A combined physical–chemical and microbiological approach to unveil the fabrication, provenance, and state of conservation of the Kinkarakawa-gami art

Elena Piacenza\textsuperscript{1,2}, Alessandro Presentato\textsuperscript{2,}\textsuperscript{*}, Francesca Di Salvo\textsuperscript{3}, Rosa Alduina\textsuperscript{2}, Vittorio Ferrara\textsuperscript{1,3}, Valeria Minore\textsuperscript{2}, Antonio Giannusa\textsuperscript{4}, Giuseppe Sancataldo\textsuperscript{5} & Delia Francesca Chillura Martino\textsuperscript{1,2,}\textsuperscript{*}

\textit{Kinkarakawa-gami} wallpapers are unique works of art produced in Japan between 1870 and 1905 and exported in European countries, although only few examples are nowadays present in Europe. So far, neither the wallpapers nor the composing materials have been characterised, limiting the effective conservation–restoration of these artefacts accounting also for the potential deteriorogen effects of microorganisms populating them. In the present study, four \textit{Kinkarakawa-gami} wallpapers were analysed combining physical–chemical and microbiological approaches to obtain information regarding the artefacts’ manufacture, composition, dating, and their microbial community. The validity of these methodologies was verified through a fine in blind statistical analysis, which allowed to identify trends and similarities within these important artefacts. The evidence gathered indicated that these wallpapers were generated between 1885 and 1889, during the so-called industrial production period. A wide range of organic (proteinaceous binders, natural waxes, pigments, and vegetable lacquers) and inorganic (tin foil and pigments) substances were used for the artefacts’ manufacture, contributing to their overall complexity, which also reflects on the identification of a heterogeneous microbiota, often found in Eastern environmental matrices. Nevertheless, whether microorganisms inhabiting these wallpapers determined a detrimental or protective effect is not fully elucidated yet, thus constituting an aspect worth to be explored to deepen the knowledge needed for the conservation of \textit{Kinkarakawa-gami} over time.

\textit{Kinkarakawa-gami} represents an important example of Japanese art and its mixture within the European culture between the XIX and XX centuries. This manufacture derived from the importation in Japan of the so-called \textit{Kinkarakawa} (i.e., foreign gilt leather) by the Dutch East India Company, which was one of the few foreign companies allowed to commercially trade with this eastern country up to 1854\textsuperscript{1}. Subsequently, the opening of Japan to the western world encouraged the exportation of authentic original Japanese goods, among which

\textsuperscript{1}National Interuniversity Consortium of Materials Science and Technology (INSTM), UdR of Palermo, 50121 Florence, Italy. \textsuperscript{2}Department of Biological, Chemical and Pharmaceutical Sciences and Technologies (STEBICEF), University of Palermo, 90128 Palermo, Italy. \textsuperscript{3}Department of Agricultural, Food and Forest Sciences, University of Palermo, 90128 Palermo, Italy. \textsuperscript{4}Vincenzo Ragusa and O’Tama Kiyohara Museum, Liceo Artistico Statale Vincenzo Ragusa e O’Tama Kiyohara, 90129 Palermo, Italy. \textsuperscript{5}Department of Physics and Chemistry (DIFC) Emilio Segrè, University of Palermo, 90128 Palermo, Italy. \textsuperscript{*}email: alessandro.presentato@unipa.it; delia.chilluramartino@unipa.it
Kinkarakawa-gami (i.e., gilt leather-like wallpapers) gained international relevance and recognition, mostly with British, Dutch and German companies. Several European sources of the XIX century documented the traditional production stream of Kinkarakawa-gami, identifying four steps: (1) hand-making of the leather-like paper, (2) its embossing with a chosen pattern, (3) gilding of the paper with metal powders or foils, and (4) its final decoration. Historically, three periods of Kinkarakawa-gami’s manufacture are distinguished: (1) the Japonist Enlightenment (1873–1884), during which the wallpapers were entirely hand-made, properly depicting Japanese patterns and featuring an “oily smell”, which did not encounter the European consumer taste; (2) the period of industrial production (1882–1889), when several British factories were established on Japanese soil, leading to faster yet traditional manufacture, as well as a change in the wallpaper design; and (3) the mass-produced commodity period (1890–1905), where the patterns were directly chosen by British designers and cheap industrial processes were applied to answer to the large demand of European customers, however resulting in the end—or almost all—of the Kinkarakawa-gami art.

The uniqueness of these Japanese wallpapers relies on their limited presence in Europe, therefore physical–chemical studies aimed to unveil the features of these artefacts are still missing, being of paramount importance to develop successful and sound conservation-restoration strategies. This aspect significantly gains importance since, to the best of the authors’ knowledge, Kinkarakawa-gami belonging to the Vincenzo Ragusa and O’Tama Kiyohara Collection of Palermo (patrimony of the Liceo Artistico founded in the XIX century by the sculptor Vincenzo Ragusa) represents the first finding of this form of Japanese fine art in Italy. Thus, four wallpapers of this collection were studied through a multidisciplinary approach aimed to overview their state of conservation for their future restoration, therefore enhancing the artistic value of these artefacts. The wallpaper manufacture was evaluated through a standard operating procedure involving non-invasive and non-destructive physical–chemical techniques—that allow to perform an extensive sampling of the artefacts without altering them or their conservation state—and microscopy observations, focusing on the (1) paper composition, (2) metallic layer on the recto of these artefacts, identification of (3) inorganic pigments, (4) organic pigments, and (5) substances (e.g., lacquers) exploited as binders for the pigments. Physical–chemical results were validated through in blind statistical analyses, which, by identifying homogeneous clusters of curves with respect to their shape, further unveiled important differences and similarities among the four artefacts. Finally, the cultivable microbiota populating these wallpapers and their potential interaction with both inorganic and organic substrates were critically discussed to better understand how to properly restore and preserve these Kinkarakawa-gami through time.

Results and discussion

Paper composition. The support material used by Japanese artisans to produce the Kinkarakawa-gami wallpapers (henceforth named as INV_11, INV_13, INV_15, and INV_20; Fig. 1, Table 1) derived from either eastern plants or wood pulp, as indicated by the ubiquitous detection, through micro Attenuated Total Reflectance-Fourier Transform Infrared (µATR-FTIR) spectroscopy, of vibrational modes typical of lignin and cellu-
lose (Fig. 2, Table 2, Supplementary Tables 1–4). The chemical similarity with wood pulp was also suggested by a statistical analysis performed on the µATR-FTIR spectra, which was carried out focusing on the 3770–2750 cm⁻¹ and 1840–720 cm⁻¹ intervals (Supplementary Fig. 1), where the shape of the curves gave information regarding the essential modes of variability among the spectra and their clustering. The cluster analysis showed the hierarchical relationship between the spectra (Figs. 3a, 4a), based on which the optimal number of clusters and the spectra allocation were determined (Figs. 3b, c, 4b, c). Principal modes of variations were derived to identify cluster effects on the shape of the curves, showing that the clusters were correctly visualized in an R² space, as the first two functional Principal Components (PCs) described the 90.8% and 82.6% of the total variability for the first and second interval, respectively (Figs. 3d, 4d). Specifically, cluster 2 (first interval) and 1 (second interval) grouped the spectra collected for wallpapers’ verso, whose high values of silhouette (average sᵢ = 0.73 and 0.76) and distribution in the R² space through PCA assessed their well-identification within the cluster (Fig. 3c–e, 4c–e). The large PC2 value of the first IR interval of INV_15_2 (Fig. 3d) indicated its greater variability than the other verso spectra, likely due to the presence of multiple –OH stretching modes characteristic of cellulose (Fig. 2c; Supplementary Table 3). The Modified Band Depth (MBD) function was used to identify a representative (i.e., deepest) curve for the clusters (Figs. 3f, 4f) describing the shape of the spectra and arrange them by resemblance from the centre (highest depth) outwards (lowest depth) (Supplementary Fig. 2, 3). INV_13_2 was the deepest curve of cluster 2 (first interval) and the other verso spectra featured high depth values, constituting the core (70%) of the cluster (Supplementary Fig. 2b). An analogous situation was observed for the second interval, although the most representative curve was INV_13_9.1 (Fig. 4f), confirming the high degree of similarity of the wallpapers’ verso. Moreover, morphological and structural features of the wallpapers’ verso were studied through Confocal Laser Scanning Microscopy (CLSM) (Fig. 5), revealing a strong fluorescence signal in the visible-light spectrum, and a tangled matrix of organic fibres with a lateral width between 10 and 20 µm for all the samples (Fig. 5a, Supplementary Video 1), which, along with the cellulose and lignin contributions detected within µATR-FTIR spectra (Supplementary Table 1–4), suggested the vegetable origin of the paper. However, it cannot be unequivocally indicated whether the paper was hand-made or made out of mechanical wood pulp, although the detection of silicon (Si) traces by X-Ray Fluorescence (XRF) spectroscopy in the verso sampling points (Fig. 6a, b) could indicate the use of sandstone grinders for the mechanical generation of the pulp. Further, the detection of Si, calcium (Ca) (Fig. 6a, b), and IR absorption bands typical of silicates or carbonates (Supplementary Table 1–4) might infer the use of quartz, calcite, or chalk as fillers.

Table 1. Description of the sampling points of the four analysed wallpapers.
Traces of Sulphur (S) and manganese (Mn) were observed on the wallpapers’ verso, although S was solely recognized in the not-written areas, except for INV_15, while a higher distribution of Ca and iron (Fe) was seen in this side as compared to the recto (Fig. 6a, b, Supplementary Fig. 4a). Since these two elements, along with lead (Pb), were preponderant in the written portions of INV_11 and INV_13 (Fig. 6b, Supplementary Fig. 4a), they may be components of the ink (e.g., iron gall ink) used for the selvages12, which are a trademark of Kinkarakawa-gami1. This conclusion cannot be drawn for INV_15_2 and INV_20_2, due to the strong presence of arsenic (As) and mercury (Hg) (Supplementary Fig. 4a), deriving from the coloured areas of the recto (Fig. 6c, Supplementary Fig. 5b) and/or a metal diffusion from one overlapping wallpaper to another, which may have hidden Ca and Fe signals.

Gilding of the paper with metals. Tin (Sn) was ubiquitously present in the wallpapers, revealing a greater signal distribution on their recto than their verso (Fig. 6), which is coherent with the gilding technique used by Japanese artisans, who covered the paper surface with a lacquer mordant and a metal foil (i.e., silver, gold or tin) and adhered to the embossed pattern. In this regard, CLSM imaging of the wallpapers’ recto showed an amorphous and heterogeneous surface attributable to the artefact’s pattern, the inorganic pigments, and/or organic pigments (Fig. 5b). To better validate whether the wallpapers featured a multilayer structure as described in the literature1,2, a 3D reconstruction of a portion of INV_15 recto only partially covered by the motif pigmentation was performed (Supplementary Video 2). As a result, two distinct layers were detected: the top layer displayed the amorphous structure characterising the pigmentation, while the bottom layer featured an underlying matrix of fibres resembling that of the verso (Fig. 5c, d, Supplementary Video 2). Since the gilding practice was first carried out by Rottmann, Strome & Co factory in Yokohama (1882) yet abandoned from 1890 onwards2, the presence of multilayers containing Sn within the wallpapers suggested their belonging to the second period of Kinkarakawa-gami production.

Figure 2. µATR-FTIR spectra performed on the wallpapers. The spectra were collected on the selected sampling points (as indicated in the legend) of (a) INV_11, (b) INV_13, (c) INV_15, and (d) INV_20. For clarity, the µATR-FTIR spectra were offset of 0.10 arbitrary units (a.u.).
Kinkarakawa-gami’s decoration. Despite the published literature on Kinkarakawa-gami art, indications regarding inorganic and organic pigments, as well as binders used for artefacts’ decoration is not reported yet, hence constituting a gap to be filled to enable their conservation-restoration.

Inorganic pigments. XRF spectroscopy revealed the recurrent presence in the artefacts of Si, S, Fe, Mn, Zn, As, whereas Hg, copper (Cu), chromium (Cr), and Pb were detected in specific coloured portions (Fig. 6), indicating the use of several inorganic pigments for the decoration. Although most of these pigments have strong IR contributions below 600 cm$^{-1}$, some absorption bands attributable to the yellow pigment orpiment (As$_2$S$_3$, Sekio; 1035, 913, and 820 cm$^{-1}$) and its degradation (-AsOFe stretching mode 830–840 cm$^{-1}$) were frequently found within the µATR-FTIR spectra (Table 2, Supplementary Table 1–4), agreeing with the detection of As and S through XRF spectroscopy (Fig. 6). Since up to 1885 the wallpaper production did not involve As-compounds, these results suggested artefacts’ manufacture between 1886 and 1889, which is a rather tight timeline unlikely

| Type of paper                  | 11_3 | 11_4 | 11_6 | 11_7 | 13_2 | 13_3 | 13_5 | 13_8 | 13_9 | 15_2 | 15_4 | 20_3 | 20_4 |
|-------------------------------|------|------|------|------|------|------|------|------|------|------|------|------|------|
| Cellulose                     | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    |
| Lignin                        | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    |
| Fillers/inorganic pigments    |      |      |      |      |      |      |      |      |      |      |      |      |      |
| Silicates                     |      |      |      |      |      |      |      |      |      |      |      |      |      |
| Carbonates                    |      |      |      |      |      |      |      |      |      |      |      |      |      |
| Organic binders               |      |      |      |      |      |      |      |      |      |      |      |      |      |
| Unidentified binders          | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    |
| Proteinaceous binders         |      |      |      |      |      |      |      |      |      |      |      |      |      |
| Waxes                         |      |      |      |      |      |      |      |      |      |      |      |      |      |
| Urushi lacquer                | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    |
| Organic pigments              |      |      |      |      |      |      |      |      |      |      |      |      |      |
| Indigo                        | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    |
| Inorganic pigments            |      |      |      |      |      |      |      |      |      |      |      |      |      |
| Orpiment                       | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    | ✓    |
| Mordants                      |      |      |      |      |      |      |      |      |      |      |      |      |      |
| Green vitriol                 |      |      |      |      |      |      |      |      |      |      |      |      |      |

Table 2. Organic and inorganic substances detected through µATR-FTIR performed on INV_11, INV_13, INV_15, and INV_20. ✓ indicates the presence of the substance within the sampling point; for clarity, “INV_X_” (X = 11, 13, 15, or 20) was omitted from the sampling points’ acronyms.

Figure 3. Statistical analysis performed on µATR-FTIR spectra for the 3770–2750 cm$^{-1}$ interval. The hierarchical relationship between the spectra is represented by the dendrogram depicted in (a), while (b, c) show the estimation of the number of clusters using the average silhouette width and the obtained silhouette of clusters, respectively. PCA results, µATR-FTIR spectra, and the most representative (i.e., deepest) curve for each identified cluster (highlighted with different colours) are illustrated in (d–f). For clarity, the most external spectra of the clusters are underlined in (d) with the grey colour.
Fe ubiquity on the recto’s background (Supplementary Fig. 4b) may instead indicate the application of natural iron oxide pigments, such as yellow—*Odo*—, brown—*Taisha*—and red—*Kincha*—ochres15, to obtain gold-bronze shades, which is another distinctive feature of this Japanese art7. Moreover, these pigments contain silicates and to be achieved through other techniques (e.g., Accelerator Mass Spectroscopy (AMS)-radiocarbon dating featuring ± 20 years of experimental error)14.

Figure 4. Statistical analysis performed on µATR-FTIR spectra for the 1840–719 cm⁻¹ interval. The hierarchical relationship between the spectra is represented by the dendrogram depicted in (a), while (b, c) show the estimation of the number of clusters using the average silhouette width and the obtained silhouette of clusters, respectively. PCA results, µATR-FTIR spectra, and the most representative (i.e., deepest) curve for each identified cluster (highlighted with different colours) are illustrated in (d–f). For clarity, the most external spectra of the clusters are underlined in (d) with the grey colour.

Figure 5. Fluorescence microscopy of *Kinkarakawa-gami* wallpapers. CLSM imaging of representative portions of INV_15 on both verso and recto are illustrated in (A, B) respectively, while (C) depicts a representation of the 3D reconstruction performed on INV_15 recto, whose multilayer structure is quantitatively reported in depth-colour code (analysis depth 75 µm) in (D). Scale bars are 100 µm.
manganese dioxide (MnO₂) as either components or impurities, supporting the frequent detection of Si and Mn in traces (Fig. 6). An additional trace element observed in INV_11_7 and INV_13_3 was S (Supplementary Fig. 4b), whose contribution might rely on the use of tin sulphide (SnS₂), which was known in Asia since 300 BC, as a bronze-gold pigment for these backgrounds. The variability of the elemental composition in recto’s backgrounds was inferred by the detection of Zn (INV_11_5, INV_11_7, and INV_15_3), As (INV_15_3 and INV_20_5), Pb (INV_11_5), or Hg (INV_15_3; Fig. 6c). In this regard, the simultaneous presence of S and Hg (INV_15_3), and S and As (INV_20_5; Supplementary Fig. 4b) suggested that red mercury- (HgS; cinnabar) or lead oxide (PbSO₄) could be present in INV_11_4 and INV_13_5, as indicated by Pb traces, while INV_13_6 displayed a high variability in elemental composition (Fig. 6e), inferring that these portions could contain multiple pigments. Particularly, Pb was present only in INV_11, whose great variability in composition was further inferred by the detection of Cr and Cu (INV_11_3; Supplementary Fig. 5b), typical components of green pigments, as chromium oxide (Cr₂O₃), malachite green [Cu₂(CO₃)(OH)₂], Verdigris [Cu₂(CH₃COO)₂Cu(OH)₂], Scheele green [Cu(CH₃COO)₂3Cu(AsO₂)₂], green (INV_11_3), or S and As (INV_20_5; Supplementary Fig. 4b) suggested that red mercury- (HgS; cinnabar or Shu) and yellow/red arsenic-sulphides (Sekio or red realgar—Yuo, AsS or AsS₄) were the main pigments of INV_15 and INV_20 recto’s backgrounds. As and Zn traces in INV_15_3 could be also linked to Shu, which generally contains these elements as impurities. Furthermore, Zn-based paint driers may have been applied to favour the pigments’ deposition and solidification on the tin foil, while Chinese white (zinc oxide; ZnO) and zinc sulphide (ZnS) could be used as white pigments. To improve the brightness of the latter, artisans even mixed them with lead white [basic lead carbonate; 2PbCO₃·Pb(OH)₂], justifying the Pb contribution observed in INV_11_5 (Supplementary Fig. 4b).

The painted areas of the wallpapers’ recto were deeply analysed focusing on the most representative colorations (i.e., red, green, brown, and black). Similar chemical compositions were observed for the red colour in INV_11_6 and INV_15_4, where the featuring elements were Hg and S, indicating the utilization of Shu, while traces of Si were detected only in INV_11_6 (Fig. 6d, Supplementary Fig. 5a). Moreover, the large signal of Hg observed in both INV_11_6 and INV_15_4 might be responsible for the Sn underestimation, as the high atomic number of Hg can mask Sn contribution. The green-coloured areas of INV_11, INV_15, and INV_20 displayed instead a high variability in elemental composition (Fig. 6e), inferring that these portions could contain multiple pigments. Particularly, Pb was present only in INV_11, whose great variability in composition was further indicated by the detection of Cr and Cu (INV_11_3; Supplementary Fig. 5b), typical components of green pigments, as chromium oxide (Cr₂O₃), malachite green [Cu₂(CO₃)(OH)₂], Verdigris [Cu₂(CH₃COO)₂3Cu(AsO₂)₂], Scheele green (a mixture of CuHAsO₄ and CuAsO₄), or malachite green [Cu(CH₃COO)₃Cu(AsO₂)₂] [17]. The presence of Pb may be instead ascribed to (1) lead white for the outline of the leaves [15], (2) lead–tin yellow [21], or (3) yellow lead oxide (PbO₂) [22] to achieve diverse green shades. Regardless, all the green areas exhibited S and As, whose distribution varied depending on the different colour shades, as noted in INV_11_3, INV_11_9, INV_13_4, and INV_13_8 (Supplementary Fig. 5b). Thus, Sekio [18] seemed to be constantly present in the green areas, suggesting (1) a combined use of Verdigris, Scheele green, or malachite green with sulphide compounds in INV_11_3 to improve both brightness and durability of the pigments [24], as well as (2) the addition of Sekio to an organic dark (blue/purple) pigment to obtain the green colour [15,25,26]. Except for INV_20, the intense Fe signal characterising the recto’s green-painted sections (Fig. 6e) could imply the combination of green/yellow inorganic pigments with natural ochres, as also supported by the detection of Si, Mn, and Zn in traces (Fig. 6e, Supplementary Fig. 5b). Nevertheless, μATR-FTIR spectra of INV_20_3 exhibited –SO₄ asymmetric (1120–1040 cm⁻¹ and 827 cm⁻¹) and symmetric (983 cm⁻¹) stretching, –FeOSOFe vibration (955 cm⁻¹), as well as –FeO stretching (830 cm⁻¹) characteristic of the Fe-based mordant green vitriol (FeSO₄·xH₂O) (Table 2, Supplementary Table 4), which was likely used in this wallpaper. Several brown drawings were part of INV_11 and INV_13 (Table 1), where Fe detection (Fig. 6f, Supplementary Fig. 5c) might infer to the utilization of Taisha, also supported by the low content of Si, Mn, and Zn, which confer diverse yellow–brown and red-brown shades to the pigment [19]. Additionally, lead white and/or lead oxide could be present in INV_11_4 and INV_13_5, as indicated by Pb traces, while INV_13_6...
and INV_13_9 showed a relatively high distribution of both As and S (Fig. 6f) attributable to the addition of Sekio or Yuo to get unique brown shades\textsuperscript{27,28}. Finally, the black colour was only present in INV_11_1–10, whose elemental analysis revealed traces of S and the three ubiquitous elements Ca, Fe, and Sn (Fig. 6g), implying the use of an organic black pigment for this colour\textsuperscript{27,28}.

**Organic pigments.** Several IR absorption bands typical of the natural purple-pigment Indigo were detected in the green areas of INV_11_3 and INV_13_8 (Fig. 2a, b, Table 2, Supplementary Table 1–2), whose similarity was confirmed by the partitioning of their spectra in cluster 4 (average $s_i = 0.56$; first interval) or 2 (average $s_i = 0.70$; second interval), as well as their PC2 values (Figs. 3c–e, 4c–e). This observation agrees with the practice of Japanese artists, from the late Edo period (1840–1860) onwards, to mix Indigo with lime, ashes, and Sekio as an alternative to malachite green\textsuperscript{24,25,26}. On the other hand, the INV_20_3 spectrum, belonging to cluster 1 for both the first (average $s_i = 0.83$) and second (average $s_i = 0.76$) intervals (Figs. 3c–e, 4c–e), suggested that diverse techniques could have been used to obtain the green colour in INV_20 as compared to INV_11 and INV_13. Although INV_20_3 featured some Indigo's IR contributions, the detection of vibrational modes typical of -SO$_4$ and -FeO groups (Supplementary Table 4) indicated that iron sulphate mordants, such as green vitriol, were supplied to Indigo and Sekio obtaining the green colour. Indeed, a similar procedure was developed by English artisans, who added pearl ashes, lime, Sekio, and green vitriol to Indigo syrup, favouring both the precipitation of Fe protoxide by the lime, as well as the deoxydgenation of Indigo, which acquired a dark green shade\textsuperscript{27,28}. Besides, only a few of Indigo’s IR contributions were identified in red and brown drawings (Table 1, Supplementary Tables 1–2, 4), reasonably due to (1) Indigo diffusion from adjacent areas containing the organic pigment or (2) its utilisation to obtain diverse colour shades. The μATR-FTIR spectra collected for brown and red drawings of INV_11 and INV_13 (Table 1) showed a high degree of similarity, as they all clustered with INV_11_3 and INV_13_8 spectra for both the 3770–2750 cm\textsuperscript{-1} and 1840–719 cm\textsuperscript{-1} intervals (Figs. 3c–e, 4c–e). INV_13_9 constituted the only exception, as it was grouped in cluster 1 (Figs. 3c, 4c), underlining that diverse organic substances, alongside the inorganic Sekio (Supplementary Fig. 5c), may be used for this brown sampling point. Since Indigo features low IR contributions in the 3770–2750 cm\textsuperscript{-1} interval\textsuperscript{29}, indications regarding its presence within red and brown drawings can be better obtained by focusing on the MBD-based analysis of the second IR interval, where INV_13_5.1 was the deepest curve (Fig. 4f), while INV_11_6, INV_15_4, and INV_20_4 were the most external and variable of the cluster (Fig. 4d, Supplementary Fig. 3b). Hence, it is reasonable to suggest that Indigo diffusion was responsible for IR contributions in INV_11_6 and INV_20_4, while it was likely used for INV_11_4, INV_13_3, and INV_13_5, being part of the core cluster (Supplementary Fig. 3b). Moreover, INV_15_4 partitioning within cluster 2 for the 1840–719 cm\textsuperscript{-1} interval (Fig. 4c–e) may be due to the presence of other organic substances having absorption bands in common with Indigo. Indeed, any of typical IR contributions of this organic pigment were observed in INV_15_4 μATR-FTIR spectrum (Supplementary Table 3), which also featured a low depth value, as well as a high PC2 score, with respect to the other spectra clustering together (Fig. 4d, Supplementary Fig. 5b).

**Organic binders.** The identification of inorganic pigments highlighted the need for Japanese artisans to use binders (e.g., proteinaceous binders, vegetable oils, waxes, and lacquers) for their painting\textsuperscript{15}. Although the type of these substances is generally determined through FTIR, the wallpapers’ complexity impaired their univocal identification, since they had similar IR absorption bands and their specific vibrational modes often overlapped with each other (Supplementary Tables 1–4)\textsuperscript{30,31}. In this regard, the clustering analysis in the 1840–719 cm\textsuperscript{-1} interval revealed that spectra featuring Indigo’s IR contributions grouped in cluster 2, while cluster 1 featured the verso spectra and those where the organic pigment was absent or less present, in which overlapping vibrational modes of cellulose, lignin, silicates, carbonates, and organic binders were ubiquitous (Fig. 4c, Table 2). Similarly, although four clusters were identified for the μATR-FTIR spectra in the 3770–2750 cm\textsuperscript{-1} interval (Fig. 3c–e), several absorption bands attributable to these substances were detected (Supplementary Tables 1–4). Nevertheless, the type of organic substances was presumed, when possible, by comparing their IR absorption bands with those characterising the organic binders (Supplementary Tables 1–4). Thus, vibrational modes typical of proteinaceous binders (INV_15) or waxes (INV_13_9 and INV_20) were detected (Supplementary Tables 3–4), which is in agreement with the Japanese traditional use of animal glue (nikawa), gelatine cubes, and beeswax as binders\textsuperscript{13,25}. Since IR absorption bands distinctive of proteinaceous binders or waxes are more present in the 3770–2750 cm\textsuperscript{-1} interval\textsuperscript{30,31}, the partitioning of μATR-FTIR spectra of INV_15 recto in cluster 3 (average $s_i = 0.66$), INV_13_9 and INV_20_3 in cluster 1 (Fig. 3c, d) statistically supported these observations. Particularly, the MBD-based analysis of cluster 1 revealed a high degree of similarity between these μATR-FTIR spectra, representing both the deepest curve (INV_13_9) and the core (70%) of the cluster itself, while INV_11_4.1 was the most external spectrum (Fig. 3f, Supplementary Fig. 2a), where IR absorption bands typical of waxes were not detected (Supplementary Table 1). Moreover, although vibrational modes characterising waxes are less present in the 1840–719 cm\textsuperscript{-1} interval\textsuperscript{30}, the comparable PC2 values of INV_13_9, INV_13_9.1, INV_13_9.2, and INV_20_3 spectra highlighted their higher variability, yet intrinsic similarity, than other curves belonging to cluster 1 (Fig. 4d).

Several μATR-FTIR spectra displayed also vibrational modes distinctive of organic vegetable lacquers (Table 2, Supplementary Tables 1–4), which grouped in cluster 3 in the 3770–2750 cm\textsuperscript{-1} interval (Fig. 3c–e). In this regard, the so-called lacquering procedure for the wallpapers’ production involved the mixing of plant-extracted lacquers—likely the Urushi one, based on the Japanese tradition\textsuperscript{27,28}, with Odo, Taitshi, and Kincha to confer the yellow-gold background characteristic of the Kinkakurokakami-gami (Fig. 3g). Indeed, the Urushi lacquer features a strong IR contribution around 3400 cm\textsuperscript{-1} attributable to –OH stretching\textsuperscript{32}, which, alongside the vibrational modes of proteinaceous binders, was a distinctive trait of the μATR-FTIR spectra belonging to cluster 3 (first
whose production and secretion of xylanases and laccases is one of their distinctive metabolic traits. For- respectively (Table 3), and autochthonous of Cladosporium cf. ramotenellum and M. chokoriensis were observed (Table 3). These results are in line with the studies reporting the isolation of cultivable microorganisms from wall or easel paintings. The microbial isolates C4(11)3, C4(13)1, C4(11)1, and C4(11)2 closely related to B. coreaensis, B. pocheonensis, M. chokoriensis, and Cladosporium cf. ramotenellum respectively (Table 3), and autochthonous of Kinkarakawa-gami, are commonly found in several environmental matrices of Korea, China, Thailand, and Japan, confirming the eastern provenance of these wallpapers. Besides, these microorganisms hold enzymatic assets (i.e., xylanases and laccases) that enable them to degrade and depolymerize lignin and cellulose compounds, lacquers, and organic pigments. Indeed, Bacillus, Staphylococcus, and Micrococcus spp. produce extracellular xylanases, which efficiently hydrolyse lignocellulosic material and the Indigo pigment, a biotic aspect that supports their presence on INV_11, INV_13, and INV_15, while any cultivable microorganisms was not retrieved from INV_20. Bacterial isolates were identified as part of the Firmicutes (i.e., Bacillus coreaensis, B. pocheonensis, Staphylococcus capitis, and S. epidermidis) and Actinobacteria (i.e., Kocuria rizophila (former Micrococcus luteus) and Micromonaspora chokoriensis) phyla, while all the detected fungi belonged to the Ascomycetes phylum, although four genera were observed (Table 3). These results are in line with the studies reporting the isolation of cultivable microorganisms from wall or easel paintings.

The microbial isolates C4(11)3, C4(13)1, C4(11)1, and C4(11)2 are proficient in degrading gelatine and starch, which were two of the most used binders in Japanese tradition. Since these enzymes are responsible for the degradation of phenol substrates, the partial identification of vibrational modes typical of urushiol polymerization or Indigo (Supplementary Tables 1–4) could be ascribed to the biotic deterioration of both Urushi lacquer and organic pigment, as Japanese lacquer IR contributions were found within sampling points where fungal strains were isolated (Tables 2, 3). A similar conclusion can be made for C4(11)3 and C4(11)2 isolates phylogenetically related to B. coreaensis and C. ramotenellum respectively, whose production and secretion of xylanases and laccases is one of their distinctive metabolic traits. Further, Bacillus [C4(13)2], Staphylococcus [C4(13)2 and C4(15)1], and Micrococcus [C4(11)1 and C4(13)1] species are among the most persistent strains capable of metabolizing honey and beeswax, while M. chokoriensis [C4(11)1] and C. ramotenellum [C4(11)2] are proficient in degrading gelatine and starch, which were two of the most used binders in Japanese tradition.
The presence of microorganisms can also be linked to their tolerance and/or resistance towards a broad spectrum of metal and metalloid compounds\(^{35}\), which are the main components of inorganic pigments. For instance, bacteria belonging to the Firmicutes and Actinobacteria phyla can detoxify their surrounding environment from toxic metal or metalloids, even using them as a terminal electron acceptor to produce energy\(^{32,33}\). Bacillus, Staphylococcus, and Micrococcus spp. are overall present in paintings where carbonates (chalk—CaCO\(_3\)) and silicates (quartz—SiO\(_2\)) are abundant\(^{46,48}\), due to their ability to overcome the challenge deriving from these minerals. Bacillus spp. are also highly tolerant towards Pb\(^{36}\) and Fe-containing compounds\(^{37}\), being able to transform lead acetate [Pb(CH\(_3\)CO\(_2\))]\(_2\)\(^{35}\) and hematite (Fe\(_2\)O\(_3\)) and its hydrated forms\(^{38}\), as well as to oxidize Fe(II) to Fe(III), as in the case of B. pocheonensis\(^{39}\). Analogously, S. epidermidis, M. chokoriensis, and K. rizophila strains were studied for their resistance against Pb-containing compounds\(^{40,42}\), while the fungal strain Hirustella spp. showed tolerance to Ca- and Fe-sulphates\(^{43}\). Thus, the elemental composition of INV_11 and INV_13 (Fig. 6) justified the presence of microorganisms (Table 3) that hold metabolic traits allowing them to survive on metal-rich wallpapers. Although INV_15 and INV_16 displayed a high amount of Shu (HgS) (Supplementary Fig. 5a), one bacterial [C\(_6\)(151)] and two fungal [C\(_6\)(152) and C\(_6\)(153)] strains related to S. epidermidis, Penicillium georgiense, and Aspergillus costaricaensis were isolated (Table 3), highlighting their great tolerance towards Hg-containing compounds\(^{46,48}\). Indeed, these fungal strains were responsible, among others, for the darkening of cinnabar (Shu), as reported elsewhere\(^{46}\). The isolation of S. epidermidis and Hirustella strains could also derive from anthropogenic/environmental contamination, as they are human and nematode pathogens respectively\(^{46,64}\), reflecting the artefacts’ poor state of conservation.

Besides, the local heterotrophic microflora (i.e., Bacillus, Micrococcus, and Micrococcus species) can produce secondary metabolites with antimicrobial properties as a defence mechanism under stress conditions\(^{34,46,47}\). Thus, Firmicutes and Actinobacteria could act as biocontrol agents for the preservation of cultural heritage\(^{34}\). These observations, along with the advanced state of deterioration visible for INV_20, may indicate the key role played by bacteria belonging to Bacillus, Micrococcus, and Micrococcus genera in controlling and preventing the further degradation of the colonized wallpapers; indeed, INV_20 was the only artefact from which any cultivable microorganism was not isolated.

Conclusion

This multidisciplinary study allowed to unveil physical–chemical features regarding the composition, manufacture, and dating of the collectively important cultural heritage represented by Kinkarakawa-gami works of art. The experimental evidence gathered was corroborated through the uncommon yet resourceful and innovative in blind functional data statistical analysis. Indeed, given the complexity of the studied wallpapers in terms of IR vibrational modes ascribed to (1) substances used for their fabrication, (2) physical–chemical degradation products, and (3) the presence of a microbiota, this statistical approach has proved greatly helpful to identify and confirm trends, differences, and similarities observed among the four Kinkarakawa-gami artefacts. Microbiological investigations supported the Eastern provenance of these wallpapers, however, whether the cultivable microbes act as detentiogen or biocontrol agents is still under debate; hence, DNA sequencing-based technology (e.g., study of the microbiome) represents the new frontier to unveil the identity of uncultivable microorganisms, and, alongside physical–chemical characterisation, will improve the development of innovative, promising, and eco-friendly restoration strategies for the conservation of cultural heritage.

Experimental section

Materials. The four wallpapers here studied belong to the V. Raga-O’T. Kiyohara collection of Palermo (Italy). Given the complexity and richness of these artefacts in terms of details and depicted colours, an extensive sampling of the wallpapers (Table 1) was performed to thoroughly analyse them.

Tryptic soy, malt extract, and agar technical were purchased from Sigma-Aldrich® (Milan, Italy), while the experimental section

X-Ray fluorescence (XRF) spectroscopy. A Tracer III sd Bruker AXS (Bruker, UK) equipped with Rhode anode and working at 40 kV and 11 μA was exploited for XRF analyses, whose acquisition time was 30 s. Element identification and XRF spectra analysis was performed by using ARTAX\(^*\) software, which was provided with the instrument, while R 3.6.1 and OriginPro\(^*\) 2016 software were used for the representation. Elemental compositions of wallpaper sampling points are reported in terms of Net Area (×10\(^4\) a.u.), which represents the integral intensity of X-ray emission characteristic of each element obtained after performing the Bayes deconvolution and deduction of the background intensity\(^{29}\). These results are to be considered as a qualitative estimation regarding the presence and abundance of diverse elements within the chosen sampling points, as any appropriate standard was not used to determine the concentration of each element.

Attenuated total reflectance-Fourier transform infrared (ATR-FTIR) spectroscopy. ATR-FTIR spectra were recorder by using a µFTIR Lumos (Bruker, UK) equipped with a Platinum ATR and an IR microscope featuring 0.1 μm as lateral resolution. The spectra were collected in the 4000–600 cm\(^{-1}\) range, with a resolution of 2 cm\(^{-1}\) and 60 scans per each sampling point, and they were subsequently analysed through the software OPUS(7.5)*, which was provided with the instrument, as well as OriginPro\(^*\) 2016.

Statistical analyses of µATR-FTIR spectra. Based on the structure of µATR-FTIR spectra, they can be assimilated to complex sets of data (i.e., functions, where each spectrum corresponds to a distinct function) varying over a continuum (i.e., wavenumber range) that, taken together, can be considered as single curves. Hence, in blind functional data analysis (FDA) was applied as the statistical methodology on these spectra\(^{11}\).
that were split, based on the wavenumber range acquisition, in 3770–2750 cm\(^{-1}\) and 1840–719 cm\(^{-1}\) intervals, which were singularly analysed by performing a hierarchical clustering of \(\mu\)ATR-FTIR curves with the respect to their shape. The quantification of each curve's cohesion to its own cluster as compared to the separation from the other clusters was obtained through a silhouette (s) measurement, while the optimal number of clusters was determined by maximizing the average s, to the configuration obtained in the hierarchical clustering.

Functional Principal Component Analysis (FPCA) was then performed to derive the PCs inside the final clusters\(^3\) and find an optimal orthogonal linear projection of the curves on a d-dimensional subspace (R\(^d\) when d = 2 or d = 3) to minimize the expected value of the squared error due to the projection. Lastly, a data depth algorithm based on the Modified Band Depth (MBD) was constructed focusing on centrality and separation of the spectra\(^2\), allowing to identify both the most representative and the most external curves of each cluster. All the statistical analyses were performed by using the R 3.6.1 software; a more extensive review of the performed statistical analyses and the R-packages used is reported in the Supplementary Methods section.

**Fluorescence microscopy.** Fragments of the wallpapers were imaged both on recto and verso sides by a Leica TCS SP5 fluorescence confocal laser scanning microscope (CLSM), using a 40X-1.25 NA objective (Leica Microsystems, Germany). Images were acquired under two-photon excitation at 830 nm in the 450–700 nm emission range. The samples were soaked in glycerol during the imaging procedure. The same setup was applied for a 3D reconstruction of the wallpaper multilayers. The data were analysed by ImageJ software.

**Microbiological analyses.** The sample areas of interest of Kinkarakawa-gami were gently swiped with sterile cotton swabs, which were then suspended in the physiological solution (sodium chloride 0.9% w/v) for 30 min. Afterward, the suspensions were serially diluted, being aliquots (100 \(\mu\)L) spread onto both tryptic soy and malt extract agar plates to isolate either bacterial or fungal strains respectively, whose biomass growth was carried out at 30 °C for 5 days under static conditions.

Polymerase chain reaction (PCR) was performed (Supplementary Methods), following thermocycler conditions described elsewhere\(^3\), on the extracted and purified genomic DNA to obtain the 16S rRNA gene product and the internal transcribed spacer (ITS) region in the ribosomal RNA operon. To identify—at genus level—the isolates retrieved from Kinkarakawa-gami’s art, PCR products were purified through QIAquick PCR purification kit, according to the manufacturer’s protocol, sequenced (BMR Genomics, Padova, Italy), and searched for nucleotide homology with other microorganisms (Supplementary Methods).

Received: 11 May 2020; Accepted: 7 September 2020
Published online: 02 October 2020

**References**

1. Wailliez, W. Japanese leather paper or Kinkarakawagami: an overview from the 17th century to the Japonist hangings by Rottmann & Co. *Wallpap. Hist. Rev.* 7, 60–65 (2015).
2. Suga, Y., Chapter, “Artistic and commercial” Japan: modernity, authenticity and Japanese leather paper. In *Buying for the Home: Shopping for the Domestic from the Seventeenth Century to the Present* (eds Hussey, D. & Ponsonby, M.) (Ashgate Publishing, Farnham, 2008).
3. Alcock, R. Lacquer ware, wall-papers, textile fabrics, and embroidery. In *Art and Art Industries in Japan* (ed. Alcock, R.) 219–236 (Virtue & Co, London, 1878).
4. Dresser, C. Chapter VII: Minor manufactures of Japan. In *Japan, Its Architecture, Art, and Art Manufactures* (ed. Dresser, C.) 450–466 (Longmans, Green, & Co, London, 1882).
5. Guimet, E., Regamey, F. *Promenades Japonaises: Tokio-Nikko* (eds. Guimet, E. & Regamey, F.) (G. Charpentier Editeur, Paris, 1880).
6. Huberman, T., Ashmore, S., Suga, Y. *The diary of Charles Holmes’s 1889 visit to Japan and North America, with Mrs Lasenby Liberty’s Japan: A pictorial record* (eds. Huberman, T., Ashmore, S. & Suga, Y.) (Global Oriental, 2008).
7. Unknown authors. Japanese leather papers. *The Decorator and Furnisher*. 6, 114 (1885).
8. Saladino, M. L. et al. A multi-analytical non-invasive and micro-invasive approach to canvas oil paintings. General considerations from a specific case. *Microchem.* J. 133, 607–613 (2017).
9. Caruso, F. et al. Micro-analytical identification of the components of varnishes from South Italian historical musical instruments by PLM, ESEM-EDX, microFTIR, GC-MS, and Py-GC-MS. *Microchem. J.* 116, 31–40 (2014).
10. Hoglund, H. Pulping technology. In *Pulping Chemistry and Technology* (eds Ek, M. et al.) 57–90 (Walter de Gruyter, Berlin, 2009).
11. Hubbe, M. A. & Gill, R. A. Fillers for papermaking: a review of their properties, usage practices, and their mechanistic role. *BioResources* 11, 2886–2963 (2016).
12. Hahn, O., Malter, W., Kanngiesser, B. & Beckhoff, B. Characterization of iron-gall inks in historical manuscripts and music compositions using x-ray fluorescence spectrometry. *X-Ray Spectrom.* 33, 234–239 (2004).
13. Vahur, S. et al. ATR_FT-IR spectral collection of conservation materials in the extended region of 4000–80 cm\(^{-1}\). *Anal. Bioanal. Chem.* 408, 3373–3379 (2016).
14. Wright, D. K. Accuracy vs precision: understanding potential errors from radiocarbon dating on African landscapes. *Afr. Archaeol. Rev.* 34, 303–319 (2017).
15. Nishio, Y. Pigments used in Japanese paintings. *Pap. Conserv.* 11, 39–45 (1987).
16. Moioli, P. & Seccaroni, C. Analysis of art objects using a portable X-Ray Fluorescence spectrometer. *X-Ray Spectrom.* 29, 48–52 (2000).
17. Eastagh, N., Walsh, V., Chaplin, T., Siddall, R. *Pigment Compendium: A Dictionary and Optical Microscopy of Historical Pigments* (eds. Eastagh N., Walsh V., Chaplin T. & Siddall R.) (Butterworth-Heinemann Springer, 2008).
18. Dalen, M. B. & Mamza, P. A. P. Some physico-chemical properties of prepared metallic soap-driers of aluminum, copper and zinc. *Sci. World J.* 4, 7–9 (2009).
19. Auer, G., et al. Pigments, inorganic, 2. White pigments. *Ulmann’s Encyclopedia Industr. Chem.* 1–36 (2017).
20. Ostmond, G. Zinc White and the influence of paint composition for stability in oil based media. In *Issues in Contemporary Oil Paint* (eds van den Berg, K. J. et al.) 263–281 (Springer, Berlin, 2014).
21. Borgia, I. et al. The combined used of lead-tin yellow type I and II on a canvas painting by Pietro Perugino. *J. Cult. Herit.* 8, 65–68 (2007).
22. Yoshimichi, E. Coloring matter used in Japanese painting (particularly those applied to architecture). In International Symposium on the Conservation and Restoration of Cultural Property (eds. Suzuki, T. & Mausa K.) 157–165 (Tokyo National Research Institute of Cultural Properties, 1985).

23. Vermeulen, M. et al. Characterization of Japanese color sticks by energy dispersive X-ray fluorescence, X-ray diffraction and Fourier transform infrared analysis. Spectrochim. Acta B 65, 321–327 (2010).

24. Zaleski, S., Takahashi, Y. & Leona, M. Natural and synthetic arsenic sulfide pigments in Japanese woodblock prints of the late Edo period. Herit. Sci. 6, 32 (2018).

25. Manso, M. et al. Characterization of Japanese color sticks by energy dispersive X-ray fluorescence, X-ray diffraction and Fourier transform infrared analysis. Spectrochim. Acta B 65, 321–327 (2010).

26. Zaleski, S., Takahashi, Y. & Leona, M. Natural and synthetic arsenic sulfide pigments in Japanese woodblock prints of the late Edo period. Herit. Sci. 6, 32 (2018).

27. Croker, T. H., Williams, T., Clark, S. Indigo. In The Dictionary of Arts and Sciences in which the Whole Circle of Human Learning is Explained (eds. Croker T. H., Williams T. & Clark S.) 74–75 (J. Wilson & J. Fell, 1765).

28. Cooper, T. Indigo. In A Practical Treatise on Dyeing and Calico Printing: Exhibiting the in the French, German, and English, and American Practice of Fixing Colours on Woolen, Cotton, Silk, and Linen (ed. Cooper T.) 45–82 (Thomas Dobson, William Fry Printer, 1815).

29. Baran, A., Fiedler, A., Schulz, H. & Baranska, M. In situ Raman and IR spectroscopic analysis of indigo dye. Anal. Methods 2, 1372–1376 (2010).

30. Meilunas, R. J., Bentsen, J. G. & Steiberg, A. Analysis of aged paint binders dye FTIR spectroscopy. Stud. Conserv. 35, 33–51 (1990).

31. Tanner, N. & Lichtenberg-Kragg, B. Identification and quantification of single and multi-adulteration of beeswax by FTIR-ATR spectroscopy. Eur. J. Lipid Sci. Technol. 121, 1900245 (2019).

32. Niimura, N. & Miyakoshi, T. Structural study of oriental lacquer films during the hardening process. Talanta 70, 146–152 (2006).

33. Naumann, D., Helm, D. & Labischinski, H. Microbial characterizations by FT-IR spectroscopy. Talanta 49, 711–745 (1995).

34. Soffritti, I. The potential use of microorganisms as restorative agents: an update. Sustainability 11, 3853 (2019).

35. Costerton, J. W., Lewandowski, Z., Caldwell, D. E., Korber, D. R. & Lappinscott, H. M. Microbial biofilms. Ann. Rev. Microbiol. 49, 399–403 (1996).

36. Ratakanhabokhai, K., Kyu, K. L. & Tanticharoen, M. Purification and properties of a xylan-binding exoαxylanase from alkalophilic Bacillus sp. strain K-1. Appl. Environ. Microbiol. 65, 694–697 (1999).

37. Ten, L. N. et al. Bacillus pocheonensis sp. nov., a moderately halotolerant, aerobic bacterium isolated from soil of a ginseng field. Int. J. Syst. Evol. Microbiol. 57, 2532–2537 (2007).

38. Tiwari, K. & Gupta, R. K. Diversity and isolation of rare actinomycetes: an overview. J. Microbiol. 140, 19–26 (1994).

39. Ayla, S., Golla, N. & Pallipati, S. Production of ligninolytic enzymes from Penicillium sp. and its efficiency to decolourise textile dyes. Open Biotechnol. J. 12, 112–122 (2018).

40. Ali, J. et al. Characterization and dye decolorization ability of an alkaline resistant and organic solvents tolerant laccase from Bacillus licheniformis LS04. Biore. Technol. 115, 35–40 (2012).

41. Lu, L. et al. Characterization and dye decolorization ability of an alkaline resistant and organic solvents tolerant laccase from Bacillus licheniformis LS04. Biore. Technol. 115, 35–40 (2012).

42. Genesse, A. & Mamo, G. Purification and characterization of an alkaline xylanase from alkalophilic Micrococcus sp. AR-135. J. Ind. Microbiol. Biotechnol. 20, 210–214 (1998).

43. Yoshimichi, E. Coloring matter used in Japanese painting (particularly those applied to architecture). In International Symposium on the Conservation and Restoration of Cultural Property (eds. Suzuki, T. & Mausa K.) 157–165 (Tokyo National Research Institute of Cultural Properties, 1985).

44. Geeser, A. & Mamo, G. Purification and characterization of an alkaline xylanase from alkalophilic Micrococcus sp. AR-135. J. Ind. Microbiol. Biotechnol. 20, 210–214 (1998).

45. Yoshimichi, E. Coloring matter used in Japanese painting (particularly those applied to architecture). In International Symposium on the Conservation and Restoration of Cultural Property (eds. Suzuki, T. & Mausa K.) 157–165 (Tokyo National Research Institute of Cultural Properties, 1985).

46. Bao, S. et al. Characterization and dye decolorization ability of an alkaline resistant and organic solvents tolerant laccase from Bacillus licheniformis LS04. Biore. Technol. 115, 35–40 (2012).

47. Petrushkova, J. P. & Lylakova, N. N. Microbiological degradation of lead-containing pigments in mural paintings. Stud. Conserv. 31, 65–69 (1986).

48. Lu, S. et al. Ecophysiology of Fe-cycling bacteria in acidic sediments. Appl. Environ. Microbiol. 76, 8174–8183 (2010).

49. Croker, T. H., Williams, T., Clark, S. Indigo. In The Dictionary of Arts and Sciences in which the Whole Circle of Human Learning is Explained (eds. Croker T. H., Williams T. & Clark S.) 74–75 (J. Wilson & J. Fell, 1765).

50. Selvin, J., Shanmugha Priya, S., Seghal Kiran, G., Thangavelu, T. & Sapna Bai, N. Sponge-associated marine bacteria as indicators of heavy metal pollution. Microbiol. Res. 164, 352–363 (2009).

51. Shen, S., Hussain, S. Z. & Rehman, A. Phenotypic and genomic analysis of multiple heavy metal-resistant Micrococcus luteus strain ASI isolated from industrial wastewater and its potential use in arsenic bioremediation. Appl. Microbiol. Biotechnol. 104, 2243–2254 (2020).

52. Lin, S. et al. Enhancement of cordyceps polysaccharide production via biosynthetic pathway analysis in Hirutella sinensis. Int. J. Biol. Macromol. 92, 872–880 (2016).

53. Palacienca, E., Presentato, A. & Turner, R. J. Stability of biogenic metalloid nanomaterials related to the colloidal stabilization theory of chemical nanostructures. Crit. Rev. Biotechnol. 38, 1137–1156 (2018).

54. Presentato, A. et al. Assembly, growth and conductive properties of tellurium nanorods produced by Rhodococcus aerophilus BCP1. Sci. Rep. 8, 3923 (2018).

55. Eger, J. et al. Utilizing the iron tolerance potential of Bacillus species for biogenic synthesis of magnetite with visible light active catalytic activity. Colloid Surf. B 177, 470–478 (2019).

56. Li, C., Chen, Y., Wang, C. & Kuang, Z. Structural study of oriental lacquer films during the hardening process. Talanta 70, 146–152 (2006).

57. Li, C., Chen, Y., Wang, C. & Kuang, Z. Structural study of oriental lacquer films during the hardening process. Talanta 70, 146–152 (2006).

58. Li, C., Chen, Y., Wang, C. & Kuang, Z. Structural study of oriental lacquer films during the hardening process. Talanta 70, 146–152 (2006).

59. Li, C., Chen, Y., Wang, C. & Kuang, Z. Structural study of oriental lacquer films during the hardening process. Talanta 70, 146–152 (2006).

60. Li, C., Chen, Y., Wang, C. & Kuang, Z. Structural study of oriental lacquer films during the hardening process. Talanta 70, 146–152 (2006).
Author contributions
E.P. designed and performed XRF and μATR-FTIR experiments, interpreted the data and drafted the entire manuscript. A.P. performed and interpreted the experiments regarding the isolation and identification of microorganisms and was the second contributor to the manuscript drafting. F.D.S. carried out the statistical analysis of μATR-FTIR spectra, while R.A. supplied a contribution in the microbiological analyses and the editing of the manuscript. V.F. and G.S. performed and interpreted the confocal fluorescence imaging, whereas V.M. contributed to the interpretation of the μATR-FTIR spectra. A.G. chose the artefacts and provided historical and artistic information. D.F.C.M. provided a major intellectual and financial contribution during the development of the study, managing and directing the research, as well as revising the manuscript.

Funding
This work is part of the PON project “Development and Application of Innovative Materials and Processes for the Diagnosis and Restoration of Cultural Heritage -DELIAS” (PON03PE_00214 2-DELIAS), which was funded by the MIUR.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41598-020-73226-6.

Correspondence and requests for materials should be addressed to A.P. or D.F.C.M.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2020