A multi-analytical study of historical coated plaster surfaces: the examination of a nineteenth-century V&A cast of a tombstone

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Abstract
A multi-analytical study was designed to characterise historical coated plaster surfaces. The method was applied to investigate the surface coatings of the nineteenth-century plaster cast of the tombstone of the Presbyter Bruno that belongs to the Victoria and Albert Museum collection. At first, selected samples of the object were examined with Visible Light Reflectance and Ultra-Violet Fluorescence Optical Microscopy (VLR- and UVf-OM respectively) and Scanning Electron Microscopy (SEM) demonstrating a consistent stratigraphy featuring a bulk, an interface and an uppermost layer. The latter layer appeared to consist of an aged coating and dirt. Overpainted and repaired areas of the object generated samples that had additional layers on top of the aforementioned stratigraphy. A layer that seemed to be an additional surface varnish or a coating that had not been absorbed to the bulk has been observed in a couple of samples. Elemental characterization was carried out with energy dispersive X-ray spectroscopy (EDS) and further analyses were performed with X-ray diffraction (XRD) and Fourier-transform infrared (FT-IR) spectroscopy with focal plane array (FPA) imaging which confirmed that the bulk of the object is made of gypsum plaster containing mostly silicate and carbonate inclusions. Gas chromatography/mass spectrometry (GC/MS) and pyrolysis-GC/MS with extraction methods based on n-propanol followed by pentafluoropropionic anhydride (PFPA), tetramethylammonium hydroxide (TMAH) and 3-trifluoromethylphenyltrimethylammonium hydroxide (m-TFPTAH) were performed to detect organic media. The results suggest that the organic medium used for the surface coating is a diterpenic resin that contained silicon, aluminium and traces of other inorganic elements. The organic medium of overpainted areas was based on alkyd resins and the in-paints were characterised as a blend of silicon and barium at varied concentrations. This multi-analytical approach can generate a better understanding of manufacturing, component materials and conservation issues of coated plaster objects.

Keywords: Coated plaster casts, Organic coatings, Multi-analytical examination, Optical microscopy, Scanning electron microscopy (SEM), Fourier-transform infrared spectroscopy (FT-IR), X-ray diffraction (XRD), Energy dispersive X-ray spectroscopy (EDS), Pyrolysis-gas chromatography/mass spectrometry (py-GC/MS)

Introduction
This work represents a pilot case study designed to establish a wider campaign of sampling and scientific analyses of the surface layers of the casts produced in the nineteenth century, in the framework of a collaborative doctoral partnership of Northumbria University with the V&A Museum (AH/R00322X/1) [1]. Part of the project...
focused on the characterisation and the ageing characteristics of the original coatings of historical plaster casts [2]. In the nineteenth century, the plaster cast replicas of famous artworks were used extensively in schools and museums [3–5]. The aim was to educate and inspire those who studied or appreciated art. They were also fashionable among the more affluent people who decorated their homes with plaster ornaments and copies of well-known sculptures [6–9].

The object of this study is a nineteenth-century plaster cast of the Tombstone of Presbyter Bruno, part of the Victoria and Albert Museum (V&A) collection (Museum Accession Number REPRO.1873-380, Fig. 1) and displayed in the Cast Courts (Gallery 46A). The ‘Copy of the Tombstone of Presbyter Bruno’ is a plaster replica of an original tombstone from the Cathedral of Hildesheim (Germany), by Friedrich Heinrich Nicolaus Küsthardt the elder (Göttingen 1830–Hildesheim 1900) [10, 11].

The original tombstone is dated 1194 and is still located in Hildesheim Cathedral. In 1873, the Museum acquired several plaster casts of key sculptural decorations in Hildesheim Cathedral, all produced by Küsthardt, among them, the tombstone, purchased directly from Küsthardt in 1873 for £6.

Küsthardt's plaster copy of the tomb slab, 218.5 cm high by 77.0 cm wide, is currently displayed between a mix of casts of different origin, as well as a variety of early Christian monuments and other reproductions from Hildesheim Cathedral. The cast is mounted on a wooden display frame, and on its reverse, is another German reproduction depicting an upper left portion of a wooden doorway from the Church of St Maria im Kapitol, in Cologne (the original was carved c. 1065 by an unknown carver) [12].

Neither cast appears in any of the early photographs of the V&A galleries. Yet later pictures of Küsthardt’s cast on its own, held in the V&A curatorial department, reveal that it was formerly mounted upright on a different backing, with metal brackets supporting its base (Fig. 2). The brackets were later removed which explains the current losses to the plaster under the relief. The photographs also reveal a series of exposed metal screws at the front to attach the plaster to its wooden or metal supporting frame (Fig. 2). Many of the casts in the collection have been fixed to their supports with similar metal screws. Often the screw-heads have been filled and retouched to conceal them, which is the case with Küsthardt’s piece. The positions of fixings are easy to recognise, as the filler is often slightly proud and, in many instances, split around the edges. This is due to the corroding metal or shrinkage of the filler. Also, the overpaint is often of a slightly different colour, either because of ageing, or the retouching was aimed to be noticed, in case the objects needed deinstalling.

A close examination shows that the cast is made of three sections, which sit directly on top of one another, with pieces of wood wedged between them. Interestingly, the original sandstone slab, which is mounted on the south wall of the choir at Hildesheim cathedral, is carved from a single block of stone, although it has additional stone sections added at the top and on its base, which have not been replicated in the V&A copy. Also, by comparing archival images of the original slab, some of the losses to the original stone surface do not show on the cast, as can be observed. It may be that some of the surface details on the original stone were lost after the cast was made, or some of the details on the cast were later reworked.
The top section of the cast has various vertical and horizontal lines under the arch surrounding the Christ’s head. The lines have been sanded down, suggesting the cast could have been made using a piece mould technique. Piece moulds tend to leave raised seamlines on casts and, they were often disguised by sanding them down [2].

The outer edges of the cast have been covered with painted plywood sheeting, to conceal their unevenness (Fig. 3). Interestingly, the edges of the original stone slab are finely finished. Many casts in the V&A collection were not meant to be viewed all around. They are often unfinished at their sides and the top and were later covered with either fabric, card, wood, or metal to disguise the exposed areas and to protect them from dust.

A close examination of the underside reveals that there is a small section of metal visible at the back of the cast, on the left side. It could be what remains of the old bracket after it was cut back, or it could be part of a larger metal support. Metal and wooden supports are commonly found on the backs of the plaster casts. Using a metal detector, it was revealed that the top and middle parts of the cast have less metal backing support than the lower part of the cast.

Küsthardt’s cast has been restored on several occasions since it came to the museum some 148 years ago. Unfortunately, there are no records of any of the previous restorations, but a close examination reveals that numerous localised areas on the surface have been repaired with a filler and touched up using different paints and colours at different times. It was fairly common before the 1970s, for the restorers, not to document their treatments. However, many casts in the collection have names and dates pencilled on discreet areas on the object’s surface. This information helps to pin down when objects were restored and by whom. Casts also might have traces of old materials on their surfaces: residue of soap or residue of casting materials such as gelatine, or items at their backs including, packages of soap, pigments, cloths, sponges, tools, newspapers, letters, food packaging etc. Unfortunately, due to the framing of the cast, it is difficult to observe the back of Küsthardt’s cast.

Although there are no signatures or dates on Küsthardt’s cast visible at the front, apart from museum numbers near the base and at the top, and Küsthardt’s company label can be found on the top right corner on the lower section. It reads: ‘Bildhauer, FR Küsthardt, Hildesheim’ in embossed lettering (Fig. 3).

There are also many tool-marks and possibly fingerprints where the plaster surface has been worked on. Several raised areas, particularly on the lower sections, are also covered in pencil marks. These were probably left by various artists and students who sketched and drew the object in the galleries ever since they were built. Similar pencil marks on the plaster surfaces have been observed by the authors in art schools’ collections, as for example the one in the Hatton Gallery in Newcastle upon Tyne.

Research on plaster cast objects often remains unpublished at large within the records of collection holding organisations and individual case studies fail to represent the breadth and complexity of the materials that can be found in the eclectic cast collections due to case
specificity [13–16], whereas often useful insight on the properties of materials can be found in studies on building decorative materials [17], such as plasterworks [18] and mortars [19]. Megens et al. [20], on the other hand, have demonstrated that systematic elemental analysis is required to uncover the provenance and composition of plaster used to replicate art and decorative objects: traces analysis, size and distribution of porosity and mineral shape and growth can be characteristic of a particular group of artefacts. Gypsum plaster (calcium sulfate, CaSO$_4$$\cdot$0.5H$_2$O) has consistently been found as the main component of the bulk of the nineteenth-century plaster casts [2]. However, organic (such as resins and gums) and inorganic compounds (such as clay, sand, lime), as well as larger structural elements (such as fabrics, wooden or metal batons or even bones) were combined and added during the plaster production, as used to improve mechanical properties, such as hardness and water resistance or to control setting time and the casting procedure [2, 20]. The complexity of the organic and inorganic blended compositions in the plaster artefacts suggests that a multi-analytical approach would be more appropriate. For example, Field emission gun–scanning electron microscopy (FEG–SEM), XRD and Particle induced X-ray emission (PIXE) allowed the examination of Renaissance stucco related materials, which indicated that the composition of the mineral phases influences its sustainability and sensitivity to moisture sorption processes [21].

The full characterization of the materials of the cast will also provide fundamental information for the deterioration processes and guide conservation decisions. Typical damage observed is related to environmental conditions or unsuitable protection and handling. Plaster objects may become heavily soiled and covered with dirt and dust layers. Exposure to high relative humidity levels

Fig. 3 The cast still retains the stamp of the workshop of Friedrich Küsthardt (a). A closer look at the object revealed the plywood sheeting used to conceal the cast’s edges unevenness (b). An accidental spillage was highlighted under UV illumination from the area where sample 12 was taken (c)
causes the migration of deposits into the plaster’s porous structure. Due to alternating swelling/shrinking cycles in time and the fragility of plaster materials, objects often lose their texture, crack or break into fragments [22].

This study aims to characterise the materials used to produce the surface layers of the plaster casts, either to provide the plaster with certain properties and to achieve the desired appearance. While VLR- and UVf-OM allowed the identification of the stratigraphy, SEM–EDS and XRD were used to monitor the inorganic composition and FT-IR and (py)–GC/MS to characterise the organic components. Trace analysis to investigate the provenance of the object were not carried in this study, being beyond the scope. The study of the object served as a pilot study for the determination of an analytical strategy for the examination of the surfaces of plaster casts that will eventually allow a better understanding of the workshops’ practices in place in the nineteenth century and define more specifically targeted conservation methods. Archival information was available in the Registry of Reproductions of the V&A Museum (1873), in the museum Collection Management System (CMS) and the V&A archive currently at Blythe House, London.

**Experimental Sampling**

A total of thirteen samples from selected areas were taken, according to British Standard BS EN 16085:2012 (ISBN 978 0 580 70588 5). Before the sampling, a careful survey was performed to prevent any risk and to minimize the quantity of sample collected, which was never larger than 1.0 mm across, and maintain the integrity of the object. The sampling procedure was fully documented [23]. The sampling areas were determined by many factors, such as accessibility and significance, but also avoiding foreground areas. The samples were taken from areas of pre-existing loss and undercuts or marginal areas. The utmost attention was given to ensure that the samples were collected limiting any contamination. Before being stored in the vials, the samples were numbered with the museum accession number and the progressive sampling number, which is used throughout the study to identify the samples, as follows: *MUSEUM ACCESSION OBJECT NUMBER_PROGRESSIVE SAMPLE NUMBER*.

**Technical photography**

Regular visible photographs were taken with a Panasonic DCM-FZ38 camera under the gallery’s normal illumination (i.e., diffuse lighting, skylight window natural light and mixed artificial illumination). Colour and dimension references were ensured through the Past Horizons® Credit Card Photography Scale. The images were processed with Adobe Photoshop® CC 19 and white-balanced through the Past Horizons® Credit Card Photography Scale. The objects were also rendered in Autodesk® AutoCAD® 2019 for mapping purposes (Fig. 4).

**Stereomicroscopy**

A StereoZoom® LEICA S6D stereomicroscope was used to observe the samples, to understand the shape of the samples, the position of the layer in the stratigraphy and to define the processing of the sample. The Leica S6D Stereomicroscope has a 10× eyepiece and the objective magnification range from 0.63× to 4.00×. When possible, samples were split into two parts: one fragment was embedded in polyester resin to allow the observation of the stratigraphy and the analysis of the layers and the other was put aside for destructive analyses.

**Samples for cross-sectional analysis**

The samples were embedded in Alec Tiranti™ Ltd clear casting resin, which required 48 h to cure and harden. Alec Tiranti™ Ltd clear casting resin consists of styrene and methyl methacrylate/polyester resin (Product Code: 405-210) and liquid hardener (BUTANOX M-50 methyl ethyl ketone peroxide, solution in dimethyl phthalate—Product Code: 405-810) in the proportion 4 mL: 1 drop.

**Visible light reflectance (VLR) and ultraviolet fluorescence (UVF) optical microscopy (OM)**

Optical microscopy was performed with an Olympus BX51 Metallurgical Microscope equipped with four objectives (magnification of ×5, ×20, ×50 and ×100), and an ×10 eyepiece. 50 µL of white spirit were added on the surface of the cross-section to improve the saturation under the microscope. The microscope is equipped with a 6-cube filter turret which allows operating the system in reflected visible light (brightfield and darkfield mode) and reflected UV light (365 nm) using a 100 W mercury burner.

**X-ray diffraction (XRD)**

The XRD analyses were performed with a Rigaku SmartLab SE equipped with a HyPix-400, a semiconductor hybrid pixel array detector and a Cu source. The analyses were performed in Brag-Brentano geometry mode, with 40 kV tube voltage and 50 mA tube current. The diffractograms were processed with SmartLab II software. The data was compared to the data available in the RUFF™ database [24] and the Crystallography Open Database (COD) Database [25].
Scanning electron microscope (SEM)—energy dispersive X-ray spectroscopy (EDS)
The SEM–EDS analyses were performed with a field emission TESCAN MIRA 3 with gigantic chamber. The SEM is equipped with a back-scatter detector (BSE) and back-scatter in-beam detector (In-beam BSE). For the EDS analytical part, it has an Oxford Instruments setup: Software: AztecEnergy, X-ray detector X-Max 150 mm² and X-ray detector X-Max Extreme, low energy detector for thin films, high resolution and low voltage. The samples were analysed by SEM–EDS Low Vacuum Mode (10–15 Pa). EDS Mapping and data processing were performed with Aztec Oxford software. A fragment of sample 2 was mounted on appropriate support, adhered with silver paint and coated with a layer of platinum (5 nm thick). This sample preparation is required when high vacuum SEM–EDS (1.5 × 10⁻² Pa)
is performed. This mode allows a higher magnification with a better definition.

Fourier-transform infrared spectroscopy (FT-IR) with focal plane array (FPA) imaging
A Perkin Elmer Frontier FT-IR spectrometer (350 cm⁻¹ at a best resolution of 0.4 cm⁻¹) was used, equipped with a UATR Diamond/ZnSe ATR accessory and combined with a Spectrum Spotlight 400 FT-IR microscope equipped with a 16 × 1 pixel linear mercury cadmium telluride (MCT) array detector standard with InGaAs array option for optimised NIR imaging. Spectral images from sample areas are possible at pixel resolutions of 6.25, 25, or 50 μm. The Perkin Elmer ATR imaging accessory consists of a germanium crystal for ATR imaging. These run with Perkin Elmer Spectrum 10™ software and with SpectrumIMAGE™ software. Baseline and Kubelka–Munk corrections were applied to the raw data acquired in diffuse reflectance (DR). The FT-IR spectrometer range is 350–4000 cm⁻¹ in the ATR mode and 650–4000 cm⁻¹ in the DR mode.

Gas chromatography/Mass spectrometry (GC/MS) and pyrolysis-GAS chromatography/mass spectrometry (py-GC/MS)
The instrument used for GC/MS is a Thermo Focus Gas Chromatographer with DSQ II single quadrupole mass spec. The column is an Agilent DB5-MS UI column (ID: 0.25 mm, length: 30 m, df: 0.25 μm, Agilent, Santa Clara, CA, USA), fitted with a Pyrolab 2000 Platinum filament pyrolyser (PyroLab, Sweden). The helium carrier gas flow rate was 1.5 mL/min with a split flow of 41 mL/min and a split ratio of 27. The temperature of the detector was set at 280 °C and the inlet injector temperature to the GC was kept at 250 °C. The pyrolysis chamber was heated to 175 °C, and pyrolysis was carried out at 600 °C for 2 s. Samples derivatization was necessary for GC/MS [26–30] and carried with n-propanol followed by pentafluoropropionic anhydride (PFPA) [31], to derivatise protein component of proteinaceous paints and adhesives, while yielding the propyl esters of fatty acids derived from lipids and diterpenoid acids derived from natural resins and thus allows the choice of a single method for the analysis of artists media which contain either oils or proteins or mixtures of both proteins and oils or even resins. 1 mg of pulvrised sample was hydrolysed in 150 μL of Hydrochloric acid (HCl) and the excess Oxygen was removed. The solution was heated at 90 °C for 3 days and placed in a vacuum desiccator for 24 h to remove the acid. After 180 μL of propan-1-ol: acetyl chloride (3:1) were added, the solution was heated at 110 °C for 45 min, then at 50 °C for 30 min. The reagent excess was then removed via nitrogen. The residue was dissolved in 50 μL 0.2% pyridine and 150 μL of dichloromethane (DCM) and 150 μL perfluoropropionic anhydride (PFPA) were added before heating at 110 °C for 15 min [31]. The sample was then injected into the column with the aid of a micro-syringe and the MS thermal Programme (3) was set as follows: Seg1 start 13.00 Scan events MS, Heated zones Ion Source 250 °C, Detector Gain 1.21-105 (Multiplier Voltage 1025 V). Oven: Initial temp 60 °C hold 2 min Ramp 1 6.0 °C/min (rate), 250 °C, hold 0 min. Ramp 2 25 °C/min, 300 °C, 20 min hold. Mode: split. A pulvrised sample of the order of 0.5 mg was either directly derivatised in an aliquot of 1 μL of 25 wt% in methanol tetramethylammonium hydroxide (TMAH) and placed on the Pt filament, or methylated with 3-trifluoroacetylmethylmethylene ammonium hydroxide (5 wt% in Methanol) CAS number 68254-41-1, C₃H₁₃F₃NO m-TFPTAH (commercially known as MethPrep II). Methylation has been widely used for the analysis of artists’ media where it is useful for the analysis of both seed oils and natural resins which contain diterpenoid acids or triterpenoid acids such as moronic acid from mastic [29, 31–33]. MethPrep II methylation was achieved either: (1) by adding to the sample 1–3 drops of MethPrep II, depending on sample size, and heating at 60 °C for 24 h, or (2) by adding 30 μL of MethPrep II each 0.3 mg of sample and heating at 60 °C for 24 h. The MS Thermal programme (1), (2) or (3) were set for the analysis, as described hereafter. Thermal Programme (1). Seg1 start 2.40 Scan events MS, Heated zones Ion Source 250 °C, Detector Gain 1.21-105 (Multiplier Voltage 1025 V). Oven: Initial temp 40 °C hold 4 min Ramp 1 10.0 °C/min (rate), until 250 °C, hold 15 min. Mode: splitless. Thermal Programme (2). Seg1 start 2.40 Scan events MS, Heated zones Ion Source 250 °C, Detector Gain 1.21-105 (Multiplier Voltage 1025 V). Oven: Initial temp 40 °C hold 4 min Ramp 1 10.0 °C/min (rate), 250 °C, hold 45 min. The acquisition was carried out in a Total Ion Count mode, where all ions in the range 40–800 m/z were monitored. The Xcalibur™ 2.2 and PyroLab™ software were used to control the instruments. The former was then supported by the library browser supported NIST MS Version 2.0 [34] which facilitated data processing.

Dataset of analysis
A comprehensive dataset for the analysis of this object was deposited in Northumbria University’s Figshare repository [23]. A full database of results in a spreadsheet format and with plots and figures was compiled. The description of the samples and images of the sampling site location can also be found in the dataset.
Results and discussion

The analyses were chosen to ensure the understanding of the coating materials, which was the scope of the research. However, a sample of incoherent dirt and dust was taken from the base of the tombstone (sample 1) and a sample of plaster bulk was taken from a deep crack (sample 2) to confirm the expected composition of the substrate. Samples 3 to 12 were taken so to be representative of the stratigraphy of the object from the surface to 0.5–1.0 mm towards the core of the object.

Dust

Dirt deposited on the tombstone was sampled (Sample 1) and analysed to have a better understanding of the contaminants present in Gallery 46A, and eventually be able to discriminate which elements found on the surface can be attributed to the environmental dust in the galleries. Under the microscope, the sample seems to be mostly made of fibres of various colours (red, green, blue, Fig. 5). FT-IR analysis highlighted that the dust is made mostly of sulfates (such as gypsum and other varieties) with the asymmetric bending modes of SO$_4$ at 435, 600 and 667 cm$^{-1}$, the SO$_4$ symmetric and asymmetric stretching at about 1005 and 1105 cm$^{-1}$ and the $\nu_2$ H$_2$O of the sulfates at about 1600 and 1680 cm$^{-1}$, due to the combination of bending and vibration modes of H$_2$O ($\nu_1 + \nu_3$ and $2\nu_3$) related to the presence of gypsum [35, 36, 45, 46]. Aluminium (Al) was detected by EDS in all the layers of all the samples analysed and, as in the samples analysed as cross-sections, Al can be also due to the use of an Alumina suspension (Agar Scientific Micro-polish Alumina 0.3 µm—B8226) to obtain the final polish. Al is likely present, together with potassium (K) and silicon (Si) as part of silicate inclusions (visible in the EDS mapping and spectra, Fig. 7), which constitute clay minerals [47] and are reported to be present as a natural contaminant of mineral gypsum [2, 48]. FT-IR spectra suggest vibrations in the 950–1100 cm$^{-1}$ region, characteristic of clay minerals (Si–O containing minerals, such as kaolin, Al$_2$Si$_2$O$_5$(OH)$_4$) [40, 47]. FT-IR peaks at 1440 and 1770 cm$^{-1}$ (v CO$_3$) and overtone centred at 2400 cm$^{-1}$ (v CO$_3$) of the sulfates, and also the sulfate overtones centred on 2220 cm$^{-1}$ area, due to the combination of bending and vibration modes of H$_2$O ($\nu_1 + \nu_3$ and $2\nu_3$) related to the presence of gypsum [35, 36, 45, 46]. Overall, the sulfates and carbonates can be due to the building works, but the fibres and organic contribution derive from the visitor interactions.

Plaster bulk

The substrate (layer 0 in samples 3 to 13, as described in Table 1) largely consists of gypsum plaster (calcium sulfate, CaSO$_4$·0.5H$_2$O), confirmed by EDS spectra and mapping (see for example sample 3 in Fig. 6). XRD analysis on all the samples also confirmed that the mineral gypsum is the main component in the sample, as shown for example in the diffractogram of sample 3 (Fig. 6) and indicated by the characteristic diffraction peaks in Table 2 and consistent with the gypsum references available in published databases [24, 25]. FT-IR spectra also showed the presence of gypsum plaster with the peaks at about 1005 and 1105 cm$^{-1}$ (SO$_4$ symmetric and asymmetric stretching) and 1600 and 1680 cm$^{-1}$ (v$_2$ H$_2$O of the sulfates), and also the sulfate overtones centred on 2220 cm$^{-1}$ area, due to the combination of bending and vibration modes of H$_2$O ($\nu_1 + \nu_3$ and $2\nu_3$) related to the presence of gypsum [35, 36, 45, 46]. aluminium (Al) was detected by EDS in all the layers of all the samples analysed and, as in the samples analysed as cross-sections, Al can be also due to the use of an Alumina suspension (Agar Scientific Micro-polish Alumina 0.3 µm—B8226) to obtain the final polish. Al is likely present, together with potassium (K) and silicon (Si) as part of silicate inclusions (visible in the EDS mapping and spectra, Fig. 7), which constitute clay minerals [47] and are reported to be present as a natural contaminant of mineral gypsum [2, 48]. FT-IR spectra suggest vibrations in the 950–1100 cm$^{-1}$ region, characteristic of clay minerals (Si–O containing minerals, such as kaolin, Al$_2$Si$_2$O$_5$(OH)$_4$) [40, 47]. FT-IR peaks at 1440 and 1770 cm$^{-1}$ (v CO$_3$) and overtone centred at 2400 cm$^{-1}$ suggest the presence of calcite (calcium carbonate, CaCO$_3$), as suggested in other studies [26, 40, 43–45]. Minor variations of the position of the
| Sample | Type | Notes | No. of layers | Layers | Plaster | Interface | Coating | Overpaint |
|--------|------|-------|---------------|--------|---------|-----------|---------|-----------|
| 1      | Dust | Fibres and mixture of organic and inorganic material | – | – | – | – | – | – |
| 2      | Plaster from the inner bulk | – | – | – | Calcium sulfate, calcium carbonate, clay minerals, Al, Mg, Si, diterpenic resin | – | – | – |
| 3      | Cross-section | 3 | 1. Yellowish and undefined layer, 2. Dark layer | Calcium sulfate, calcium carbonate, clay minerals, Mg, diterpenic resin | Intermediate of plaster and dark layer | Mostly Si and Al, Fe, Ti and traces of Na, Cl and K, diterpenic resin | – |
| 4      | Cross-section | Edge area of sampling | 4 | 1. Yellowish and undefined layer, 2. Dark layer, 3. Purple layer | Calcium sulfate, calcium carbonate, clay minerals, Mg, diterpenic resin | Intermediate of plaster and dark layer | Mostly Si and Al, traces of K, Ba, Zn, Cl and P, diterpenic resin | Fe, Si and Ti, traces of K, Ba, Zn, Cl and P, drying oil modified alkyd paint |
| 5      | Cross-section | – | 1. Yellowish and undefined layer, 2. Dark layer | Calcium sulfate, calcium carbonate, clay minerals, Mg, Ti, diterpenic resin | Intermediate of plaster and dark layer | Mostly Si and Al, traces of Mg, Ti, Na, Fe and K, diterpenic resin | – |
| 6      | Cross-section | Area of repair | 3 | 1. Yellowish and undefined layer, 2. Dark layer | Calcium sulfate, calcium carbonate, clay minerals, Mg, Ti, diterpenic resin | Intermediate of plaster and dark layer | Mostly K, Fe, Si and Al, traces of Na, Pb and Cl, organic medium | – |
| 7      | Cross-section | Area of repair | 2 | 1. Yellowish and undefined layer with large orange and black particles | Calcium sulfate, calcium carbonate, clay minerals, organic medium | – | Mostly Fe and Ba, traces of Mg, K and Cl, organic medium | – |
| 8      | Cross-section | Fluorescent varnish | 4 | 1. Yellowish and undefined layer, 2. Dark layer, 3. Varnish | Calcium sulfate, calcium carbonate, clay minerals, Mg, Cl, diterpenic resin, possibly additional triterpenic resin or birch | Intermediate of plaster and dark layer | Mostly Si and Al, traces of K, Fe and Na, diterpenic resin, possibly additional triterpenic resin or birch | – |
Table 1 (continued)

| Sample Type | Notes | No. of layers | Layers | Plaster | Interface | Coating | Overpaint |
|-------------|-------|---------------|--------|---------|-----------|---------|-----------|
| 9 Cross-section | 3  | 2. Dark layer | Calcium sulfate, calcium carbonate, clay minerals, Mg, Al, Sr and Fe, diterpenic resin, possibly additional triterpenic resin or birch | Intermediate of plaster and dark layer | Mostly Si, Al, Mg, Sr and Fe, traces of K and Ti, diterpenic resin or birch | – |
| 10 Cross-section | 3  | 2. Dark layer | Calcium sulfate, calcium carbonate, clay minerals, Mg, organic medium | Intermediate of plaster and dark layer | Mostly Si and Al, traces of K and Cl, organic medium | – |
| 11 Cross-section | 4  | 3. Varnish | Calcium sulfate, calcium carbonate, clay minerals, Mg, organic medium | Intermediate of plaster and dark layer | Mostly Si and Al, traces of Fe and Mg, organic medium | – |
| 12 Cross-section | 4  | 3. Varnish | Calcium sulfate, calcium carbonate, clay minerals, Mg | Intermediate of plaster and dark layer | Mostly Si and Al, traces of P, Cl and Fe, organic medium | – |
| 13 Cross-section | 4  | 3. Varnish | Calcium sulfate, calcium carbonate, clay minerals, Mg, diterpenic resin | Intermediate of plaster and dark layer | Mostly Si and Al, traces of Fe, Mg, Cl, Ti and Na, diterpenic resin | – |
Fig. 6 EDS mapping and spectra (a) showed that sample 3 is made of calcium (Ca), sulfur (S) and oxygen (O) (calcium sulfate). Carbon (C) is present as detected from different sources (environment, casting polyester resin and organic media). XRD diffractogram (b) shows the characteristic peaks of gypsum, as also confirmed by comparison with the RUFF database [24] and COD Database [25].
FT-IR peaks related to the inorganic components can be observed for several reasons, one of which is the local substitution of elements such as Magnesium (Mg) and lead (Pb) in the gypsum and other minerals’ structure [35, 40, 44, 45, 47]. Mg was also detected in all the samples by EDS and it is possibly an exchangeable element of the sulfate variety MgSO₄ (more or less hydrated, chalcanthite CuSO₄·5H₂O, kieserite MgSO₄·H₂O, starkeyite MgSO₄·4H₂O, hexahydrite MgSO₄·6H₂O, epsomite MgSO₄·7H₂O, kieserite MgSO₄·H₂O, epsomite MgSO₄·7H₂O and meridianite MgSO₄·11H₂O). Small quantities of sulfate varieties such as barite (BaSO₄), celestite (SrSO₄), anglesite (PbSO₄) and the Mg varieties mentioned above can be naturally present in the gypsum quarries or form after the rehydration of the gypsum plaster that occurs after the addition of water to the calcined powdered gypsum plaster [2, 47, 48]. These secondary mineral phases are commonly found in minerals [47] and, for example, suggested by trace studies on stucco objects [21]. This composition of the plaster was consistently found in layer 0 of samples 3–8 and 10–13 (Table 1). Sample 5 shows additional small inclusions of calcium carbonate are present in the bulk, the object is largely made of gypsum plaster (made from gypsum) rather than lime plaster (made from calcite), which is instead often used for outdoor architectural details [4, 6], and was also found in the analysis of other casts [35]. Sample 2 was taken from the inner plaster bulk (about 5 cm from the surface), exposed on a deep crack of the tombstone, to understand whether the organic coating visible on the surface was also used as an additive in the plaster mixture, as suggested as possible by the historical literature [2, 48–50]. No organic materials were detected in sample 2 by FT-IR. Py-TMAH-GC/MS analysis indicated the presence of abietane skeleton diterpenoids due to the occurrence of the peaks at m/z 315, 299, 285, 253 and 239 (Fig. 8 and Table 3) [29, 51]: 7-Oxodehydroabi-etic acid, methyl ester at retention time, RT, 25.47 min and methyl dehydroabietate at 25.57 min in the chromatogram. The fragments at m/z 314 and 253 suggest the presence of a compound formed by oxidation of pine resin biomarkers, as suggested in several case studies [29, 51]: 7-Oxodehydroabi-etic acid, methyl ester at retention time, RT, 25.47 min and methyl dehydroabietate at 25.57 min in the chromatogram. The fragments at m/z 314 and 253 suggest the presence of a compound formed by oxidation of pine resin biomarkers, as suggested in several case studies [29, 51]: 7-Oxodehydroabi-etic acid, methyl ester at retention time, RT, 25.47 min and methyl dehydroabietate at 25.57 min in the chromatogram. The fragments at m/z 314 and 253 suggest the presence of a compound formed by oxidation of pine resin biomarkers, as suggested in several case studies [29, 51]: 7-Oxodehydroabi-etic acid, methyl ester at retention time, RT, 25.47 min and methyl dehydroabietate at 25.57 min in the chromatogram.

### Table 2 XRD peaks observed in the diffractograms of samples 2 and 3 and found in the RUFF database gypsum references R060509 and R040029

| Sample 2 | Sample 3 | R060509 | R040029 |
|----------|----------|---------|---------|
| 11.6304  | 11.67299 | 11.6688 | 11.6553 |
| 20.6882  | 20.74514 | 20.7682 | 20.7582 |
| 23.3816  | 23.37026 | 28.1588 | 28.1498 |
| 26.8718  |          |         |         |
| 29.121   | 29.10739 | 29.1507 | 29.1396 |
| 31.0975  | 31.12382 | 32.09465|         |
| 33.4643  | 33.34622 |         |         |
| 34.53655 | 34.6194  |         |         |
| 35.94149 | 35.9997  |         |         |
| 36.65667 | 36.0027  | 36.65   |         |
| 37.3434  | 37.4189  | 37.4124 |         |
| 40.5746  | 40.63457 |         |         |
| 43.4695  | 43.38031 | 42.2079 |         |
| 44.1625  |          | 44.2309 |         |
| 45.8294  | 45.51888 | 45.5453 | 45.5268 |
| 46.495   |          | 46.4606 |         |
| 47.8711  | 47.85724 | 47.8955 | 47.8848 |
| 48.35782 | 48.4221  | 48.3983 |         |
| 48.72981 |          |         |         |
| 50.32856 |          |         |         |
| 50.73068 |          |         |         |
| 51.34237 |          |         |         |
| 54.41229 |          |         | 54.4676 |
| 55.14868 | 55.2183  |         | 55.191  |
| 55.81309 |          |         | 55.8618 |
| 56.6775  | 56.7291  |         |         |
| 58.2058  | 58.19157 |         | 60.3599 |
| 64.75298 |          | 65.8196 |         |
| 67.0222  | 66.7291  |         | 66.7077 |
| 68.9538  | 68.66935 |         |         |
| 70.65909 |          |         |         |
| 71.4685  | 71.22731 |         |         |
| 74.11747 |          |         |         |
| 76.56741 |          |         |         |
| 77.39589 |          |         |         |
| 79.61351 |          |         |         |
sources of lipids or naturally present in the resin [28, 29, 51]. It was suggested in other studies [52, 53] that the intensity of the FA peaks can change, as affected by matrix effects due to the presence of inorganic pigments. This would change the P/S ratio and therefore invalidate the correlation which allows the determination of the type of lipid. Due to the lack of previous studies on the effects of the predominance of the inorganic portion over the organic component on the areas of the FA peaks in the chromatogram, further research on this topic is required. Due to the complexity of the mixture, as often happen in the case of cultural heritage materials, separation methods are required prior the mass analysis so that several components arrive in the ion source one at a time [29]. For this reason, other peaks that can be assigned to small fragments of amine and lipids between 4.18 and 16.99 min in the chromatograms have not been considered diagnostic in this sample (Fig. 8). The full record of GC/MS results is provided in the dataset [23]. The inconsistency of the FT-IR and py-TMAH-GC/MS results in sample 2 may suggest that a very small quantity of resin was added in the plaster [2, 48] or that such small amount had penetrated deep in the plaster from the surface. It is also possible that the organic material derived from accidental contamination in the studio of the plasterer or it might have even been transferred from the mould during the casting process.

An interface layer, that appears yellow under visible light reflectance and possibly consists of a portion of the lower layer soaked with the surface coating(s), showing characteristics of both layers, is visible in samples 3–6 and 8–13 (Fig. 9). The tabular crystalline structure typical of gypsum plaster can be seen in the substrate layer and the interface layer in the BSE image [21, 47, 54]. In samples 10 and 11 the casting resin homogeneously penetrated the stratigraphy and in samples 3, 4, 5, 6, 8, 12 and 13 layer 1 appears ‘denser’ than layer 0 (Table 1), suggesting that the coating layer had penetrated the plaster, filling the pores and impeding the casting resin occupying the voids and possibly preserving the mineral structure of the gypsum plaster (Fig. 9). This was visible in the BSE image as well as through the EDS mapping (casting polyester resin and catalyst are mainly made of organic compounds C, H and O). The presence of the casting resin (polyester resin) was also detected in all the layers by FT-IR spectra (\(\delta\) OH phenol at about 1312 cm\(^{-1}\) and \(\delta\) CH aromatic at about 1760 cm\(^{-1}\), as can be seen in the reference in Fig. 10).

**Coating**

Samples 3–5 and 8–13 show a coating layer, that appears dark under visible light reflectance (layer 2 in these samples as described in Table 1), and it likely consists of a pigmented organic medium and dirt. EDS mapping and spectra show that this dark layer contains calcium sulfate but consists mostly of Si and Al (Fig. 11). Traces of other elements can be seen and the composition varies from sample to sample (Table 1), suggesting that the trace elements in the surface layers derive from both the plaster substrate and the dirt deposited onto the surface. The presence of the polyester casting resin (Fig. 10) and gypsum plaster (CaSO\(_4\)_0.5 H\(_2\)O, characteristic peaks as described above) was detected in all the layers by FT-IR, including the surface. The presence of an
organic medium is indicated in all the FT-IR spectra and the position of the peaks suggest that is likely a wax or resin (CH bending and stretching); however, the crowded appearance of the spectra and the broadness of the peaks impede the unique assignment of such contributions. The several peaks that can be identified in the area over

![Fig. 8 py-TMAH-GC/MS chromatogram of sample 2 in the full retention time range, 0–40 min (a) and a narrower range, 16–30 min (b). The peaks up to 16.99 min are due to smaller lipid and amine fragments which are not diagnostic in this sample. The identification of pine resin was possible upon determination of the following markers: dimethyl azelate 1, methyl palmitate 2, methyl stearate 3, 7-oxo-dehydroabietic acid methyl ester 4 and methyl dehydroabietate 5. The mass spectra of compounds 4 (c) and 5 (d) are also shown.](image)
3000 cm\(^{-1}\) cannot be considered uniquely diagnostic, as OH and NH stretches occur in this region and once again due to the complexity of the mixture, the water present in the crystals and the pores of the plaster, as well as in the organic components will add up in this area. Calcium oxalate might be also present in layer 2, but it was not possible to uniquely assign its peaks, as usually close to the vibrations characteristic of calcium sulfate, as also suggested in other studies [46]. Py-TMAH-GC/MS of samples 3, 4, 5, 8, 9 shows the markers characteristic of a diterpenic resin (similarly to sample 2, Fig. 8), such a resin or pine resin, possibly mixed with a non-drying oil or another source of lipids, similar to what is suggested in relevant literature [27–29].

Py-TMAH-GC/MS of samples 8 and 9 shows peaks that were assigned, as based on previous studies [29, 55], to a betullin-like triterpene marker (at \(t = 30.08–33.49\) min, Table 3), suggesting that, as rarely are diterpene and triterpene molecules are rarely found together in a plant resin, a birch, dammar or mastic is additionally

### Table 3 Most relevant py-TMAH-GC/MS mass spectrum fragmentation peaks for the materials characterization across the samples

| RT (min) | m/z          | Assignment                                      |
|---------|--------------|-------------------------------------------------|
| 16.25   | 55, 69, 74, 83, 87, 111, 129(100), 138, 171, 188 | Dimethyl phthalate                              |
| 17.50   | 74(100), 87, 97, 111, 143, 152, 178, 185      | Dimethyl azelate                                |
| 21.68   | 74(100), 87, 101, 129, 145, 185, 227, 270     | Methyl palmitate                                |
| 23.60   | 74(100), 87, 129, 143, 199, 255, 298          | Methyl stearate                                 |
| 25.47   | 187, 207, 253(100), 313, 328                   | 7-Oxodehydroabietic acid, methyl ester          |
| 25.57   | 141, 155, 197, 239(100), 253, 314              | Methyl dehydroabietate                          |
| 30.08   | 79(100), 121, 138, 160, 189, 205, 442          | Betullin                                         |

![Fig. 9](image) High vacuum SEM–EDS (1.5 × 10\(^{-2}\) Pa) of sample 2 allowed a higher magnification; BSE image (c) shows the tabular structure of the gypsum plaster. SEM–EDS Low Vacuum Mode (10–15 Pa) of the samples cast in resin allowed to acquire the BSE image of the full stratigraphy. In sample 12 the tabular structure can be observed in layers 0 and 1 and layer 1 appears denser in the BSE image (b) and has a yellow tone under VLR OM (a). In sample 10 the BSE image (d) shows that the casting resin was absorbed evenly in the pores.
present in the stratigraphy. The full fragmentation patterns of the markers characteristic of dammar and mastic were not detected, but this can be due to the natural degradation of the molecules or to the derivatization processes applied to the already degraded organic material, as suggested in [29, 56, 57]. The small quantity of the organic material, when compared to the inorganic portion, also contributed to the small relative abundance of the di- and tri-terpene molecules in the chromatogram. The full record of GC/MS results is provided in the dataset [23].

The contributions of the organic material as shown in the FT-IR spectra of all the layers indicates that either the material was added to the wet gypsum plaster wet mixture or that the coating has also penetrated in the bulk. The latter seems also possible as the average depth of the samples is about 0.5 mm. The characterization of unknown organic materials in aged samples has been recognised as the most challenging application of GC/MS techniques [29]. Moreover, in sample 8 an additional layer (layer 3) fluoresces white under UV illumination, suggesting that these could be an additional layer or that layer 2 has not been absorbed evenly, as seen in the other samples. The same was also observed in samples 11, 12 and 13.

**Areas of retouching**

Sample 4 is characterised by the same stratigraphy described above but presents an additional layer (layer 3), purple under visible illumination (Fig. 12). EDS mapping suggested that this layer is mostly made of iron (Fe), silicon (Si) and titanium (Ti). Py-TMAH-GC/MS
Fig. 11  Sample 11: the surface layer appears dark under VLR OM (a) and the interface layer is yellow under VLR OM. No characteristic fluorescence was observed in the UV-f OM (b). EDS mapping shows that the surface layer is mostly made of Si and Al. FT-IR analysis (c) shows a broad band that suggests the presence of gypsum, casting resin and an organic medium.

Fig. 12  Sample 4's VLR OM (a) and BSE images (b) and EDS mapping. The sample was taken from an area of overpaint. Al = aluminium, Si = silicon, Ti = titanium, Fe = iron.
of this sample highlighted that a drying oil modified alkyd paint is the medium in this layer, suggesting that they were applied after the 1920s [30, 58, 59]. Dimethyl phthalate was detected at t = 16.25 min in the chromatogram (Table 3) and was considered a marker for aged alkyd paints mixtures, as it is reported to be forming in these paints upon ageing [30, 31]. Phthalic compounds can indicate the presence of a range of different twentieth-century materials. In art conservation they can be found, for example, in alkyd paints, polyvinyl adhesives and BEVA®371 and cellolyn [60]. Given that the phthalates were detected in the sample showing an additional refill and retouching layer and that V&A conservators have confirmed that polyvinyl adhesives, BEVA®371 and cellolyn are not documented as used in the Cast Courts, it is postulated that the dimethyl phthalate is due to the presence of alkyd paints. Samples 6 and 7 broadly present the same stratigraphy of the other samples, however, the coating layer is made of K, Fe, Si and Al and traces of Na, Pb and Cl in sample 6 and Fe and Ba and traces of Mg, K and Cl in sample 7. In samples 8, 11, 12 and 13 an additional surface layer (layer 3 in these samples) fluoresces milky-white under UV illumination. No differences can be seen in the FT-IR spectra of these samples nor the GC/MS chromatograms, suggesting that either layer 3 is an additional layer of varnish made of the same material of the medium of layer 2 (diterpene resin) or layer 3 is an unabsorbed portion of the medium of layer 2 visible on the surface.

Manufacturing as suggested by the stratigraphy
In summary, the stratigraphy of the samples is quite consistent, featuring a ‘substrate’ layer, an ‘interface’ layer and a coating ‘dark’ layer, likely a combination of aged coating and dust. On top of these, some samples feature additional layers, having been overpainted or having been sampled from an area of repair. The bulk of the object is made of gypsum plaster, which contains several types of inclusions (including silicates and carbonates). By looking at the results, it is possible to hypothesise that in the surface layer, containing silicon and aluminium, but also traces of other elements, the medium is a diterpene resin. Areas of repairs consist of overpaints made of alkyd paint, suggesting that they were applied after the 1920s [30, 58, 59], and inpaints containing silicon or barium. Areas showing an additional varnish layer might have locally highlighted or the additional layer might be due to a local difference in the surface absorbance or to an accidental spillage (as for example in sample 12, see Fig. 3).

A summary of the results can be seen in Table 1 and has also been compiled in a comprehensive dataset [23].

Conclusions
A multi-analytical approach allowed the characterization of the surface coatings of the object. A ‘substrate’ layer made of gypsum plaster, an ‘interface’ layer and a coating ‘dark’ layer, likely a combination of aged diterpenic resin coating and dust were identified. Overpaint made of alkyd paint and areas of repair were also highlighted. Due to the immense variety of recipes for the manufacturing of the nineteenth-century plaster casts, the stratigraphy of a plaster cast might result in a complex combination of organic media and inorganic features, even more complex when modern treatments have been applied for the care of the cast. As a multi-analytical approach in studies on similar materials has been proven effective, the combination of techniques for the characterization of inorganic and organic components is fundamental in plaster artefacts. This study demonstrated that the suggested methodology for the characterisation of the coatings of historical plaster casts can provide information on nineteenth-century manufacturing. This case study suggests an analytical protocol that combines diverse methods to characterise the manufacturing of such artefacts. The investigation is ongoing and additional elements are still needed to have a comprehensive understanding of the cast manufacture and history, but the results discussed and summarised here demonstrate that despite the many analytical challenges of studying the complex composition of the cast, a thorough and comparative study can unveil the secrets of this nineteenth-century art. Further investigation is required to study the deterioration of those compounds on the sub-molecular level.

Abbreviations
BSE: Back Scattered Electron; EDS: Energy Dispersive Spectrometry; FT-IR: Fourrier-Transform Infra-Red; FPA: Focal Plane Array; GC/MS: Gas Chromatography–Mass Spectrometry; OM: Optical Microscopy; PL: Proper Left; PR: Proper Right; py: Pyrolysis; SEM: Scanning Electron Microscopy; TMAH: Tetramethylammonium hydroxide; UVF: Ultraviolet Fluorescence; VLR: Visible Light Reflectance; XRF: X-ray Fluorescence.

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Authors’ contributions
Analyses were performed at the Northumbria University laboratories by VR under the supervision of C.T. The project was initiated by CT and CH, who supervised the work in the Galleries. JP provided technical observations on the object. All authors read and approved the final manuscript.

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Availability of data and materials
All data generated during this study are discussed in this published article and are fully available at the Figshare permanent data link: https://doi.org/10.25398/rd.northumbria.13469925.v3 [23].

Declarations
Competing interests
The authors declare that they have no competing interests.

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