body	Unidentified	EPMA	C
SAT.7	Dark blue	Yellow, white and turquoise	Body	Unidentified	EPMA	C
SAT.8	Dark blue	Yellow and white/trails	Body	Unidentified	EPMA	C
SAT.9	Dark blue	Yellow and white/trails	Rim	Unidentified	EPMA	C
SAT.10	Dark blue	Yellow and white/trails	Body	Unidentified	EPMA	C
SAT.11	Dark blue	No	Rim	Unidentified	EPMA	outlier

paper of various grits (600, 800, 1200, 2500) and then polished using diamond paste of 6–3 and 1 μm .

Scanning electron microscopy/energy dispersive X-ray spectrometry (SEM/EDX)

Scanning electron microscopy was the analytical technique used for the detection of major and minor elements of 45 glass samples (42 vessels and 3 beads). A JEOL (JSM-6510LV) scanning electron microscope, coupled with an

energy-dispersive X-ray spectrometer Oxford instruments, was used. All samples were analysed under high vacuum, with an operating voltage at 20 kV and working distance for each sample of 15 mm. The calibration of the system was performed with geological standards and the accuracy/precision it was established by analysing standard reference materials (NIST SRM620, SRM1831 and SRM612). The analyses are in close agreement with the expected values and are presented in Table 2. The relative error between the expected and measured values is approximately 5 % for most



Fig. 2 Sample fragments from Satricum. Clockwise from top left: **a** Cobalt blue core-formed alabastron, **b** Cobalt blue core-formed alabastron, **c** Brown core-formed amphoriskos, **d** Blue green (turquoise) rim from a core-formed alabastron, **e** Cobalt blue bead with opaque yellow zigzag trail

of the oxides. Due to submicrometer beam size, 5 analyses of 300 s were performed on each sample, and the mean value was calculated for each element.

Electron probe micro analysis (EPMA)

Electron probe microanalyses were carried out by Dr. Andy Tindle in order to detect major and minor elements in 11 fragments of core-formed vessels (see Table 5). A Cameca SX50 is located at the Department of environment, earth and ecosystems, The Open University was run at 20 kV accelerating voltage and 20 nA beam current. Quantitative analyses were produced for all major, minor and trace elements detected. Analyses were performed using a defocused electron beam of 20 μm in diameter so as to prevent the volatilisation of light (low atomic number) elements such as soda and magnesium. Multi-element glass standards (Corning glass standards and NIST) were analysed on a regular basis so as to establish and monitor the accuracy and precision of the machine. Table 3 provides a comparison of quoted oxide results compared with our measured results with associated standard deviations. The levels of detection varied from 170 ppm for CaO to 1200 ppm for CuO. A ZAF programme was used to correct and quantify the results.

Laser-ablated inductively plasma mass spectrometry (LA-ICP-MS)

Trace element analysis was performed by laser ablation—inductively coupled plasma—mass spectrometry (LA-ICP-MS). The LA-ICP-MS instrument consisted of a NewWave UP193FX excimer (193 nm) laser system with built in microscope imaging coupled to an Agilent 7500 series ICP-MS. Laser ablation craters were set at 70 μm , the laser being fired

Table 2 Measured and expected values of major and minor oxides for standard reference materials (in % wt.)

SAMPLE	Na ₂ O	MgO	Al ₂ O ₃	SiO ₂	SO ₃	K ₂ O	CaO
SRM620	14.19	3.66	1.72	73.59	0.21	0.39	6.27
Expected	14.39	3.69	1.8	72.08	0.28	0.41	7.11
SRM1831	13.67	3.67	1.20	74.03	0.27	0.29	6.9
Expected	13.32	3.51	1.21	73.08	0.25	0.33	8.2
SRM612	13.94	–	2.06	72.93	–	–	11.08
Expected	13.7	–	2.03	72.1	–	–	11.9

The expected values for SRM612 were provided by GeoRem (Jochum et al. 2011)

for 45 s at 10 Hz and a typical fluence of 2.8 Jcm^{-2} . Data was collected in a time resolved analysis mode, with a gas blank being measured before a series of ablations on glass samples, calibration standards and quality control standards, were carried out. Calibration standards bracketed the samples and QC over a period of 1 h or less.

Calibration of the system was performed using NIST SRM610 trace element glass standard. The measured and expected values are presented in Table 4. NIST SRM612 was used for quality control purposes.

Results and discussion

Type of glass

All Satricum samples are of a soda-lime-silica type (Sayre and Smith 1961). Silicon dioxide (SiO₂), the main glass former, varies between 68.61 % wt. and 74.01 % wt. with a mean value of 70.58 % wt. The main source of SiO₂ can be either sand or crushed quartzite pebbles. Sand, a less pure source than quartz, exhibits elevated amounts of impurities such as aluminium oxide (Al₂O₃) and iron oxide (Fe₂O₃) (Nicholson and Henderson 2000). In the present study, the mean values of Al₂O₃ and Fe₂O₃ are 2.08 % wt. and 0.24 % wt., respectively, close to the typical values of such impurities found in sands; although we are not suggesting this as a source, sand from the Belus river on the Levantine coast is thought to have been a suitable sand source used during antiquity, containing 2.98 % wt. Al₂O₃ and 0.325 % wt. Fe₂O₃, values that do not change significantly the base glass composition (Brill 1988). When evaluating potential sand sources, it should be considered that they exhibit variability in their chemical composition and not all sands are suitable for glassmaking (Brems et al. 2012; Degryse 2014).

Potassium oxide (K₂O) and magnesium oxide (MgO) are found in concentrations below 1 % wt., indicating that natron was the flux used to introduce a mean sodium oxide (Na₂O) level of 17.55 % wt. The evaporite deposits consist of sodium-rich minerals such as natron (Na₂CO₃.10H₂O) and/or trona

Table 3 The recommended composition for the Corning B standard (Wagner et al. 2012) compared to average analytical results ($n = 22$) and associated standard deviations using the electron microprobe (in % wt.)

	SiO ₂	Al ₂ O ₃	Na ₂ O	K ₂ O	CaO	TiO ₂	Fe ₂ O ₃	MnO	MgO	CoO	CuO	P ₂ O ₅	Sb ₂ O ₃
Measured	60.76	4.18	17.35	1.02	8.51	0.1	0.33	0.24	1.02	0.05	2.95	0.81	0.67
Quoted	62.02	4.63	16.5	1.3	8.75	0.099	0.31	0.24	0.99	0.043	2.82	0.61	0.42
St. dev.	1.03	0.04	0.17	0.03	0.27	0.02	0.03	0.03	0.05	0.02	0.09	0.11	0.08

(Na₂CO₃·NaHCO₃·2H₂O) (Shortland 2004; Henderson 2013). Calcium oxide (CaO, with a mean value 6.90 % wt.) can be introduced in the glass either but unlikely as a deliberate additive or usually as in our case as an impurity in the form of marine shells in coastal sands (Henderson 2013) or freshwater shells in river sands.

The majority of samples have a deep blue colour. Their colouration is attributed to the simultaneous presence of both cobalt (Co) and copper (Cu) which have mean values of 1132 and 1713 mg/kg, respectively. Two samples have a turquoise colour because of the elevated amount of Cu (5542 and 5307 mg/kg), while four samples are brown but without significantly high values of colourant elements (e.g., Fe, Mn, Cu). Their colouration is probably due to varied furnace atmospheres (Henderson 2000). These samples have elevated boron concentrations (above 350 mg/kg), which is an additional chemical discriminant, while the mean value in boron of the remaining samples is 177 mg/kg. Opacity in yellow and white glasses is due to the presence of antimony Sb and lead Pb which have mean values of 717 and 2049 mg/kg, respectively. Calcium and lead antimonate crystals are responsible for the opacity of ancient glass (Lahlil et al. 2008).

Sand source

Al₂O₃ is present in varying amounts in most glass artefacts in antiquity. It is introduced in the glass batch most of the time as an impurity in the sand used in glassmaking (Jackson et al. 2005; Nicholson and Henderson 2000); we should not exclude its deliberate addition (Beltsios et al. 2012) which is not the case in the samples in this study. Therefore, Al₂O₃ is a possible way to differentiate between primary sand sources. A bi-plot of SiO₂ and Al₂O₃ in Satricum samples and coeval samples from Spina, Italy (Arletti et al. 2011) and Rhodes Island, Greece (Triantafyllidis et al. 2012) is shown in Fig. 3. Furthermore, in the same figure, the concentrations of two hypothetical glasses derived from suitable Italian sands for glassmaking, according to Brems et al. (2012), are plotted.

All samples can be divided in 3 basic groups according to their SiO₂ and Al₂O₃ levels. The first group (group A) has values of SiO₂ and Al₂O₃ above 68 % wt. and below 1.5 % wt., respectively, consisting of 7 Satricum samples and 2 samples from Spina (low Al₂O₃ group). The second group (group B) has values above 68 % wt. and 1.5 % wt. respectively,

consisting of the majority of Satricum samples (40 samples out of 56) and 6 samples from Spina, showing elevated values in both elements (Table 5). Finally, the third group (Group C) has values below 68 % wt. and above 1.5 % wt., respectively, consisting of all the Rhodian samples with the addition of 8 Satricum samples and two samples from Spina. The sample close to group A can be considered either as a group A sample with marginal SiO₂ content or as an outlier (SAT.11).

Evidence for the use of different sand sources can be derived from trace element characterization. Forty-five out of the total of 56 samples were analysed by means of LA-ICP-MS for the determination of their trace element compositions. The 45 samples analysed are part of group A and B samples (Fig. 3, Tables 1 and 6). Unfortunately, due to sampling criteria, none of group C samples were analysed by LA-ICP-MS to identify their trace element composition. By plotting Nd against Sr, an interesting correlation can be seen (Fig. 4). There are two groups of samples with positive correlations between the two elements but with different slopes. Samples with a higher slope are part of the samples that form group A (see above) with only one addition (sample Sa.23), which has a basic glass composition similar to group B samples (high Al₂O₃, high SiO₂). A similar positive correlation between these two elements was found in plant ash Late Bronze Age glass from Egypt and Greece (Henderson 2013). It has been suggested that the latter correlation was derived from the fact that plant ash was contributing to the excess of Nd and therefore the combination of plant ash and sand created this correlation (Henderson et al. 2010; Henderson 2013). This study is different because natron-based glasses have no contribution from plant ash so a different explanation needs to be sought. Sr is an element associated mainly with Ca and by implication primarily derived from shells in the sands used; however, a contribution from feldspars or/and heavy minerals in the silica source cannot be excluded (Henderson et al. 2005; Degryse et al. 2006, 2010). Nd is presumed to be related to accessory minerals such as zircon present in sand (Brems et al. 2014). Thus, differences in the Nd/Sr ratio between the two groups could reflect use of sands with different proportions of “contaminant” minerals.

Strontium (Sr), barium (Ba) and zirconium (Zr) are three elements, associated with various minerals in rocks or sediments and hence in derived sands. Their concentration varies and reflects the local geology of the sand precursors.

Table 4 Measured and expected values (mg/kg) of trace elements for standard reference glass NIST612 by LA-ICP-MS

Sample	Li	B	Ti	V	Cr	Mn	Co	Ni	Cu	Zn	As	Rb	Sr	Y	Zr	Sn	Sb	Ba	La	Ce	Nd	Pb	Th	U
NIST 612	39.5	34.9	40.7	37.8	35.0	37.6	34.2	38.8	37.2	43.4	33.3	31.4	77.8	38.2	38.1	38.4	33.1	39.3	35.6	37.7	34.4	38.0	36.6	36.3
Expected	40.2	34.3	44.0	38.8	36.4	38.7	35.5	38.8	37.8	39.1	35.7	31.4	78.4	38.3	37.9	38.6	34.7	39.3	36.0	38.4	35.5	38.6	37.8	37.4

The expected values for SRM612 were provided by GeoRem (Jochem et al. 2011)

Zirconium is expected to be primarily present as zircons in sands while Sr is connected with the presence of Ca which derives from shells (aragonite) or/and limestone. Barite (BaSO_4), which is found as concretions in sands and sandstones, is the likely main mineral in sands. These three elements can therefore be possible independent markers for differentiating sand sources used to make glasses.

In Fig. 5, a bi-plot of Sr versus Zr concentrations, samples divide into two groups. The majority of the samples form a group with low Zr values (29–42 mg/kg) and varying Sr compositions (270–562 mg/kg). Nine samples have elevated values of Zr (above 50 mg/kg). Five of them (yellow diamonds) are part of group A samples and show a positive correlation. This clear distinction, between part of group A and the majority of group B samples, strengthens the idea that they were manufactured with different raw material sources. Four samples have values “between” the two main groups. This could reflect a third group or could be a result of recycling. This distinction is also obvious in our Ba against Zr plot (Fig. 6) where the majority of samples have elevated Ba and lower Zr values while there are some samples with elevated Zr values ($\sim 50 < \text{Zr} < 130$ mg/kg) suggesting use of different sands (Shortland et al. 2007; Henderson 2013). The five samples (yellow diamonds) are again positively correlated while the four “recycled” have a distinct position in the plot. A comparison with relative levels of barium and zirconium in late Hellenistic and early Roman vessel glasses shows a clear distinction from many Satricum samples, especially for

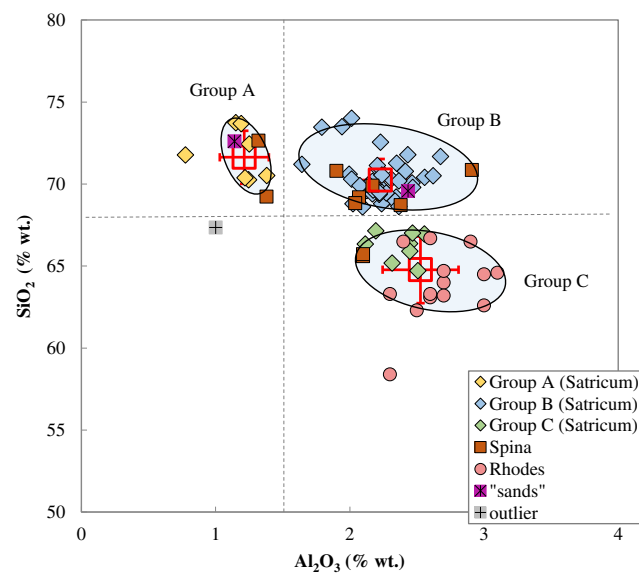


Fig. 3 Al_2O_3 versus SiO_2 concentrations in glasses from Satricum (this study), Spina and Rhodes samples. The samples are divided in 3 main groups (group A high SiO_2 /low Al_2O_3 , group B high SiO_2 /high Al_2O_3 and group C low SiO_2 /high Al_2O_3). In the same plot, mean values with 1 s.d. of each group are presented as well as the concentrations of two hypothetical glasses using sand suitable for glassmaking (Brems et al. 2012)

Table 5 Major and Minor oxides detected by SEM-EDX (Sa.1-Sa.45) and EPMA (SAT.1-SAT.11) including min. max. mean and standard deviation (s.d.) values (in % w.t.)

Sample	Inventory No.	Colour	Type	Group	Na ₂ O	MgO	Al ₂ O ₃	SiO ₂	SO ₃	Cl	K ₂ O	CaO	Fe ₂ O ₃
Sa.4	CAT.7	Dark blue	Unidentified	A	19.11	0.63	1.25	72.45	0.18	1.08	0.29	4.66	0.35
Sa.5	CAT.9	turquoise	Alabastron	A	18.23	0.57	1.25	70.26	0.51	1.05	0.33	7.13	0.67
Sa.16	CAT.52	Dark blue	Alabastron	A	16.69	0.57	1.15	73.74	0.18	1.17	0.29	5.67	0.55
Sa.21	CAT.64	turquoise	Alabastron	A	17.92	0.48	1.22	70.38	0.46	1.06	0.27	7.48	0.72
Sa.35	CAT.94	Dark blue	Aryballos	A	16.37	0.63	1.19	73.69	0.23	1.19	0.4	5.72	0.57
Sa.40	CAT.104	Brown	Oinochoe	A	17.91	0.62	0.78	71.77	0.46	1.35	0.27	6.51	0.34
SAT.5	V565	Blue	Unidentified	A	16.51	0.59	1.38	70.52	0.17	1.22	0.34	7.15	0.9
Sa.1	CAT.1	Dark blue	Alabastron	B	16.45	0.32	1.79	73.47	0.37	1.05	0.62	5.05	0.95
Sa.2	CAT.4	Dark blue	Alabastron	B	16.21	0.44	2	70.6	0.36	1.23	0.48	7.95	0.75
Sa.3	CAT.5	Dark blue	Alabastron	B	16.62	0.63	2.47	70.07	0.14	1.21	0.39	7.65	0.84
Sa.6	CAT.11	Dark blue	Alabastron	B	19.16	0.73	2.25	69.68	0.27	1.17	0.29	6.07	0.57
Sa.7	CAT.13	Dark blue	Alabastron	B	17.47	0.53	2.13	69.6	0.2	1.33	0.5	7.43	0.81
Sa.8	CAT.15	Dark blue	Alabastron	B	17.71	0.58	2.24	69.49	0.24	1.42	0.48	7.19	0.77
Sa.9	CAT.16	Dark blue	Alabastron	B	17.83	0.64	2.28	69.13	0.26	1.36	0.53	7.32	0.72
Sa.10	CAT.36	Dark blue	Alabastron	B	17.1	0.36	2.35	71.28	0.39	0.96	0.56	6.27	0.73
Sa.11	CAT.40	Dark blue	Alabastron	B	15.91	0.67	2.42	70.78	0.16	1.27	0.44	7.54	0.83
Sa.12	CAT.42	Dark blue	Alabastron	B	16.94	0.53	2.31	70.31	0.22	1.17	0.44	7.18	0.9
Sa.13	CAT.44	Dark blue	Alabastron	B	17.01	0.6	2.37	70.19	0.29	1.15	0.4	7.23	0.76
Sa.14	CAT.45	Dark blue	Alabastron	B	17.22	0.93	2.21	70.91	0.29	1.01	0.62	6.01	0.8
Sa.15	CAT.50	Dark blue	Alabastron	B	16.7	0.42	2.01	70.3	0.17	1.19	0.48	8.16	0.61
Sa.17	CAT.54	Dark blue	Alabastron	B	16.6	0.38	1.94	73.49	0.4	0.91	0.59	4.79	0.82
Sa.18	CAT.55	Dark blue	Alabastron	B	16.83	0.59	2.15	70.08	0.27	1.27	0.45	7.64	0.77
Sa.19	CAT.57	Dark blue	Alabastron	B	19.43	0.69	2.35	69.22	0.15	1.17	0.36	6.12	0.53
Sa.20	CAT.58	Dark blue	Alabastron	B	19.52	0.66	2.34	68.96	0.22	1.25	0.42	6.08	0.57
Sa.22	CAT.66	Dark blue	Alabastron	B	21.53	0.76	2.37	68.64	0.24	0.99	0.51	4.7	0.26
Sa.23	CAT.68	Dark blue	Alabastron	B	16.53	0.3	2.01	74.01	0.47	0.77	0.49	4.84	0.58
Sa.24	CAT.72	Dark blue	Alabastron	B	17.74	0.58	2.17	70.17	0.19	1.13	0.46	7.16	0.44
Sa.25	CAT.76	Dark blue	Amphoriskos	B	20.17	0.65	2.24	68.81	0.21	1.22	0.51	5.86	0.34
Sa.26	CAT.78	Dark blue	Amphoriskos	B	17.63	0.66	2.27	69.46	0.25	1.28	0.46	7.24	0.76
Sa.27	CAT.79	Dark blue	Amphoriskos	B	17.57	0.54	2.22	69.37	0.26	1.33	0.45	7.26	0.79
Sa.28	CAT.81	Brown	Amphoriskos	B	18.58	0.5	2.1	68.61	0.3	1.47	0.55	7.55	0.34
Sa.29	CAT.82	Brown	Amphoriskos	B	17.61	0.55	2.17	69.39	0.22	1.43	0.63	7.72	0.27
Sa.30	CAT.83	Brown	Amphoriskos	B	18.22	0.56	2.02	68.81	0.33	1.54	0.54	7.74	0.24
Sa.31	CAT.85	Dark blue	Hydriskos	B	17.72	0.61	2.21	69.49	0.11	1.42	0.42	7.24	0.84
Sa.32	CAT.87	Dark blue	Hydriskos	B	18.57	0.68	1.64	71.2	0.23	1.2	0.36	5.57	0.55
Sa.33	CAT.89	Dark blue	Hydriskos	B	17.74	0.54	2.14	69.97	0.21	1.41	0.54	6.69	0.76
Sa.34	CAT.92	Dark blue	Hydriskos	B	17.03	0.5	2.21	71.14	0.3	1.15	0.47	6.54	0.66
Sa.36	CAT.98	Dark blue	Oinochoe	B	16.99	0.76	2.56	70.41	0.49	0.83	0.54	6.88	0.53
Sa.37	CAT.99	Dark blue	Oinochoe	B	17.99	0.72	2.17	69.63	0.49	1.05	0.58	6.64	0.73
Sa.38	CAT.101	Dark blue	Oinochoe	B	18.16	0.6	2.08	69.91	0.57	0.97	0.59	6.46	0.64
Sa.39	CAT.102	Dark blue	Oinochoe	B	16.85	0.72	2.62	70.51	0.45	0.73	0.53	6.95	0.65
Sa.41	CAT.105	Dark blue	Oinochoe	B	16.1	0.33	2.43	71.8	0.45	1.04	0.5	6.91	0.45
Sa.42	CAT.107	Dark blue	Oinochoe	B	16.44	0.44	2.23	72.55	0.36	1.03	0.51	6.17	0.35
Sa.43	V1091	Dark blue	Bead	B	17.23	0.56	2.68	71.69	0.19	1.02	0.33	5.23	1.09
Sa.44	Tomb 29a (V75)	Dark blue	Bead	B	18.66	0.52	2.22	69.47	0.38	1.31	0.4	6.74	0.3
Sa.45	Tomb 19 (V71)	Dark blue	Bead	B	17.79	0.59	2.24	70.55	0.41	0.89	0.58	5.42	1.53
SAT.2	V2088	Dark blue	Oinochoe	B	16.15	0.4	2.47	69.79	0.29	1.01	0.49	7.18	0.67
SAT.1	V528	Dark blue	Amphoriskos	C	16.83	0.59	2.45	66.38	0.18	1.31	0.41	8.76	1.22

Table 5 (continued)

Sample	Inventory No.	Colour	Type	Group	Na ₂ O	MgO	Al ₂ O ₃	SiO ₂	SO ₃	Cl	K ₂ O	CaO	Fe ₂ O ₃
SAT.3	V22718	Dark blue	Unidentified	C	17.22	0.62	2.19	67.14	0.36	0.92	0.59	8.1	1.03
SAT.4	V2160	Dark blue	Alabastron	C	16.87	0.62	2.45	65.91	0.18	1.18	0.39	8.88	1.17
SAT.6	V730	Dark blue	Unidentified	C	17.55	0.41	2.56	66.95	0.3	1.12	0.37	8.47	0.83
SAT.7	V2331	Dark blue	Unidentified	C	17.6	0.64	2.12	66.35	0.37	1.02	0.6	8.5	0.96
SAT.8	V525	Dark blue	Unidentified	C	17.78	0.67	2.47	67.01	0.22	1.32	0.45	8.79	0.9
SAT.9	V10443	Dark blue	Unidentified	C	17.84	0.41	2.51	64.72	0.37	0.96	0.48	6.89	0.93
SAT.10	V2537	Dark blue	Unidentified	C	18.7	0.57	2.32	65.19	0.24	1.46	0.52	8.75	0.3
SAT.11	V759	Dark blue	Unidentified	Outlier	15.76	0.52	1	67.36	0.28	1.03	0.35	8.62	1.01
Min					15.76	0.3	0.78	64.72	0.11	0.73	0.27	4.66	0.24
Max					21.53	0.93	2.68	74.01	0.57	1.54	0.63	8.88	1.53
Mean					17.55	0.57	2.1	69.87	0.29	1.16	0.46	6.9	0.7
s.d.					1.1	0.12	0.44	2.09	0.11	0.18	0.1	1.12	0.26

Satricum samples containing below c. 200 mg/kg (Thirion-Merle 2005; Henderson 2013, Fig. 8.3).

Among the 56 glass samples discussed in this paper, there are 53 fragments of glass vessels and 3 fragments of glass beads, the balance being sub-samples from the same objects of different colours. The 53 fragments are dark blue (47), brown (4) and turquoise (2); the 3 bead samples are dark blue. The deep blue colour, in all the vessel and bead samples, is associated with the presence of cobalt (Co) and copper (Cu) in varying concentrations, while the turquoise colour in 2 fragments is associated with high Cu levels. The 4 brown samples do not show anomalous values in any of the colourant elements so it can be assumed that their colouration is due to controlling the furnace atmosphere and therefore will be excluded from the following analysis.

As it can be seen in Fig. 7, the majority of samples have elevated values of both Co and Cu suggesting that they belong at the CoCu cobalt colouring glass category identified by various scholars for earlier glass samples (Shortland and Eremin 2006; Shortland et al. 2007; Smirniou and Rehren 2013). They are positively correlated with a Co:Cu ratio close to 1. There is also a cluster of samples (yellow ellipse) (Sa.14, Sa.36, Sa.37, Sa.38 and Sa.39) which have lower Co values. Also, samples Sa.5, Sa.21 show low Co (91 and 86 mg/kg) and high Cu contents (5542 and 5307 mg/kg) which is to be expected since they are turquoise. Five samples (Sa.4, Sa.16, Sa.41, Sa.42 and Sa.45) can be considered as Co-blue glasses since their Cu values are below 850 mg/kg, as suggested by Smirniou and Rehren (2013). The two glass beads (samples Sa.43–44) show high Cu content (above 3000 mg/kg) and especially Sa.44 show high Co content (2683 mg/kg). The two beads, as it is seen on Fig. 7, do not fall in the same CoCu line like the majority of samples. Triantafyllidis (2001) has suggested that glass used to make beads was often recycled. Furthermore, Pliny the Elder in Natural History (XXXVI. 199) writes about recycling activities associated

with the manufacture of glass beads. Therefore, the glass used for these two beads could be a result of recycling.

There are various minerals rich in cobalt, that have been proposed as colourant sources in ancient glasses, such as cobaltite (CoAsS), absolane (a mixture of MnO and CoOOH), trianite (2Co₂O.CuO.6H₂O) and skutterudite ((Co,Ni,Fe)As₃) (Henderson 2013). Another common source of cobalt is cobaltiferous alums which are found mainly in Egypt but there are also alum ores in Iran, Turkey and Germany as well (Kazmarczyk 1986; Henderson 2013). Therefore, elements that can be associated with either minerals or alums include Al, Mn, Ni, Zn, As, Fe. In Fig. 8, the correlation between Mn and Co for vessels and beads of deep blue and turquoise colours is presented.

According to the plot, three major groups of samples can be distinguished, while there are five samples which are “outliers” (falling in the upper, lower and left part of the plot). The first group ($n = 9$) has low Mn values (between 100 and 200 mg/kg) and variable Co values (700–1550 mg/kg). The majority of the samples ($n = 19$) fall in the centre of the plot having elevated values of both Co and Mn while the third group ($n = 5$) has low Co and high Mn contents. We can show that three different cobalt sources were used for the coloration of these samples.

Concerning the “outliers”, all of them are group A samples except from the blue bead (Sa.44), which has high Mn content (2341 mg/kg) and elevated amount of Co (2683 mg/kg Co). Combined with the fact that it has a notable level of Cu (4495 mg/kg), its composition can be considered to be a result of recycling. Among group A samples, there are two turquoise samples (Sa.21 and Sa.5) having low Co values (their coloration is due to high levels of Cu: 5307 and 5542 mg/kg, respectively) and lastly there are two samples with the highest amount of Mn (Sa.4 and Sa.16), which is another indication that group A samples were manufactured not only with different primary raw materials (sand, see Fig. 3) but also with

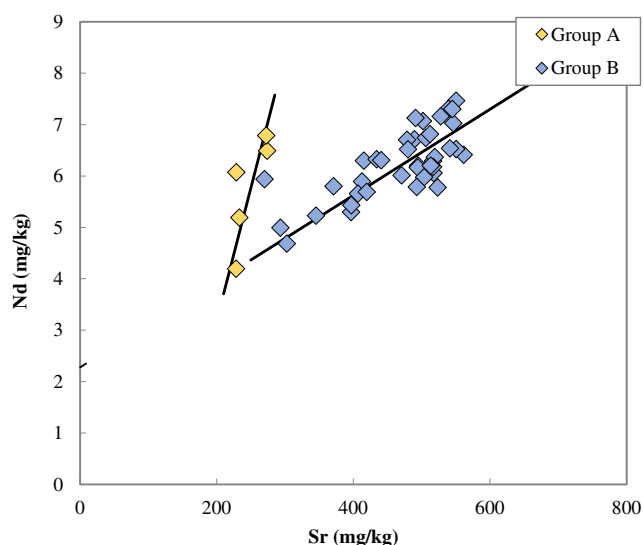


Fig. 4 Nd versus Sr concentrations in 45 Satricum glass samples. The samples are divided in two groups with differing slopes which were calculated by maximum likelihood functions. The first group consists of six samples (part of group A samples see Fig. 3) and the second with the rest (group B samples see Fig. 3)

different secondary additives, indicating a totally different glassmaking tradition.

Excluding the outliers, if we plot the rest on a ternary diagram, the correlation between Mn, Co and Ni becomes clear. In this diagram, we can note that the samples fall into three different groups with low, middle and high values of Mn. The high and low Mn samples seem to coincide and fall almost precisely into the two groups that Abe et al. (2012) have identified (the yellow ellipse in Fig. 9). The yellow ellipse in the middle of the graph is for samples coming from Egypt of the eighteenth Dynasty and therefore we can assume they have a totally different source of Co. According to Abe et al. (2012),

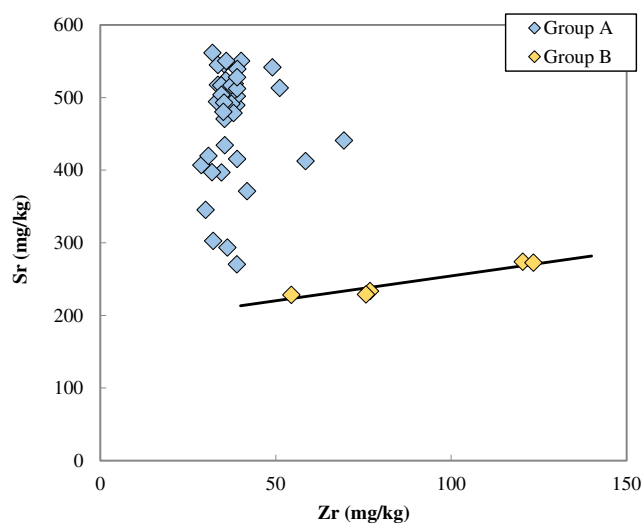


Fig. 5 Zr versus Sr concentrations in Satricum glass samples. The samples are divided in two groups with high and low Zr levels and/or Sr levels. The samples with the positive correlation are part of group A samples (see Fig. 3)

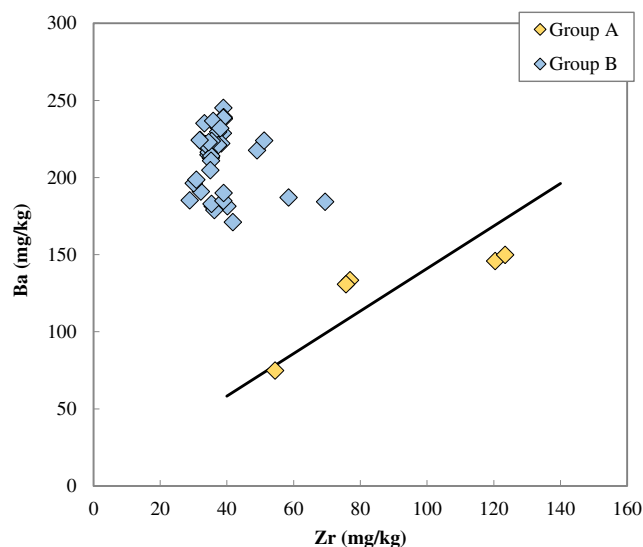


Fig. 6 Zr versus Ba concentrations in Satricum glass samples. The samples are divided in two groups with high and low Ba levels. The samples with the positive correlation are part of group A samples (see Fig. 3). Colourants

Co in the lower group of samples is derived from Mn-rich cobaltiferous ores such as asbolane which was mined at Iran. It is rather unclear what the Co source of the other two groups (with middle and low Mn content) is but cobaltiferous alums from the Dakhla and Kharga oases in Egypt should be excluded since they show different chemical compositions in these trace elements (Shortland et al. 2006). Furthermore, differences in Mn values of cobalt blue glasses were also noted by Henderson (2000), where the chemical analyses of European Iron Age blue glass revealed a change in Co source in the second c. BC, from a Sb-rich source to a new Mn-rich source. In our assemblage, there is no correlation between Sb and Co,

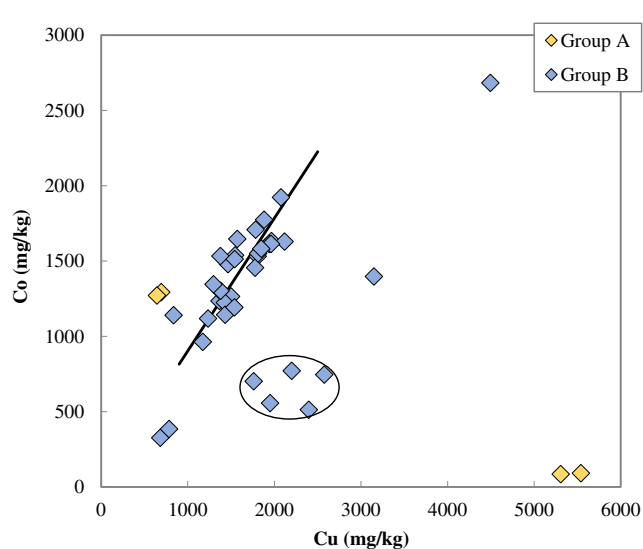


Fig. 7 Cu versus Co concentrations in Satricum glass samples. The majority of Satricum samples are positively correlated (blue diamonds), and they are coloured with the Cu and Co

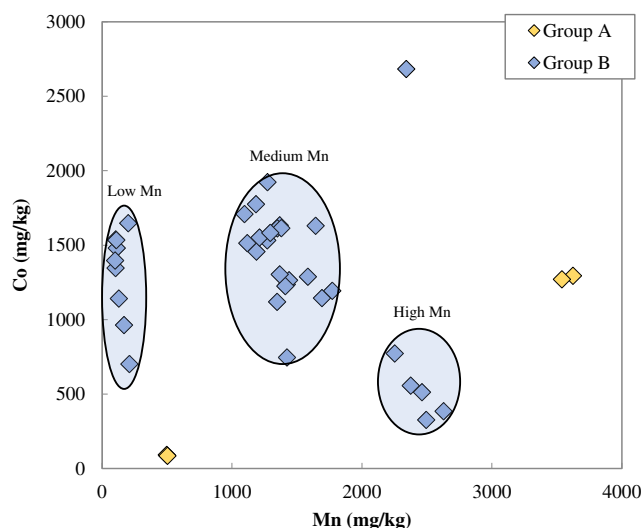


Fig. 8 Mn versus Co concentrations in Satricum glass samples. There are three distinguishable groups from left to right with low Mn high Co, high Mn high Co and high Mn low Co content

and we can assume that Sb was introduced in the glass as an opacifying agent rather than as an impurity of the Co source.

Discussion

From our previous discussion of the basic compositions of Italian glasses and the colourants used, we can extract useful conclusions concerning their technology. From the analysis of major minor and trace elements, we can suggest a model of production that involves the manufacture of the base glass and the coloration of it as two different processes. Therefore, we

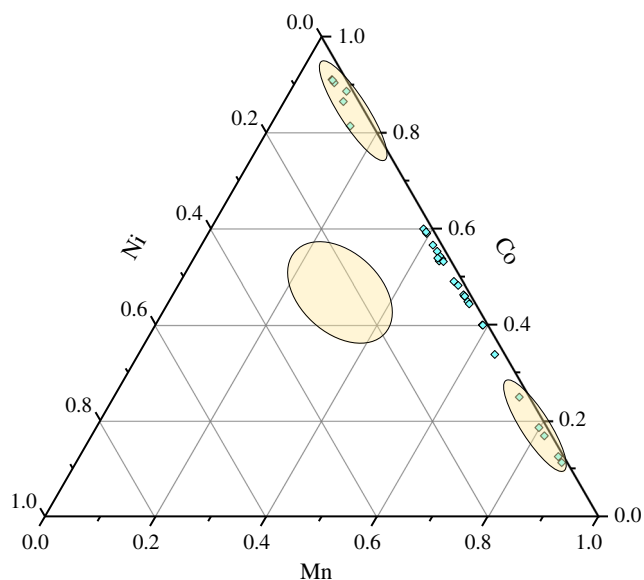


Fig. 9 Ternary plot of Mn, Co and Ni in cobalt blue glasses. Satricum samples do not correlate with Egyptian coeval samples (yellow ellipse in the middle) (Abe et al. 2012). A third group of samples exists between the two elongated ellipses suggesting a possible third source of Co

can suggest a two-stage procedure. In the first stage, the base glass was fused using sand and natron as an alkali source. In the second stage, which probably formed part of a secondary glassmaking process when the base glass was reheated, colourants were mixed with the base glass; the glass vessels were then created (Fig. 10). This can be suggested because all group B samples can be divided into 3 broad groups depending on their Mn content which is associated with the colouring agents, mainly Co. Even if manganese oxide is present in the base glass, the use of variable Mn impurities in different cobalt-bearing minerals would produce the compositional variations that we have observed.

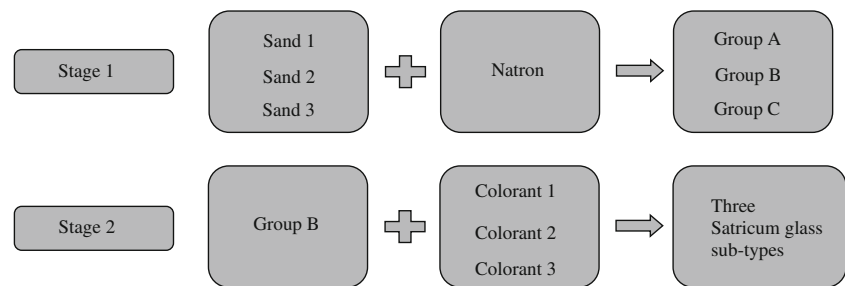
In the first stage, three different base glass compositions can be distinguished according to their relative Al_2O_3 and SiO_2 levels, as discussed in relation to Fig. 3. Therefore, we probably have evidence for the use of three different sand sources and a common natron source. Group C samples have similar values to those found in Rhodian glass and, therefore, the silica can be considered as deriving from the same origin as Rhodian samples. The island of Rhodes Island played an important role as a glassmaking centre of core-formed vessels (especially Mediterranean core-formed group I vessels) during the late sixth and fifth c. BC (Harden 1981; Grose 1989; Triantafyllidis 2003). The continuity of the core-formed industry in the succeeding centuries can be attested by various finds (products of incomplete firing) (Weinberg 1966; Triantafyllidis 2000a, b). Therefore, group C samples can be either attributed to a “Rhodian” manufacturing centre or one that produced the glass that was used to make Rhodian vessels. It was certainly a different manufacturing centre from our group A and B samples.

It is quite interesting that 3 hypothetical glasses, manufactured from sands suitable for glassmaking of Roman period, have common characteristics with group A and B samples—and these are plotted with the data in Fig. 3. The concentration of these hypothetical glasses was calculated by Brems et al. (2012). Taking into consideration that the source of sand probably did not change significantly in the era after the Mediterranean Group II industry, we could hypothesise that group B vessels were manufactured most likely with sands coming from Italy, especially sands from Salentina peninsula (South Italy) and/or Tuscany (Brems et al. 2012).

This suggestion fits well with the archaeological record according to which the provenance of most of the Mediterranean group II vessels is southern Italy (Harden 1981; Grose 1989; Stern and Schlick-Nolte 1994). However, alternative regions in mainland Greece, such as Macedonia, have been suggested (McClellan 1984).

Supporting evidence for this conclusion is that Mediterranean group II vessels are found far more frequently in Italy and Magna Graecia than in the Aegean, Egypt, the Levant and Western Asia (Grose 1989). Our group B samples form a consistent group of 40 samples with the majority of

Fig. 10 A proposed model for the manufacturing processes of Satricum samples



them being a dark blue colour. We could hypothesise the same for group A samples since the Italian “sand” has the same chemical characteristics. Group A samples though do not form a homogeneous group; two samples have turquoise colour (Sa.5, Sa.21), 1 is brown (Sa.40), 4 are blue (Sa.4, Sa.16, Sa.35, SAT.5) and 1 (a kohl tube) that comes from Spina is attributed to Egyptian origin according to Arletti et al. (2011), while all of them have totally different decorative patterns to the samples that fall into group B. Therefore, even though group A samples share common chemical characteristics, due to their visual inconsistency and their small number, it seems more likely they were imported to Italy.

Turning to colourants, group B samples are coloured using Co. The analytical results suggest that either they use 3 different sources or most likely they use the same Co source which contains variable proportions of minerals. The alum deposits in Egypt (Dakhla and Kharga oases) should be excluded because they have different trace element concentrations, such as the levels of Ni, Mn and Co; the most likely source of Co should be Iran. There is an Iranian Co deposit which occurs at Qamsar, southwest of Kashan, being exploited as late as the 1960s can be suggested. This deposit contains various copper and cobalt minerals such as erythrite ($\text{Co}_3(\text{AsO}_4)_2 \cdot 8\text{H}_2\text{O}$), nickel sulphides, spinel ($(\text{Co}, \text{Ni}, \text{Fe},)_3\text{S}_4$) and absolane (Henderson 2013, 72) which can be linked to our three different (sub-) groups according to Mn levels and the positive correlation between Co and As (not shown here). Therefore, high Mn samples can be attributed to the use of an absolane mineral while the rest can be attributed to a mixture of absolane (for the medium Mn samples) and/or erythrite (for the low Mn samples).

Conclusions

The analysed samples from Satricum show interesting technological differences not only between the samples analysed but also with coeval glass artefacts (Arletti et al. 2011; Triantafyllidis et al. 2012).

Satricum samples can be differentiated into three groups based on the relative elemental concentrations associated with minerals in the sands used to make the glasses indicating that different sand sources were used.

One of the groups (group A) can be considered as being imported from a currently unattributed source; a second group (group B) has characteristics consistent with an Italian origin; a third set of samples (group C) can probably be attributed to a Rhodian origin. There is a strong indication that group B samples were manufactured somewhere in Italy taking into account both archaeological and scientific/technological factors. Even though there is not any archaeological evidence for primary glass production during the Hellenistic period in Italy, certain scholars believe that production of core-formed vessels type II (after Grose 1989) took place somewhere in southern Italy. This interpretation combined with the fact that glass samples have similar compositions with Italian sands suitable for glassmaking provide a solid indication for a production in this region or/and in Southern Italy.

A two-stage manufacturing process can be suggested for Satricum G group B samples. In the first stage, the basic glass composition would have been produced and in the second stage colourants were added to the glass melt. The majority of samples, as characterised by trace elements, have a common cobalt colourant source which can probably be located in Iran especially in central Iran, rather than Egypt, while the outliers exhibit totally different trace element characteristics enhancing the belief that they were manufactured using a totally different set of raw materials or/and a different glass technology.

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