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First insights into the archaeometric analysis of the *Los Amores* Mosaic in Cástulo (Linares, Spain): the Judgement of Paris

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Abstract

This paper discusses results obtained from in situ analysis of the tesserae of the Roman mosaic of Los Amores (Cástulo site, Linares, Spain) dating back to the turn of the 1st to the 2nd century AD. Specifically, it focuses on the scene The Judgment of Paris. In view of the exceptional state of preservation of the mosaic, from which very few tesserae had fallen off, non-invasive methods with portable Micro Raman Spectroscopy (MRS) and hand-held X-ray fluorescence (hXRF) and data assessment by use of principal component analysis and binary representations were selected. The results obtained allow to evaluate both the analytical method and the portable equipment used, as well as to classify the raw materials, the colouring agents and the opacifiers used. MRS analysis proved crucial for the identification of stone tesserae (ironstones, carbonate and siliciclastic rocks) and for the identification of the type of glasses used (soda-lime-silicate and lead type glasses) based on the analysis of two detached tesserae. hXRF analysis of the glass tesserae identified both colouring agents (Co, Cu, Pb, Zn) and opacifiers (calcium antimonate). The data obtained lend themselves to an assessment of the degradation process that threaten the integrity of the mosaic. The identification of tessera made of specific stone materials (especially ironstone) and of lead glass tesserae suggest the existence of a mosaic workshop in the Upper Guadalquivir (Eastern Andalusia, Spain).

Keywords: Roman, Mosaic, Cástulo, In situ, Portable MRS, hXRF

Introduction

The Cástulo Archaeological Zone is located on the right bank of the River Guadalimar, 5 km from the city of Linares (Spain) (Additional file 1: Figure S1). During its Ibero-Roman phase Cástulo would have been one of the ancient capitals of the southern Iberian Peninsula, as attested both by the size of its walled area (50 ha) and its strategic position at the head of the Guadalquivir Valley. Cástulo has a history of more than four thousand years spanning from the Chalcolithic to the fifteenth century AD. It is a place in which diverse native cultures

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Punic War in 206 BC, it was definitively conquered by Rome under the general Publius Cornelius Scipio. From then on it was part of the Roman Empire until the crisis that began in the third century AD led to its fall. The Roman town attained considerable splendour and importance in the late first century AD. It had a theatre adorned with statues, baths, latrines, an aqueduct, a forum and an amphitheatre in which gladiator spectacles were held. Cástulo also built a water supply network that included aqueducts, pipes, tanks and fountains. This period of great splendour in Cástulo during the early Roman period contrasts with the decline it suffered in parallel with the general crisis in



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the Empire during the third century AD [1, 2]. The city rose from its ashes in the fourth and fifth centuries AD [3] and was inhabited during the Islamic occupation and after the Castilian conquest at the beginning of the twelfth century AD. After several attempts to repopulate it, the town was definitively abandoned in the mid-fifteenth century AD [1, 2].

After several decades of research inactivity, annual excavations with a notable methodological renovation [4] resumed in 2011. Thanks to this, the significance of Cástulo from an archaeological, artistic and tourism point of view has been reactivated. This aspect is very important as the archaeological site is one of the main tourism and economic assets of the town of Linares, which has been immersed for many years in a profound economic depression.

There can be no doubt that one of the most spectacular outcomes of this new activity was the excavation of the so-called Building D, a Roman-period building of approximately 33×12 m dated to the late first century AD [5, 6]. It was built in honour of Emperor Domitian (51-96 AD) and would have been a public building. Its construction would have been part of an imperial project, a representation of the *maiestas imperii*, and also of the emperor himself as a representative of them. The excavations begun in 2011 have so far identified ten rooms in Building D (Additional file 1: Figure S1), several of them decorated with mosaics and mural paintings [7]. The most outstanding is Room 1 with its splendid decoration known as the Los Amores Mosaic (Additional file 1: Figures S2) [6, 8]. The 11.65-m-long, 5.75-m-wide mosaic is almost completely preserved. Its composition is a variation on the so-called compass or *a oculi* scheme, incorporating two central circles instead of the more normal one [6]. The origin of the compass scheme can be found in Italy, in black and white designs in Pompeii, Ostia and Lucera. From Italy it would have been disseminated to the provincial workshops in the rest of the empire, including Hispania, in the late first century AD [9].

The right-hand circle contains a depiction of the Judgement of Paris (Fig. 1) and that on the left the myth of Selene and Endymion. The half circles on the longest sides are decorated with four Erotes in profile offering bunches of grapes to a partridge and a pheasant. The other two half circles on the shorter sides are also illustrated with Erotes, although in this case they are hunting a hare. The quarter circles in the corners contain allegoric busts of the seasons with their characteristic attributes. They represent the passage of time and annual fertility, a theme that was not just philosophical but also linked to wellbeing. This was a very common subject in Roman mosaics and was particularly prevalent in the *Baetica* region. The six oblong quadrangular spaces are occupied by mammalian, herbivorous and carnivorous animals in an attitude of running through a rocky landscape: on one side a wild boar, a lion and a horse, and on the other, a tigress, a deer and a lioness.

In general, the Cástulo mosaic has a dual significance. On the one hand it has been suggested that the figurative scenes have a moralising content linked to the excesses of love and the imbibing of wine [6]. On the other, it would have formed part of an architectural and iconographic discourse related to the legitimation of power and the consolidation of the legitimate, continuous and eternal nature of Roman and imperial domination. In this respect, the presence of the Judgement of Paris connects the origin of Rome to the history of Troy, as, according to one Roman legend, the foundation of the former was linked to Trojan hero Aeneas [10, 11]. This second interpretation would be in keeping with the building in which it was found. All the evidence points to this having had a public, propagandistic and legitimating function: its size and internal structure, the quality of the materials used, the presence of quality mosaics and mural decorations, the absence of productive and/or domestic activities, the location near the urban centre, and its unique nature in the town of Cástulo [5].

Aims

In 2018, an archaeometric analysis of the *Los Amores* Mosaic was undertaken as part of the "Cástulo: Archaeometric Analysis and Social Transfer" Research Project. From all the scenes, the Judgement of Paris was chosen as the most representative for demonstrating the initial results of the project and to evaluate the mosaic's main characteristics. Thus, the objectives proposed with this study are two-fold.

Firstly, given the artistic, archaeological and historical importance of the *Los Amores* Mosaic and the scene of the Judgement of Paris, the first objective proposed was the archaeometric analysis of the tesserae. This research can be added to the stylistic, iconographic and historical research already undertaken [5, 6, 12, 13], thus completing the comprehensive study of the mosaic. In this respect, the mineral and elemental analysis should conclude with the spectroscopic characterisation of the tesserae to identify the raw materials (stone, glass, ceramic), colouring agents and opacifiers. This study should also provide fundamental information for evaluating the state of the mosaic's conservation and the possible maintenance and restoration measures.

Depictions of the Judgement of Paris are documented in only two mosaics in Hispania: one in the villa of El Alcaparral, Casariche (Seville) [14] and the other in the villa of Noheda (Cuenca), both from the fourth century



AD [15]. The scene also appears in the mosaics of the Atrium House in Antioch on the Orontes (Turkey) from the first half of the second century AD; in the baths on the island of Kos (Greece) from the late second and early third centuries AD [16]; in the colony of *Ulpia Traiana Augusta Dacica Sarmizegetusa* (Romania) from the second-third centuries AD; and in Caesarea (Algeria) from the fourth century AD. Of all of them, only the Noheda mosaic has been subjected to an archaeometric analysis, although that focused solely on the glass tesserae [17, 18].

The methodological dimension of the second objective was aimed at exploring the capacity and enhancing the use of portable equipment in the *in-situ* analysis of mosaics. Given the undesirability of disturbing them by removing a significant repertory of the tesserae for study, non-invasive strategies need to be developed to analyse them in situ. Thus, this proposal comprises the use of portable equipment able to carry out readings of the mineral (portable MRS) and elemental (handheld XRF) compositions. A joint analysis undertaken with both techniques allows us to analyse both stone and glass tesserae with greater guarantees.

The spectroscopic techniques selected have been used frequently and successfully on Roman mosaic tesserae from all over the Mediterranean, although in those cases by analysing separate tesserae with high precision laboratory equipment [19–23]. In contrast, the combined application of portable equipment to analyse mosaics has so far been infrequent [24–26]. Nevertheless, it is necessary because in many cases it is the only option for undertaking a comprehensive study of a mosaic. *Los Amores* Mosaic is an excellent example of this situation. Our investigation also had the advantage of preliminary studies in which the effectiveness of various types of portable Raman and hXRF equipment were tested, with the specific portable devices used showing good applicability and performance [27, 28].

Experimental

Sampling

Prior to the identification of the colours and the analysis, the tesserae were treated by removing all the dust with a soft bristle brush and cleaning the surface with a solution of water and methanol. Following this operation, the tesserae were identified on the mosaic and selected according to their colour. A high-resolution gigapixel image available in GigaPan was also used as a complementary tool [29]. We analysed 92 tesserae from The Judgment of Paris scene using portable MRS and hXRF instruments (Additional file 1: Figures S3–S7). The tesserae were chosen to account for all the colours and chromatic variations (Fig. 2) in the scene. This large number allowed us to minimise identification and measurement errors caused by the difficulty of accurately directing the laser and/or X-ray beam over the mosaic surface.

In addition to these 92 samples, two loose vitreous tesserae found during the archaeological excavation of the Paris mosaic were further analysed using MRS laboratory equipment, with the aim of completing the analytical characterisation (Fig. 3).

Micro Raman spectroscopy

All the equipment and the experimental parameters are as described in Rousaki et al. [27], Tuñón et al. [30] and Sánchez et al. [31].

The portable equipment used was a BWS445-785S innoRam[™] Raman spectrometer (B&WTEK, Inc., Newark, USA) with a 785 nm excitation laser (maximum power of 300 mW) and a ~4 cm⁻¹ spectral resolution. The spectrometer was attached to a TE-cooled, back-thinned, 2D binning CCD detector. For measuring the tesserae from the *Los Amores* Mosaic, a 1.5-m fibre optic probe was focused by hand. The lens used had a 5.9-mm working distance regulator that ensured proper focusing while minimising contamination and



Fig. 2 Colours analysed in the Judgement of Paris and Munsell classification. a White 2.5Y 8/0. b, c Greys N 6/0, N 5/0. d Pinkish-beige 2.5YR 7/4. e, f Reds 7.5R 4/8, 7.5R 3/3–3/4. g Black N 1.7/0. h, i Blues 5 PB 5/10, 5 PB 8/2. j–I Turquoise-green 5 PB 8/4, 7.5B 7/8, 5BG 6/4. m Orange 5YR 7/8. n, o Yellows 10YR 6/6–7/6, 5Y 9/8



degradation of the lens material. The spot size was 85 μ m. The laser power was kept low and the measurement time and accumulations were adapted in order to acquire acceptable signal-to-noise (S/N) ratio. No special calibration was applied. Before each measurement, a dark scan was executed to improve the S/N ratio in the Raman spectra. The experimental conditions were exposure time ranging from 100 to 1000 ms, maximum power of 40%, maximum 60 acquisitions and a spectral range of between 65 and 3000 cm⁻¹. This equipment was powered both by a Pb-ion battery (MICROBEAM S.A., Barcelona, Spain) and a portable generator. Raman spectra were recorded using the BWSpecTM 3.26 software version (B&W Tek Inc.).

For the analysis of the loose tesserae, the laboratory equipment used was a Renishaw inVia Qontor Spectrometer (University of Málaga, SCAI-UMA) coupled with a Leica confocal microscope equipped with three lasers: solid state (473 nm, 8 mW on sample), Nd:YAG (532 nm, 27 mW on sample) and diode (785 nm, 140 mW on sample). The spectrometer was attached to a Peltier-cooled CCD detector. This device consisted of two gratings of 2400 l/mm for the 473-nm and 532-nm lasers and 1200 l/mm for the 785-nm laser. The spot size was ~1 µm. The spectra were acquired using the $50 \times$ objective lens in the 100–3000 cm⁻¹ region with a spectral resolution of ~1 cm⁻¹. Acquisition time was set at between 1 and 20 s per accumulation and the maximum number of accumulations was 20.

The interpretation of the spectra was carried out by comparison with related literature, freely available RRUFF and Raman databases [32–35].

X-Ray Fluorescence (hXRF)

The equipment and the experimental parameters are as described in Costa et al. [28]. The in-situ analyses were performed with an Olympus Innov X Delta Premium commercial instrument coupled with a rhodium (Rh) anode-based X-ray tube and a 20-mm² silicon drift detector (SDD). An aluminium (Al) filter was applied for measuring the higher Z-elements (from Al onwards) with a voltage of 40 kV and a current of 38.7 µA. Without filtering (low-Z elements), a voltage of 10 kV and a current of 50.8 µA can be applied. The measurements were performed in air for 150 s (live time) with the Geochem mode of the instrument. All measurements were performed with a collimated polychromatic X-ray beam for excitation $(3 \times 3 \text{ mm}^2)$. This equipment also had a camera that allowed the correct positioning of the instrument and viewing of the analysed area. The calibration of the spectrometer was automatic using an alloy 316 stainless steel. Each sample was analysed in one location. The portable equipment was powered by removable Li-ion batteries. The collected spectral data were processed using the dedicated XRF spectrum evaluation software AXIL (Analysis of X-ray Spectra by Iterative Least Squares) [36, 37]. The hXRF values shown in this

work mean the raw net peak intensity of K α and L α lines of detected elements in XRF spectra.

Data analysis

To obtain an objective classification of the tesserae, a principal component analysis (PCA) was carried out using the hXRF data. All the PCA analyses were run on the correlation matrix of the raw net counts of K α and L α lines of the detected elements using the statistical tools from OriginPro v.2019 (OriginLab Corp). Previously to statistical analysis, the elements values were scaled and normalised. Scores were standardised by default with a correlation matrix using OriginPro. Those elements that were detected only in a few tesserae have not been included in the PCA.

The first analysis of all 82 tesserae was performed on the data from 14 elements (Mg, Al, Si, P, Cl, K, Ca, Ti, Mn, Fe, Cu, Zn, Sr, Pb). Principal components PC 1 and PC 2, with eigenvalues of 5.18 (percentage of variance: 37.02%) and 2.61 (percentage of variance: 18.66%), account for an explained variance of 55.67% and were represented on scores and loading plots graphs. Based on the results of this PCA another two were carried.

The second PCA was performed on the 35 stone tesserae using data from the 15 most common elements in the characterisation of this type of tessera [26] (Mg, Al, Si, P, S, Cl, K, Ca, Ti, Mn, Fe, Cu, Zn, Sr, Pb). Principal components PC 1 and PC 2, with eigenvalues of 9.49 (percentage of variance: 63.27%) and 1.54 (percentage of variance: 10.29%), account for an explained variance of 73.56% and were represented on scores and loading plots graphs.

The third PCA was applied to the 47 glass tesserae based on the data from 9 characteristic elements of the

most common chromophore and opacifier groups [18] (Si, Ca, Mn, Fe, Ni, Cu, Zn, Sb, Pb). In some cases it has been described how high lead values can interfere with the measurement of certain elements [38–41]. This effect was only clearly corroborated in the Rb, which was therefore eliminated as a variable of the PCA. Principal components PC 1 and PC 2, with eigenvalues of 4.52 (percentage of variance: 50.26%) and 1.71 (percentage of variance of 69.27% and were represented on scores and loading plots graphs.

Based on the PCA and for the detailed analysis of small groups of tesserae of the same typology or colour, an analysis of the characteristic chemical elements was undertaken through binary representations and/or based on characteristic indices of pairs of chemical elements with ability for discrimination.

Results

Types of tesserae

As shown in Fig. 4, PCA offers an impartial criterion based on hXRF data to classify the tesserae into three main groups: two made up of stone and one of glass.

The first group was made up of 28 white, black, grey, red, pinkish-beige (also known as nudes) and mustardyellow stone tesserae characterised by high Ca, Mg, Fe contents. The second group included 7 black tesserae with significantly higher levels of Al, K and Ti than the rest.

The colours of the group of 47 vitreous tesserae were black, red, blue, green, turquoise, yellow and orange with high levels of Si, Co, Cu Sn, Sb and Pb. In contrast to other Roman mosaics [42–44], no ceramic tesserae were identified in the Judgement of Paris mosaic.



Types of glass matrix

In the case of the glass tesserae, not only is the nature of the chromophore interesting but also the type of glass. In this respect, considering the limitations of *in-situ* MRS and hXRF analyses, we carried out a thorough analysis using MRS laboratory equipment of two individual tesserae (one green and one black) that were found detached from but close to the Judgement of Paris mosaic.

The Raman spectrum of an amorphous silicate is composed of two broad bands at around 500 and 1000 cm⁻¹ that are assigned to the symmetric bending of the SiO₄ tetrahedra (δ_s SiO₄) and the symmetric stretching of the Si–O bonds (v_s SiO₄), respectively [45]. Observed bands are indicative of certain additives, the presence of which allows the classification of the glass. Thus, the Raman spectrum of the green tessera presents a clear profile associated with soda-lime-silicate glass corresponding to "Glass Family 3" in the Raman signature guide [34, 45, 46]. In the Raman spectrum we can also clearly identify the main bands at 235, 324, 335, 669 cm⁻¹ (Fig. 5a). All these peaks are characteristic of the calcium antimonate (Ca₂Sb₂O₇) orthorhombic phase that acts as an opacifier agent [45, 47].

In the case of the black tessera, the Raman spectrum profile fits the model or pattern described for lead glass classified in "Glass Family 7" (Table 1) [34, 46]. In general, a lead-rich glass is easily identified by a strong stretching mode at 959 cm⁻¹ (Fig. 5b). In the case of this tessera, the variation in the value of the stretching mode is linked to the concentration of Pb. An increase in the amount of lead causes a red-shifting of this vibrational mode [34, 47].

From the data obtained from both types of tesserae we can see that at least two types of glass are present in the group of 47 vitreous tesserae analysed. Judging from the Pb values obtained in the hXRF analysis, only one group of 6 black tesserae shows extremely high Pb values that would place them in the lead glass category. The rest of glass tesserae could be classified as alkali-silicate glass type.

Colours and raw materials

The applied methodology, based on the joint analysis by MRS, hXRF and on the statistical treatment from PCA and analysis of two variables, has allowed the identification and classification of the types of rock, chromophores and opacifiers used in the manufacture of the tesserae.

The PCA of the stone tesserae succeeded in differentiating a group of red tesserae with high Fe values (18, 19, 20, 23 45, 59, 60, 87, 88) and two groups of black tesserae, while the white, grey, red, black and yellow tesserae remained grouped with little definition (Fig. 6). In those cases, the MRS mineralogical analysis and the analysis of two variables provided fundamental information for improving their classification.

The PCA of the glass tesserae identified the orange, light blue, black and intense green groups of tesserae, while the dark blue and red tesserae and the others whose colour and opacity depends on their lead content (dark and light turquoise and yellow) were less well defined (Fig. 7) [40, 41]. In those cases, once again the analysis of two variables was necessary to improve their characterisation.

White

The Raman spectra of the white tesserae (5, 41, 75) present the typical bands of calcite at 155, 282, 712 and 1086 cm⁻¹ [48, 49], meaning they were made mainly of limestone (Fig. 8a) (Table 1). The Ca values registered in the hXRF analysis make them the tesserae with the highest Ca content in the whole scene and confirm their



Table 1 Raman spectroscopy results

Colour	Tesserae	Raman bands (cm ⁻¹)	Compound
White	5	280, 1086	Calcite [48, 49]
	41	282, 711, 1086	
	75	155, 282, 712, 1086	
Beige	14	175, 300, 725, 1097	Dolomite [35, 48]
	52	175, 299, 724, 1098	
	56	174, 300, 1097	
	71	175, 299, 1097	
	72	175, 300, 1098	
Red	18	224, 291, 410, 609	Hematite [35, 50]
	19	224, 291, 409, 610	
	20	224, 242, 291, 410, 611	
	23	224, 292, 410	
	45	223, 291, 410, 613	
	59	224, 291, 409, 610	
	60	224, 290, 410	
	87	224, 290, 409	
	88	224, 292, 410	
	90	154, 282, 711, 1086	Calcite [48, 49]
	40	282, 1084	
	54	281, 1086	
	63	154, 177, 282, 299, 712, 1086, 1098	Calcite [48, 49], dolomite [35, 48]
	76	156, 282, 710, 1086	Calcite [48, 49]
Black	34	281, 1086	Calcite [48, 49]
	42	282, 1086	Calcite [48, 49]
	43	142, 397, 463, 515, 636	Anatase [32, 33], quartz [33, 35]
	78	142	Anatase [32, 33]
	79	141	Anatase [32, 33]
	BT	140, 473, 635, 766, 959	Lead glass Raman signature [34, 46, 47]
Blue	9	670	Calcium antimonite [24, 45]
	10	670	
	11	670	
	12	670	
	47	670	
	68	670	
	91	670	
Green	GT	235, 324, 335, 483, 550, 669, 781, 952, 992, 1090	Soda-lime-silicate glass Raman signa- ture [34, 45, 46] Calcium antimonate [24, 45]

BT loose black tessera, GT loose green tessera

classification among the calcareous rocks known as limestones (Table 1).

Grey

nuance the result for the type of rock used and bring it closer to the dolomitic limestone category (Table 4).

Two types of grey tesserae were analysed (Fig. 2b, c): clear (33) and dark (44). Neither case provided a well-defined Raman spectrum, although their high Ca values indicate that both were made of limestone (Table 2). The somewhat more notable presence of Mg in Tessera 33 could

Pinkish-beige

The set of analysed tesserae (14, 52, 56, 71, and 72) repeatedly shows the characteristic Raman bands of dolomite rock at 175, 300, 725, 1097 cm⁻¹ [47] (Table 1) (Fig. 8b). The elemental analysis by hXRF, characterised by high relative amounts of Mg and, to a lesser extent,





Ca, gives them a different profile to the other calcareous tesserae made with limestone (white and grey tesserae) (Fig. 9a).

Red

Twenty-one red tesserae, differentiated mainly by the intensity of their colour, were selected for study (Fig. 2e, f). Their Raman profiles, the PCAs (Figs. 4, 6, 7) and the Mg/Ca ratios obtained in the hXRF analysis (Fig. 9) place them in three differentiated groups (Tables 1, 2, 3 and 4):

Group 1. Nine tesserae (18, 19, 20, 23 45, 59, 60, 87, 88) with Raman spectra that coincide with that of hematite $(Fe_2O_3, 224, 291, 410 \text{ and } 609 \text{ cm}^{-1} \text{ bands})$ (Fig. 8b) [50]

and with a high Fe content. The PCA also classifies them differently (Fig. 6). The tesserae in this group would have been made from ironstone (Fig. 9b).

Group 2. Nine tesserae (21, 22, 40, 54, 58, 63, 76, 89, 90) that present the Raman spectra of calcite (bands ca. 154, 282, 711, 1086 cm⁻¹) and higher Ca values than the rest of red tesserae. They would have been made from limestone rocks although some of them (40, 54, 58, 63, 76) present a high enough Mg content to catalogue them in the dolomitic limestone rock category (Fig. 9c, d).

Group 3. In three tesserae (13, 15, 46) (Fig. 9e) classified as vitreous in the grouping shown in Fig. 4, a particularity was detected based on the classification established by



the PCA of the glass tesserae (Fig. 7). Tesserae 13 and 15 present higher values of Cu, indicating that this element acts as the chromophore in form of cuprous oxide (Cu₂O)

or metallic copper (Cu^0) [51]. In the case of Tessera 46, which is opaquer, the combination of Cu and Pb (acting as a stabiliser for colouring crystals) that was introduced

Tesserae	Mg-Ka	Al-Ka	Si-Ka	P-Ka	S-Ka	CI-Ka	K-Ka	Ca-Ka	Ti-Ka	V_Ka	Mn-Ka	Fe-Ka	Co-Ka	Ni-Ka	Cu-Ka	Zn-Ka	Rb-Ka	Sr-Ka	Zr-Ka	Sn-L	Sb-L	Pb-L
5-W	221	2827	20,334	p.n	n.d	11,232	8624	6,190,000	p.u	n.d	5797	17,664	p.u	127	60	721	p.u	8778	p.u	h.n	n.d	357
14-Be	1429	3012	31,041	n.d	8779	p.u	17,828	4,050,000	7263	n.d	11,807	143,011	n.d	166	158	830	249	8821	p.u	n.d	n.d	342
18-R	p.n	4020	55,991	2889	11,027	21,464	36,084	3,180,000	15,829	n.d	16,899	2,630,000	p.n	131	306	1295	227	13,117	1833	n.d	h.n	h.n
19-R	p.n	4268	68,730	3684	8809	19,924	44,387	2,870,000	18,693	n.d	15,978	2,660,000	n.d	143	281	1476	303	12,561	461	n.d	n.d	n.d
21-R	p.n	4991	47,986	2235	7205	10,582	48,871	5,160,000	3234	p.u	8572	83,330	n.d	112	191	1571	443	7466	800	n.d	n.d	924
22-R	p.n	5052	50,536	2101	7866	12,608	53,784	5,480,000	2979	p.u	8349	92,272	n.d	141	188	1138	657	8374	854	n.d	h.n	1988
33-G	1134	3584	48,549	1774	7391	15,711	19,954	3,800,000	p.u	3755	4772	39,441	n.d	46	101	1005	1775	7143	8378	n.d	n.d	319
34-B	405	5558	54,215	3441	10,016	15,938	33,724	5,260,000	2235	p.u	5233	64,358	n.d	159	103	1181	p.u	15,469	p.u	n.d	n.d	546
39-Ү	211	4036	95,009	1644	9308	10,470	39,060	4,790,000	3017	9734	10,846	108,081	n.d	146	248	1461	1515	20,405	11,427	n.d	1784	1148
40-R	211	3373	22,353	1026	6738	10,747	22,819	6,050,000	p.u	p.u	5351	57,792	6853	160	95	902	444	6253	p.u	n.d	n.d	339
42-B	321	2842	23,914	n.d	13,203	13,179	9020	5,890,000	p.u	n.d	6923	35,013	n.d	125	104	983	229	15,879	p.u	n.d	n.d	253
43-B	p.n	21,794	227,770	3508	1591	6731	275,678	199,590	96,887	n.d	10,249	746,650	n.d	129	255	2055	p.u	8661	p.u	n.d	n.d	733
44-G	p.n	3320	33,292	463	9547	8751	27,259	5,680,000	p.u	16,140	5402	61,359	n.d	231	169	1287	15,265	15,951	10,152	p.u	n.d	316
45-R	p.n	3829	43,905	1361	13,617	22,453	16,191	3,490,000	p.u	p.u	54,560	2,740,000	n.d	106	177	1976	531	7487	p.u	n.d	n.d	3968
52-Be	1111	2862	31,260	n.d	4851	7596	23,598	3,710,000	674	p.u	19,496	88,422	p.u	271	1009	1331	382	7782	p.u	n.d	n.d	4426
54-R	300	4406	42,092	3334	8397	10,177	41,389	5,560,000	2833	p.u	8792	76,697	n.d	114	466	1333	584	9210	p.u	p.u	n.d	1007
56-Be	1500	2677	27,268	401	5060	10,194	16,564	4,000,000	n.d	n.d	7523	63,901	n.d	p.u	148	946	326	8299	p.u	n.d	h.n	359
58-R	284	3700	41,146	1852	n.d	10,858	22,380	5,480,000	1431	n.d	7541	118,596	p.u	p.u	247	1750	420	41,779	p.u	n.d	h.n	1268
59-R	249	5042	81,622	4113	p.u	27,820	31,876	2,010,000	20,497	n.d	67,883	4,010,000	n.d	137	748	1847	141	5504	p.u	n.d	h.n	2497
60-R	274	4551	65,480	n.d	21,459	31,403	25,051	2,730,000	14,767	n.d	56,205	3,320,000	n.d	n.d	277	1520	p.u	5397	p.u	n.d	p.u	2736
63-R	594	4000	30,313	1645	7703	13,118	26,242	5,380,000	2942	n.d	4295	80,704	n.d	150	181	1062	265	3008	658	n.d	h.n	540
71-Be	1456	2068	18,646	n.d	4924	11,128	8857	3,950,000	n.d	11,288	15,554	55,136	n.d	150	170	669	p.u	7136	2641	n.d	n.d	230
72-Be	1648	2094	17,484	h.n	5276	13,078	6803	4,010,000	p.u	2079	28,276	43,044	n.d	197	113	635	p.u	5700	p.u	p.u	h.n	179
74-Y	252	4036	97,952	1690	9710	12,421	40,065	4,800,000	1408	956	10,676	150,613	p.u	199	185	1426	1135	18,645	p.u	p.u	p.u	715
75-W	213	3175	22,609	n.d	9942	13,290	8460	6,250,000	p.u	p.u	5770	22,995	n.d	102	360	949	472	4564	n.d	p.u	n.d	758
76-R	277	4043	33,425	n.d	8555	15,110	37,179	5,600,000	2969	p.u	4265	92,096	n.d	173	123	1459	p.n	2380	n.d	n.d	n.d	422
78-B	p.n	25,011	252,165	3337	p.u	11,701	299,962	126,172	111,882	p.u	18,087	872,170	n.d	344	277	1888	p.u	9248	p.u	n.d	n.d	585
79-B	p.n	25,654	259,463	3595	p.n	8050	303,759	143,275	102,012	22,505	11,412	803,772	p.u	265	835	1622	15,765	10,320	10,141	p.u	n.d	2013
80-B	p.n	26,077	252,331	3384	p.u	9822	307,207	210,173	110,672	18,440	21,733	975,834	n.d	318	276	1982	15,691	8841	9782	n.d	n.d	985
88-R	n.d	5394	87,216	3781	11,886	28,160	42,314	2,500,000	17,684	6720	58,884	3,180,000	n.d	133	285	2157	n.d	6934	n.d	n.d	n.d	2252
89-R	n.d	4045	37,240	1931	9746	14,783	28,816	5,760,000	2312	4804	6846	51,402	n.d	130	192	1545	295	6057	588	n.d	n.d	322
90-R	p.u	4709	45,778	2883	11,118	16,392	37,508	5,430,000	5393	p.u	3786	84,565	n.d	135	222	1396	257	2732	373	p.u	n.d	564
93-B	p.n	18,942	177,155	6773	8853	13,033	502,650	1,290,000	89,615	p.u	28,390	972,776	n.d	205	1262	2627	p.n	120,839	n.d	n.d	n.d	3246
94-B	p.n	16,412	150,352	8318	7208	10,051	515,081	1,530,000	62,518	17,332	15,606	732,325	n.d	170	626	2766	13,704	78,575	30,525	p.u	n.d	1322
95-B	p.n	19,469	185,139	5928	3118	13,910	364,764	806,811	87,431	9152	18,586	871,385	p.u	182	607	3011	23,584	105,190	22,937	p.u	n.d	4449

Raw net counts of Ka and La lines of detected elements in stone tesserae *W* white, G grey, *Be* beige-pinkish, *R* red, *B* black, *Y* yellow, *n.d.* non detected

Table 2 hXRF in situ results



into Roman tesserae from the first century AD on would explain that particular characterisation [52–54]. That notable presence of Pb would be the reason it is classified among the black lead tesserae. This notwithstanding, a second option aimed at cataloguing that tessera as a lead glass type would be feasible. In the future, a larger number of analyses of red tesserae of this type would be able to settle the question of their classification.

Black

Fifteen black tesserae were analysed. As mentioned above, there are black glass tesserae and black stone-type tesserae (Fig. 2g). A more detailed analysis of both categories allows us to identify 4 types according to their composition:

Group 1. Six glass tesserae with a high Pb content (24, 27, 49, 77, 81, 82) (Table 3) (Figs. 7, 10a).

Group 2. Two tesserae made of black limestone (34, 42). The Raman spectra of calcite and the high Ca values in the hXRF analysis validate this classification in both tesserae (Fig. 10b).

Group 3. Three tesserae (93, 94, 95) manufactured from rock with a high Si and Al content (Figs. 6, 10c). They can be classified as coming from siliciclastic rocks and, considering the more than plausible presence of

aluminosilicates, in the mudrock category, probably black shales [55].

Group 4. Four tesserae (43, 78, 79, 80), that judging by their high Al and Si values were also made from mudrocks, but with a considerable Ti content (Table 2) (Figs. 6, 10d), as reflected in the presence of the 140 cm⁻¹ band associated with that oxide (TiO₂) (Table 1) (Fig. 8d). The presence of Ti is common in shales [56], meaning that this group could be a variation of the previous one.

Blue

The blue tesserae were made with vitreous paste and are differentiated in two groups according to whether their tonality is dark or light (Fig. 2h, i).

Group 1. The five dark blue tesserae analysed (1, 2, 3, 69, 73) contain cobalt as the chromophore responsible for that colour (Table 3). It is usually added as Co(II) to the glass matrix of the tesserae by using some kind of mineral salt or another material rich in cobalt [31, 57–59]. It is known that the colouring power is five times greater than that of other transition metals: to produce a deep blue colour only a few hundred *ppm* are needed [54, 60]. Despite the intensity of the blue provided by the cobalt, the values of Sb lead us to consider, as has been shown in other cases [60, 61], that calcium antimonate was added as an opacifier.

Table 3	hXRF i	n situ i	results																			
Tesserae	Mg-Ka	Al-Ka	Si-Ka	P-Ka	S-Ka	CI-Ka	K-Ka	Ca-Ka	Ti-Ka	V_Ka	Mn-Ka	Fe-Ka	Co-Ka	Ni-Ka	Cu-Ka	Zn-Ka	Rb-Ka	Sr-Ka	Zr-Ka	Sn-L	Sb-L	Pb-L
1-DB	n.d	6917	297,597	1279	5610	36,388	74,182	561,038	3354	n.d	95,862	178,327	46,214	810	45,128	1169	p.u	10,585	5917	n.d	5061	276,770
2-DB	p.u	7603	300,133	1459	3669	42,931	79,266	554,964	6584	1257	23,102	201,889	34,523	565	43,167	1737	1349	4480	6457	n.d	5564	335,828
3-DB	356	6214	304,266	1238	5497	43,022	57,950	578,120	4185	p.u	19,082	148,304	18,369	276	14,280	2319	1441	p.u	5381	n.d	4261	608,752
9-IG	255	7328	285,889	1693	11,499	34,152	90,613	659,684	6566	820	31,875	100,596	2969	462	235,179	7208	652	23,911	n.d	457	5706	54,196
2-IG	279	7008	249,772	3298	23,733	43,026	74,137	813,139	12,034	p.u	22,507	129,561	n.d	341	295,373	4008	724	27,902	p.u	n.d	3100	51,023
8-IG	p.u	8002	270,802	3646	8257	27,175	96,824	710,691	11,243	p.u	15,839	122,722	p.u	375	268,338	4579	752	27,669	n.d	n.d	3390	54,625
9-LB	p.u	7851	297,408	2247	p.n	46,076	85,908	607,565	14,477	p.u	73,456	134,552	p.u	218	3782	1714	1209	36,246	5822	n.d	2066	13,062
10-LB	381	7451	287,885	2511	6436	34,304	94,598	574,610	12,679	3106	61,541	127,206	p.u	206	16,552	2943	1130	35,348	6684	n.d	1558	13,723
11-LB	518	8195	343,464	1370	6199	42,408	93,433	526,221	11,497	1543	73,893	123,711	n.d	211	20,389	2166	1449	37,709	0609	n.d	1674	14,461
12-LB	p.u	8073	301,546	1928	6163	p.u	104,685	549,038	12,702	p.u	84,412	121,717	n.d	180	2918	2146	1247	32,025	6614	n.d	1197	83,586
13-R	p.u	6159	320,325	2034	6467	55,418	55,861	649,485	12,996	p.u	47,233	562,692	p.u	142	136,711	1522	670	28,335	5715	n.d	737	9673
15-R	p.u	8036	296,666	1941	2087	26,788	122,700	422,372	24,626	p.u	57,655	457,639	p.u	133	117,468	5577	1485	32,242	6456	h.n	532	298,418
24-B	p.u	9798	221,709	18,473	p.n	28,447	105,152	318,517	13,878	p.u	7673	155,728	p.u	n.d	1912	1723	n.d	n.d	n.d	h.n	h.n	2,480,000
27-B	p.u	6265	139,759	6479	p.n	27,793	62,099	367,665	11,193	p.u	11,587	177,355	p.u	n.d	2541	2371	p.u	n.d	n.d	n.d	p.u	2,300,000
28-DT	p.u	6983	330,444	2918	p.n	64,057	56,252	575,767	11,490	3985	70,579	94,574	p.u	n.d	121,312	1464	n.d	9103	7652	n.d	957	375,648
29-DT	p.u	7466	345,096	4031	7614	63,206	62,983	774,823	14,597	4387	50,729	138,364	p.u	84	85,462	2065	509	38,613	4230	p.u	684	6367
30-DT	p.u	7114	306,647	2810	3534	55,733	61,567	674,058	13,252	p.u	66,256	115,620	n.d	92	117,429	2476	1699	6920	7505	n.d	996	371,235
32-LT	p.u	6010	320,776	1206	p.n	53,073	55,181	570,247	10,653	p.u	42,727	102,269	n.d	111	115,878	1351	1897	n.d	8366	257	1577	691,081
35-Y	298	7568	329,561	2549	3689	47,222	65,264	631,495	11,890	p.u	33,055	98,382	n.d	144	662	1186	n.d	15,134	p.u	n.d	1182	190,663
36-O	p.u	0669	187,280	8036	p.n	32,988	78,022	653,771	26,312	p.u	19,158	226,372	n.d	158	328,387	17,478	1613	n.d	6787	643	474	1,640,000
37-DT	306	7638	241,227	3612	31,870	38,904	99,860	732,061	27,309	p.u	85,745	247,450	n.d	n.d	97,341	3478	p.u	37,074	n.d	n.d	772	19,908
38-DT	220	9320	228,704	5727	10,334	35,772	122,407	889,268	37,595	p.u	113,102	346,858	n.d	144	100,859	5767	830	18,586	6955	n.d	416	228,659
46-R	p.u	6598	157,111	4014	p.u	16,660	58,528	197,611	14,664	h.n	8782	121,034	n.d	83	6883	2729	n.d	2719	n.d	n.d	p.u	2,490,000
47-LB	p.u	5484	293,702	1300	p.n	34,644	69,253	646,351	11,881	p.u	47,320	99,354	p.u	n.d	5935	1769	n.d	35,352	n.d	n.d	1674	47,638
48-IG	p.u	6257	277,216	1553	8285	29,578	60,596	565,297	9493	p.u	21,607	117,025	p.u	280	297,835	3589	1030	24,272	p.u	n.d	4277	48,493
49-B	p.u	8031	211,422	3300	p.n	15,229	80,837	140,220	10,353	p.u	5397	105,421	p.u	341	5848	1829	679	2656	p.u	p.u	p.u	2,430,000
50-Y	228	6076	213,898	2457	p.n	22,947	58,374	397,143	8589	p.u	16,662	86,706	p.u	251	3346	2042	n.d	n.d	p.u	p.u	1579	1,620,000
51-Y	p.u	5249	235,664	3003	p.u	27,979	46,076	444,208	8898	n.d	39,352	117,416	n.d	181	3811	2405	n.d	n.d	n.d	259	1094	1,670,000
53-Y	164	5718	203,568	4738	p.u	34,315	55,609	390,810	12,850	h.n	56,176	306,543	h.n	198	2139	2400	647	n.d	n.d	119	1396	1,730,000
55-0	p.u	4900	162,049	3134	p.u	26,199	51,002	371,331	15,123	h.n	24,197	381,923	11,804	159	356,396	4620	896	n.d	n.d	350	611	1,560,000
57-0	199	6162	184,068	7190	p.n	34,203	65,531	487,680	22,099	n.d	22,433	259,026	n.d	166	301,055	17,878	807	n.d	n.d	481	627	1,670,000
61-Y	167	5409	224,371	4294	p.n	39,860	52,336	680,692	8568	h.n	32,648	117,501	n.d	130	3810	2249	827	n.d	n.d	n.d	1338	1,570,000
62-Y	307	6946	218,695	3896	30,170	30,674	73,091	759,422	17,262	3714	28,012	160,561	n.d	126	2176	2990	2124	n.d	6253	n.d	927	1,050,000
64-LT	283	6204	275,671	2153	6457	49,677	61,721	627,308	12,351	4368	39,841	105,369	n.d	163	102,704	2398	1787	n.d	7267	n.d	1435	634,801
66-DT	180	6087	279,817	3953	8179	54,751	55,487	677,691	11,898	3199	60,240	97,706	n.d	150	108,267	2059	1418	7465	6977	n.d	666	354,546
68-LB	243	8380	312,399	3561	14,044	67,152	98,453	710,226	13,248	2763	74,029	141,535	n.d	233	21,399	2982	1392	39,818	7114	n.d	2016	16,166
69-DB	147	5873	316,029	1564	9253	50,422	63,758	687,070	7282	2241	107,455	228,715	36,851	625	23,245	1177	1392	33,015	n.d	n.d	4513	56,245
0-02	249	6330	184,268	10,300	1773	52,621	99,329	486,979	26,214	3963	17,328	278,251	8589	148	378,168	21,026	673	n.d	n.d	991	763	1,380,000

Table 3	(contii	nued)																				
Tesserae	Mg-Ka	Al-Ka	Si-Ka	P-Ka	S-Ka	Cl-Ka	K-Ka	Ca-Ka	Ti-Ka	V_Ka	Mn-Ka	Fe-Ka	Co-Ka	Ni-Ka	Cu-Ka	Zn-Ka	Rb-Ka	Sr-Ka	Zr-Ka	Sn-L	sp-L	p-L
73-DB	302	5855	282,917	1673	5300	45,324	55,788	643,571	4054	3558	157,260	250,820	33,483	755	25,938	1142	p.n	24,415	p.u	p.u	3559 2	32,180
77-B	p.u	6106	137,369	4896	n.d	27,864	56,114	316,090	16,914	p.u	10,108	143,198	n.d	p.u	3869	3541	462	n.d	942	n.d	2 p.c	,440,000
81-B	p.u	6348	135,509	4715	n.d	28,336	65,455	328,926	15,384	26,190	9451	124,495	n.d	p.u	1845	2526	15,888	2867	9890	n.d	2 p.c	,390,000
82-B	p.u	6890	139,087	5433	p.u	29,228	67,193	334,980	15,944	3688	9644	126,125	n.d	146	2194	2371	p.u	2966	n.d	n.d	2 p.r	,400,000
83-LT	p.u	7615	270,500	2684	p.u	53,685	96,127	532,234	15,637	3722	31,939	247,147	n.d	p.u	105,786	3608	p.u	12,426	n.d	n.d	1229 1	,310,000
84-DT	p.u	7303	278,932	4751	p.u	70,547	70,048	735,715	13,355	5734	65,523	113,258	p.u	150	117,873	2708	p.u	31,151	n.d	n.d	1152 4	01,836
85-DT	p.u	6151	301,246	2077	p.u	58,046	56,496	626,326	14,081	3241	70,682	102,989	p.u	174	118,463	2543	p.u	30,417	n.d	n.d	1003 4	08,636
91-LB	p.u	7523	289,990	3226	0770	59,051	83,787	734,293	14,299	p.u	73,385	126,780	p.u	254	20,449	2080	378	35,698	951	n.d	2708 1	6,303
92-LT	p.u	6860	282,438	5674	9127	48,596	68,544	699,677	11,570	p.u	35,535	120,787	n.d	100	92,976	2496	1215	23,575	5238	253	1547 6	47,885
Raw net co	ounts of K	α and Lα	lines of det	tected el	ements in	glass tess	serae															

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R red, B black, DB dark blue, LT: light turquoise, DT dark turquoise, IG intense green, O orange, Y yellow, n.d. non detected

Table 4 Summary table

Material	Colour	Characterization	Tesserae	Location
Stone	White	Limestone	5, 41, 75	Background of the scene and Hera's dress
	Grey	Limestone	33, 44	Folds in Hera's dress, the stalks of the plant motifs and the shadow of Hermes' foot
	Beige	Dolomitic rock	14, 52, 56, 71, 72	Skin and clothing
	Red	Ironstone	18, 19, 20, 23, 45, 59, 60, 87, 88	Clothing, the ear of an ox, the dog at Paris' side
		Limestone	21, 22, 89, 90	Clothing, the mouth of an ox, Paris' hand, Hermes'
		Dolomitic limestone	40, 54, 58, 63, 76	hair, Athena's helmet
	Black	Limestone	34, 42	Tree and the ox close to Paris
		Mudrocks (black shales)	93, 94, 95	Heras' hair and outer perimeter around the scene
		Mudrocks (shales) + TiO_2	43, 78, 79, 80	Shadow lines of Aphrodite and Athena`s feet
	Yellow	Silicified limestone	39, 74	Hera's headwear, Athena's shield, one of the oxen
Glass	Red	Cu(l) (chr.)	13, 15, 46	Paris' clothing and Hermes' chlamys
	Black	Pb-rich glass	24, 27, 49, 77, 81, 82, BT	Vegetable and clothing
	Blue	Dark blue: Co(II) (chr.) + Calcium antimonate (op.)	1, 2, 3, 69, 73	Paris and Hermes' chlamys
		Light blue: Cu(II) (chr.) + Calcium antimonate (op.)	9, 10, 11, 12, 47, 68, 91	Paris, Hermes, Aphrodite, Athena and Hera
	Green	Light turquoise: Cu(II) + lead antimonate (high)	32, 64, 83, 92	Clothes of Athenea and Aphrodite, and in
		Dark turquoise: Cu(II) + lead antimonate (low)	28, 29, 30, 37, 38, 66, 84, 85	vegetable motifs (trees, schrubs) depicted near
		Intense green: Cu(II)	6, 7, 8, 48, GT	Palis
	Orange	Cu from bronze and/or brass (chr.)	36, 55, 57, 70	Hermes' chlamys and the apple, also found in Athena's helmet and the dog at Paris' side
	Yellow	Lead antimonite (chr.)	50, 51, 53, 61, 62	Hera's headwear, in details on Hermes' head
			35	Paris' clothes and the frame around the Medusa's head on Athena's breast

Colours and characterization of tesserae. chr. chromophore, op. opacifier, BT loose black tesserae, GT loose green tesserae

Group 2. In the Raman spectrum, the group of light blue tesserae (9, 10, 11, 12, 47, 68, 91) present the 670 cm⁻¹ band characteristic of calcium antimonate (Fig. 8e), although no other intense bands (Table 1). In this group, Cu in its divalent state was used to produce a light blue colour in the translucent glass [54] (Figs. 7, 11a).

Green

We have identified three groups of green tesserae: light turquoise, dark turquoise and intense green (Fig. 2j, k, l). Their manufacture depends on the amounts of Cu and Pb added during the preparation of the tesserae vitreous paste [21, 24, 62–64]. The PCA was especially conclusive in the separation of intense green tesserae (Fig. 7).

Group 1. According to the hXRF analyses of the four tesserae (32, 64, 83, 92) (Table 3), the light turquoise colour would have been the result of a mixture in diverse proportions of Cu (blue chromophore) and Pb (yellow chromophore), present in the form of lead antimonate $(Sb_2O_7PbO_2)$ [65]. The Cu/Pb ratio would have caused a variation ranging from lightest turquoise (Tessera 83)

to less light turquoise (Tesserae 32, 64, 92) (Fig. 11b) (Table 3).

Group 2. This same argument is valid for the dark turquoise tesserae (28, 29, 30, 37, 38, 66, 84, 85) (Fig. 11c). In this case, the reduction is based on the Pb values, resulting in tesserae with a darker, bluer tone due to the lack of the yellow chromophore.

Group 3. In contrast to the previous groups, the intense green coloured tesserae (6, 7, 8, 48) are characterised, according to the hXRF analysis, by the significant presence of Cu (Fig. 11d).

Orange

All the orange tesserae (36, 55, 57, 70) (Fig. 2m) were made of glass. In this case the chromophore agent was exclusively Cu, a very common element in this type of tessera [21, 62]. This group recorded the highest Cu values of all the tesserae analysed from the Judgement of Paris. The small amounts of Sn and even of Zn they all displayed indicate a bronze and/or brass smelting product as a source for the Cu-based colourant [54, 62] (Fig. 7) (Table 3).





Yellow

This group presents two types of tesserae according to the yellow tonality and the raw material used (Fig. 2n, o). Two of them (39 and 74) with a mustard-yellow tonality were made of limestone with an appreciable presence of Si. In fact, these are the limestone tesserae with the largest Si content of all those analysed in the mosaic. The Raman spectra of both samples were not conclusive, but the hXRF analysis indicated a significantly higher Ca/Si ratio (Fig. 12a).

The rest of the yellow tesserae (35, 50, 51, 53, 61, 62) are shinier and made of glass. Raman spectra was not



conclusive (Fig. 7), but, judging by the values of Pb and Sb present, lead antimonate must be considered their main colouring agent [66, 67] (Table 3) (Figs. 7, 12b). In this group, the composition of Tessera 35 shows a lower Pb content, which produces a type of tessera with a more matt, less glossy tone [67]. In general, although the origin of the lead antimonate is difficult to establish, different studies have proposed various options, among which the most likely are galena for the Pb and stibnite for the Sb [60, 67].

Discussion

Several aspects can be highlighted from the results presented, both from a methodological point of view and from an archaeometric and archaeological perspective.

Assessment of the methodology

The procedure followed is not exempt from the problems inherent to both techniques. Fluorescence background is a limiting factor in the use of MRS and, in the case of portable equipment, this is even more acute, especially when analysing glass tesserae [68–70]. The incorporation of loose tesserae and their analysis using high-resolution, large spectral window instruments can, in part, overcome this handicap.

In the case of the hXRF analysis, one of the clearest limitations is the detection of low Z elements (due to the strong absorption of low energy X-rays by air and by the Be window of the detector) [40, 41] Adlington 2017 y 2020. The inability of hXRF equipment to generate

high vacuum hinders its ability to correctly detect and quantify these elements. This problem affects particularly important elements such as Na, which is commonly present in glass and is necessary for assessing the manufacturing and degradation processes of this material [25, 26, 71]. Thanks to the two-beam (40 and 10 kV) measurement mode of the OlympusTM Innov-X Delta Premium instrument, it was possible to measure major low-Z elements [28].

Although the tesserae were fairly well preserved and were cleaned to remove any attached dirt or dust, we should not ignore the fact that some of them may have undergone superficial physical, biological and chemical alterations (especially those made from carbonate rocks) that could affect the measurements [72]. However, in cases of initial degradation, the penetrating power of the X-Ray beams used in this study, that is often considered a limitation for the use of the XRF technique in purely superficial measurements, would allow obtaining information of the chemical composition below the external surface.

Despite the aforementioned limitations, the impossibility of obtaining a representative number of detached tesserae and the need to maintain the integrity of the mosaic made it necessary to instigate an in-situ analytical strategy that, judging by the results obtained with the Judgement of Paris, has been effective.

The analysis of tesserae revealed the intrinsic difficulty of selection according to colour. Subjective as it may appear, the analysis based on objective, physical and chemical parameters bring to light issues in the chromatic criterion. Thus, tesserae initially selected according to well-differentiated shades (two shades of red, two shades of yellow and one black), reveal different chemical compositions that eventually lead to a more fine-grained classification: three shades of red, three shades of yellow and four shades of black. By contrast, tesserae that are clearly different in colour may reveal, upon analysis, the use of the same raw material (e.g. pink tesserae made of dolomites).

Stone tesserae

The *in-situ* analysis has not allowed us to perform petrological analysis on the Judgement of Paris, but the results obtained in the MRS and hXRF analyses of the stone tesserae have provided an enough level of detail to propose a first classification into three categories (Table 4):

- Carbonate rocks. Within this type we differentiated limestones of various colours (black, white, grey and red tesserae), silicified limestone (yellow), dolomitic limestones (grey, red tesserae), dolomites (beige/pink tesserae).
- Siliciclastic rocks. This group of rocks includes the mudrocks with the presence or absence of Ti for the black tesserae.
- Ironstone. This rock category includes the red tesserae with a recurring and abundant presence of hematite.

Of all the types of stone used, the most abundant are from carbonate rocks, particularly limestone, a type widely used in all the analysed mosaics dated between the second and the fifth centuries AD in the Mediterranean area [26, 44, 67, 73]. This type of rock is easy to cut, as it has a very compact texture and low porosity, thus facilitating the work of the artisans. To this property we have to add the uniformity of colour, a very important aspect for the creation of the mosaic, both for the composition of the figures and for the geometries, given that any veins in the tesserae deform their colouring and the portrayal. Another important characteristic of this material is the ease with which it can be polished to achieve a relatively smooth, non-shiny surface for the mosaic [74].

Regarding the possible origin of the stone tesserae, Cástulo is located in a geologically diverse region with a variety of sedimentary rocks, including conglomerates, limestones, dolomites, sandstones and granite outcrops [28, 75]. With this in mind, it is quite likely that the mosaic artisans made the stone tesserae using the various types of rocks available in the nearby territory of Cástulo. Particularly interesting is the identification of the use of ironstone to make numerous red tesserae. The use of this raw material is practically unknown in most of the Roman mosaics analysed in Europe. Exceptions include those from *Complutum* (Alcalá de Henares, Spain) [76] and Pompeii, although in the latter it is an imitation with limestone tesserae decorated with a layer of the reddish iron oxide mineral [26]. In the case of Cástulo, its use appears to indicate ease of access to this type of rock and, therefore, an especially differentiating aspect of the *Los Amores* Mosaic. Outcrops of ferruginous minerals in the Upper Guadalquivir and the surrounding mountains are common and their exploitation to obtain metal, pigments and building materials has been documented since Prehistory [77, 78].

Given that the majority of the stone tesserae are made from carbonate rocks (limestone and dolomite), they are susceptible to physical alteration due to the formation of soluble salts, fractures caused by thermal factors, the development of biological patinas (lichens), and chemical changes caused by dissolution in an acidic medium. In unpolluted environments, such as that of Cástulo, the acidity would have been caused by carbonic acid formed by the dissolution of rainwater in atmospheric CO_2 [72, 79–81].

Glass tesserae

Concerning the glass tesserae, in addition to identifying at least two types of glass used, this research has been able to study a considerable number of tesserae and obtain information about seven colour types (black, red, blue, turquoise, green, orange and yellow) (Table 4), the chromophore agents and opacifier used:

- Cobalt (Co⁺⁺) for dark blue.
- Copper for light blue (Cu⁺⁺), red (Cu⁺ or Cu⁰) and orange (metallic copper likely obtained from bronze and/or brass).
- Copper (Cu⁺⁺) for greens and copper and lead antimonate for turquoises.
- · Lead antimonate for yellow.
- The opacifier identified was calcium antimonate which is found mainly in the green, dark and light blue.

This expansion of the colour repertory investigated is particularly important, as yellow, orange and red tesserae had not been analysed in the studies of vitreous tesserae from pre-fourth-century-AD Spanish mosaics [44, 82– 87]. Although the information provided by hXRF does not allow any definitive conclusion to be reached about the source of the glass used in any of the tesserae (local or of eastern Mediterranean origin) [88], it is interesting to point out the case of the lead tesserae. The surrounding area and the mountains north of Cástulo are renowned for their rich Pb-Ag ore mineral veins. Cástulo was a major source of the extraction and management of the Upper Guadalquivir's mineral riches between the second century BC and the second century AD. The proximity of the lead mining areas and the presence of black vitrified lead slags on the surface of Cástulo suggest local manufacture. This proposal, which would challenge the model of specialized glass-colouring workshops, will be subject to verification with the expansion of the mosaic study and the analyses of the vitrified black fragments collected.

The important collection of glass tesserae identified also hints the most common pathologies associated with archaeological glass objects, such as craters and dealkalisation layers. The former are frequent and tend to appear as small spherical or oval holes that may eventually merge. Some are visible in the detached green tesserae under analysis (Fig. 5a). Dealkalinisation causes iridescent silica gel layers on the surface and is associated with phases of humidity and dryness. This humid/arid cycle would have affected the microstructure of the glass, giving rise to physical stresses that would have eventually fractured and separated the surface layers [83, 89].

Selection of colours

Another aspect to emerge from the analyses is the choice of the colour for each figure and the types of tesserae used to obtain it. The selection or intentional use for symbolic or practical reasons of tesserae in certain parts of the scene is contradictory and irregular. In some cases, it is to maintain specific identifying signs in the figures; in others it appears that a shortage of some colours compelled the use of similar pieces.

In general terms, stone tesserae were mainly used in the background of the scene, on animals and on some clothing and bodies. The vitreous tesserae were used for specific items of clothing characteristic to each figure, as well as on complements and plant motifs.

More precisely, the use of white in the figure of Hera on a background of white limestone tesserae can only be explained by the fact that white is a colour associated with that goddess. The same is true of Hermes, whose chlamys was depicted in red and orange tones from the fifth century BC on. The turquoise tesserae that make up Venus' clothing, which is normally silk, are also intentional. Centuries later, the mosaics of Casariche (Seville), Antioch (Turkey) and even Noheda (Cuenca) repeat wholly or partially the decorative schemes seen in Cástulo [14, 16, 18].

In other cases, the composition of a colour was achieved by combining tesserae made with different materials. The reason for this could be explained on artistic grounds, the deliberate exploitation of the different optical properties of the material, random selection of tesserae or an attempt to make up for a shortage of certain types. The case of the red, black and yellow tesserae is quite clear in this respect. In the red decoration of Hera's chest, we identified red limestone (89) and ferruginous origin (88) tesserae. The black tesserae used on the lines of Venus' dress contain a combination of glass (81 and 82) and black mudrock tesserae (80). In the case of the yellow tesserae, whereas Hera's headwear is created with silicified limestone and glass paste tesserae, the frame around the Medusa's head is a combination of two types of glass tesserae (Fig. 13).

Faced with the difficulty of establishing the origin of the glass tesserae from the hXRF data, the combination of glass and stone tesserae is of some help when it comes to the question of their supply and their possible foreign origin. The artisans responsible for the manufacture of tesserae, probably from an itinerant workshops operating in different areas of Andalusia [9], did not have a regular supply of glass tesserae. This could indicate a foreign rather than a local origin, otherwise they would not have had to substitute them in some places with other types of a similar appearance (yellow tesserae) or those made of stone (yellow, red and black tesserae). We will be able to reach more detailed and in-depth conclusions regarding these hypotheses when we evaluate all the analysed tesserae from the *Los Amores* Mosaic.

The Cástulo workshop

The particularity of some of the raw materials used in Cástulo calls for some considerations and explanations regarding the existence of schools and workshops specialising in the manufacture of mosaics in the south of the Iberian Peninsula. Research into the artistic and iconographic aspects has identified in this region a school with its main focus around the Guadalquivir (from Córdoba to the river mouth). The Cástulo mosaic shows iconographic and stylistic similarities that have favoured proposals to include it in that school, although represented by a differentiated, itinerant workshop operating in Jaén province from the second century AD [6, 9].

The data obtained from the archaeometric analysis supports this hypothesis if we compare it with other studies of mosaics in Andalusia of a similar chronology. Mosaics from Cártama (Málaga), Itálica (Seville), Carmona (Seville), Puerto Real (Cádiz), Écija (Seville) and Málaga have been chemically and petrographically analysed and all have been dated to between the second and the third centuries. Most of the results obtained from their study only refer to stone tesserae, in which limestone of various colours (white, red, black and yellow) is the predominant raw material, together with some tesserae manufactured with sandstone, volcanic rocks and pottery [44, 82–87]. To this group we could



Fig. 13 Details of Aphrodite, Hera and Medusa's head on Athena (from left to right)

add the mosaics from the Maritime Baths at Baelo Claudia (Cádiz), although their chronology remains undefined between the second and the fourth centuries AD [90].

The only thing the Judgement of Paris mosaic has in common with the previously mentioned examples is the use of diverse types of carbonate rocks, as there are no tesserae made with sandstone, volcanic rocks or pottery. As an example of the particularity of this mosaic and as a possible expression of the aforementioned workshop operating in Jaén province, of particular note are the tesserae made with dolomite, siliciclastic rocks and, above all, the previously mentioned ironstone.

Data referring to glass tesserae in the south of Spain are much sparser and it is difficult to make comparisons. Only twelve blue and green tesserae from the mosaics of Itálica and Cármona with Cu and Fe identified as chromophore agents belong to this period [83, 87]. In Cástulo, on the other hand, it has been possible to study one of the most comprehensive repertoires on the Iberian Peninsula. One differentiating element of this hypothetical workshop, which would have to be confirmed with further research into glass tesserae, would be the use of the aforementioned lead tesserae. The use of this type of glass has not been documented in the archaeometric analyses undertaken on the Iberian Peninsula, either in mosaics contemporary to Los Amores Mosaic or in those to which later datings are attributed [17, 18, 83, 87]. This singularity, linked to the types of stone tesserae, would therefore add credence to the stylistic and iconographic identification of a workshop of artisans who would have been operating in the Upper Guadalquivir with Cástulo as their main place of work.

Conclusions

The results achieved in this study are mainly important in several respects. A particularly relevant aspect with significant repercussions is related to the effectiveness of in situ analysis using portable non-invasive spectroscopic techniques. Despite the aforementioned limitations, the quantity and quality of the information obtained justifies its use. This is especially interesting for overcoming problems caused by the shortage of loose tesserae and for tesserae sampling based only on colour types.

The results obtained from the mineralogical and elemental analyses of the tesserae in the Judgement of Paris position the *Los Amores* Mosaic as a benchmark for our knowledge of the stone or vitreous raw materials, chromophores and opacifiers used in creating the mosaics of Roman Hispania. There are no archaeometric studies on the Iberian Peninsula of mosaics dated between the late first and third centuries AD that can offer such a full and varied catalogue of materials as the *Los Amores* Mosaic. In it we find the combination of stone and glass tesserae. The latter were mainly of the alkali-silicate glass type, although some of the black tesserae were also found to have been made of lead glass.

The characterisation undertaken is of special importance, as the dissemination, conservation and restoration strategies have to take this information into account. Of particular note is the combination of tesserae of the same colour manufactured with similar glass and stone as a consequence of the scarcity of certain types of tesserae. It is also important because more mosaics have been excavated in other rooms of Building D and it may be possible in the future to establish similarities and differences between the diverse types, depending on their location and the importance of the area. Finally, the proposal for a workshop of mosaic artisans operating in the Upper Guadalquivir and Cástulo, open a new line of archaeometric and archaeological research.

Supplementary Information

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Additional file 1: Figure S1. a Location of Cástulo (Linares, Spain). b Virtual reconstruction of Cástulo city at Roman times. c Building D with *Los Amores* Mosaic. Figure S2. *Los Amores* mosaic in Cástulo (Linares, Spain). Figure S3. Areas 1, 2, 3 and 4 of Judgement of Paris in *Los Amores* mosaic. Figure S4. In situ analysed tesserae of Judgement of Paris in *Los Amores* mosaic (Area 1). Figure S5. In situ analysed tesserae of Judgement of Paris in *Los Amores* mosaic (Area 2). Figure S6. In situ analysed tesserae of Judgement of Paris in *Los Amores* mosaic (Area 3). Figure S7. In situ analysed tesserae of Judgement of Paris in *Los Amores* mosaic (Area 4).

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Authors' contributions

JT, AS, MM. and PV: coordination, conceptualization, wrote the main manuscript texts and analysed the overall results. BC and PA: selected samples, prepared figures, tables and analysed the results. JT and AS: conducted EDXRF and analysed the results. MM and PA conducted portable MRS and analysed the results. AR, SL, DS and MC: conducted handheld XRF and analysed the results. All authors read and approved the final manuscript.

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Availability of data and materials

All data generated or analysed during this study are included in this published article (and its supplementary information files).

Competing interests

The authors declare that they have no competing interests.

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