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Introduction

The use of analytical techniques for the study of objects of historical and artistic interest has increased in the last thirty vears, providing useful information about artists' techniques, the provenance of materials, the nature of degradation processes,1 authentication and dating. The priceless nature of works of art has driven scientists to employ non-invasive and non-destructive techniques. This, coupled with the high insurance values of the articles, essentially prevents the analysis of objects in host laboratories: work has to be done within the conservation studios of the host institution. Over the past decade, this has been achieved with the advent of portable, instrumentation that can be taken to the host libraries or institutions for *in situ* investigations on the fly. Among the different techniques that are routinely employed for the analysis of artefacts or manuscripts, micro-Raman spectroscopy has proven to be an extremely powerful, due to its portability, specificity,

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Raman spectroscopic library of medieval pigments collected with five different wavelengths for investigation of illuminated manuscripts[†]

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Raman spectroscopy is widely applied in the cultural heritage field to perform non-destructive measurements *in situ*, in order to identify materials, specifically pigments. The spectra collected can be challenging to interpret because certain source laser wavelengths may be absorbed by specific pigments, leading to large fluorescence backgrounds which obscure the weak Raman signals, or worse cause photodegradation of the sample. Furthermore, the reference spectra for a specific pigment obtained from a particular laser wavelength is not always available and is a crucial step in the detective work of pigment identification, especially when the resonance Raman effect can enhance some signals. As the range of lasers available increases, spectral libraries do not always record spectra acquired with the same wavelength used to carry out the measurements in field. In this work, reference spectra of 32 different wavelengths were used as excitation sources. The aim is to provide a useful and more complete reference source to enable better planning of which laser wavelength is the most appropriate to study a specific set of pigments, and to allow comparisons between spectra acquired with the same wavelength, leading to the unequivocal pigment identification in a step by step manner.

spatial resolution and non-contact, non-destructive nature.²⁻³¹ The first effective application of Raman spectroscopy on cultural heritage objects was on illuminated manuscripts to identify pigments,32 and it has since been used to study a wide variety of materials including paper,33 binder media,34 inks,35,36 glass,5,9,10,18 ceramics and pottery,7,8 gemstones,20 stones and rocks from archaeological sites,^{19,30} degradation products.³ The availability of a spectral library is therefore essential to help identify the materials deployed.^{34,37-39} A few portable Raman spectrometers are now commercially available, but these should be used with extreme caution: in many examples, the power of the laser light source is much higher than we would advocate and may cause damage to the artefact. Furthermore, commercially available portable instruments usually have a single laser source, in rare cases two, which are not necessarily the best ones to investigate the wide variety of pigments that can be found on the same artefact, positive identifications can often be difficult.26,31 Indeed, pigments respond differently to laser irradiation according to their nature (dyes or pigments, colour, etc.). In certain cases, for example, where the laser wavelength matches or is close to the absorption bands of the pigment being analysed, the Resonance Raman effect can lead to excellent sensitivity, but, in some cases, some absorption can also lead to large interfering, luminescent background signals which hide the low intensity Raman signals. The selection of the most appropriate wavelength for the identification of pigments then is of great interest and importance.



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[†] Electronic supplementary information (ESI) available: Figures of pigments' spectra, from 1 to 30; the data of the Raman spectra collected at the different wavelengths. See DOI: 10.1039/c8ay00016f

Analytical Methods

Portable equipment means compromises have to be made in comparison with fixed laboratory equipment. Portable systems tend to sacrifice spectral resolution and sensitivity.³¹ This work provides an updated library of pigments' Raman spectra acquired using different laser wavelengths in order to supply the best spectrum possible for each pigment for comparison to the data collected *in situ*. The pigments have been chosen as representative of those used in illuminated manuscripts between Vth–XVIth centuries in Europe.^{32,36,40–67}

Experimental

Instrumentation

Two different Raman spectrometers have been used, equipped with different lasers. The first was a Horiba Jobin Yvon LabRAM HR confocal Raman microscope, equipped with a Peltier-cooled CCD and $50 \times$ LWD Leica objective. The instrument has four different laser sources available, 488 nm, 532 nm, 632.8 nm and 785 nm providing a diffraction limited laser spot from 1 to 2 μ m diameter. Each laser source has different maximum power values that can be reduced using neutral density filter (100% 50%, 25%, 10%, 1%, 0.1% and 0.01%). Reported laser powers were measured after the objective lens in the sample plane. A 600 l mm⁻¹ grating was used for measurements using the 532 nm, 632.8 nm and 785 nm laser, and an 1800 l mm⁻¹ grating used for the 488 nm laser. The minimum wavenumber for each laser wavelength, dictated by the edge filters in the spectrometer were 200 cm⁻¹, 120 cm⁻¹, 70 cm⁻¹, 100 cm⁻¹ respectively for the 488 nm, 532 nm, 632.8 nm and 785 nm lasers. All the acquisition operations were controlled by Lab Spec 6-Horiba Scientific software.

For excitation at 830 nm, a Renishaw InVIa micro-Raman spectrometer (Rutherford Appleton Laboratory, Harwell Campus, Didcot), with 830 nm laser source, a silicon CCD detector and Nikon L-PLAN SLWD $50 \times / 0.45$. The resulting laser spot was circa 2 μ m in diameter. The spectral range recorded was 70–1800 cm⁻¹, with a 1200 l mm⁻¹ grating. The maximum laser power was 55 mW, and this could be reduced to lower levels (50%, 25%, 10%, 5%, 1%, 0.5%, 0.1% and 0.05%). Again, the laser power was recorded after the objective in the sample plane.

A silicon standard sample was used as reference for calibration (520 cm^{-1}). The time of acquisitions and number of accumulations was decided on a sample-by-sample basis.

To be able to compare spectra acquired with different devices, spectra were corrected for the instrumental response by comparison to the spectra obtained for a broad-band source.^{68,69} A stable white light source (HL-2000-CAL Ocean Optics), whose spectral distribution was known (and can be approximated by a black body radiator of 2939 K), was used to generate a correction curve. The lamp emission spectrum was provided as photons/($\Delta\lambda t$), where $\Delta\lambda$ is the bandwidth detected by the spectrometer, and *t* the time unit. It had to be multiplied by λ^2 (where λ is the emission wavelength), in order to obtain the spectrum in terms of photons/ $t \times \Delta \bar{\nu}$ of Raman shift. The calibration lamp spectrum was recorded for every wavelength laser using the same scan-conditions as used for the sample

measurements and covering the same wavenumber range. Using this correction curve, all of the Raman spectra were corrected for instrument response. No background subtraction was performed, since one of the goals of this work was to provide a library that helps to decide which the best wavelength to investigate a certain pigment is, so any luminescent background that may detract from the signal or prove diagnostic was recorded. All the spectra have been normalized to a maximum intensity of 1 in the graphs (Fig. 1–30) available in the ESI (ESI†). In the ESI,† the raw ASCII data of the spectra can be found. Corrected for the instrument response. Thus, relative intensities of peaks may be obtained and compared directly.

Classification of the spectra

A classification of the quality of the spectra collected (Table 1) was made, in order to establish the optimum wavelength for investigating the presence of a specific pigment, and which wavelength provides the maximum number of positive identifications. To classify them, a criterion to value the quality of the spectrum had to be established and the signal to noise ratio was considered appropriate. Even though this parameter is meaningful if applied to a single peak or band it does not give an evaluation of the whole spectrum, since to identify a Raman spectrum a "finger printing" approach is often used; analysing the most intense peak and then the following less intense ones to confirm or refute the identification. In this study only the signal to noise ratio of highest peak of every spectrum was used to evaluate the whole spectrum. However, spectra showing only one peak were not included because a single peak is insufficient to unequivocally identify a pigment.⁷⁰ In spectroscopy, the signal to noise ratio, SNR, is defined as

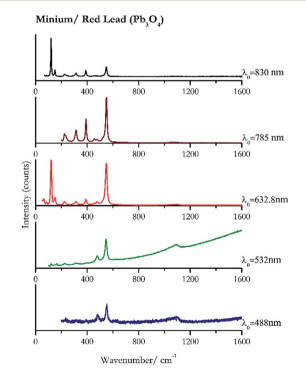


Fig. 1 Raman spectra of minium or red lead.

Table 1Classification of the spectra. ++ very good spectrum; $\pm =$ good spectrum; $\pm =$ spectrum with peaks but pigment not identifiable;- = no spectrum

Pigment	488 nm	532 nm	632.8 nm	785 nm	830 nm
Minium	+	+	++	++	++
Haematite	\pm	+	+	+	++
Red ochre	+	+	+	+	+
Caput mortuum	+	+	+	++	++
Vermillion	+	+	++	++	++
Cinnabar	+	+	++	++	++
Realgar	_	±	++	++	++
Kermes	_	_	_	-	_
Cochineal	±	+	_	+	_
Orcein	+	+	+	+	+
Brazil wood	-	-	_	_	_
Purple madder	+	+	+	+	+
Alizarin crimson	+	±	+	+	+
Alizarin purple	+	_	_	+	+
Raw umber	±	+	\pm	_	+
Sepia	_	+	+	+	+
Indigo	±	+	+	+	++
Azurite	+	+	+	+	+
Ultramarine	+	++	+	+	+
Orpiment	+	++	++	++	++
Lead tin yellow I	++	++	++	++	++
Yellow ochre	+	+	+	+	++
Massicot	++	++	++	++	++
Gamboge	+	\pm	_	+	+
White lead	+	+	++	++	++
Verdigris	+	+	+	±	+
Malachite	+	+	+	-	+
Carbon black	+	+	+	\pm	+
Ivory black	+	+	+	+	+
Lamp black	+	+	+	+	+
Iron gall	-	+	_	±	±
Bistre	+	\pm	+	+	+

$$SNR = \frac{I}{\sigma}$$

where *I* is the average intensity (net peak height – background) of the signal and σ is its standard deviation.⁷¹ However, the noise is the result of different sources. So that

$$\sigma = \sqrt{(\sigma_{\rm s}^2 + \sigma_{\rm b}^2 + \sigma_{\rm d}^2 + \sigma_{\rm r}^2)}$$

where σ_s is the uncertainty of the measurements known as signal shot noise. σ_b is the noise due to the background, that includes fluorescence of the sample and stray light. σ_d is the noise generated by the dark current of the detector and the σ_r is the readout noise caused by the conversion from the electronic signal to a digital value in the CCD camera (and subsequent transfer from the detector to the computer).⁷¹ The signal shot noise is the square root of the signal intensity according to the Poisson statistic, since the emission and detection of photons are random event.^{72,73} However, background and the dark noise are detected in the same way as the Raman signal and similarly they are equal to the square root of the background intensity and dark current⁷¹ respectively. Finally, the read out noise is the standard deviation of the numerical value the electrons from the detector device are converted into when digitized.⁷¹ Thus, the overall signal to noise ratio is defined as

$$\text{SNR} = \frac{I}{\sqrt{\left(\sigma_{\text{s}}^2 + \sigma_{\text{b}}^2 + \sigma_{\text{d}}^2 + \sigma_{\text{r}}^2\right)}}$$

To calculate the signal to noise ratio, the contribution of the dark noise and the read out noise were considered negligible, which is appropriate for the scientific grade CCD camera employed in the spectrometers, while the background noise was the result of the square root of the difference of intensities between the spectra before and after background correction. A peak may be defined as at least 2 or 3 times the intensity of the noise.^{70,72} So that the spectra were classified as "very good" spectrum (++) when the SNR > 100; "good spectrum" (+) when 3 < SNR < 100; "spectrum not identifiable" (\pm) when SNR < 3 and/ or the spectrum presented only one peak; "no spectrum" (–) when no spectrum at all was recorded.

Materials

Pigments and inks investigated were both pigments and dyes, chosen in accordance with the literature, most commonly used in manuscripts between Vth-XVIth centuries, supplied by L. Cornelissen & Son (London) and Kremer Pigmente GmbH & Co. KG (Aichstetten, Germany). Iron gall ink, Brazil wood and kermes were made following ancient recipes.74 Analysis of pure pigments using the 532 nm and 632.8 nm lasers was made by sampling through the wall of a glass vial containing the pigments. Indeed, the use of a confocal microscope allows collecting radiation coming only from the focal plane, so that there is no signal related to the glass.⁷² However, using the 785 nm excitation source the spectra presented a large background at around 1400 cm⁻¹ (ref. 75) so pellets of pigments were prepared to obtain Raman spectra without glass contribution. The measurements performed with 488 nm and 830 nm excitation were also run on pellets. They were prepared by pressing a mixture of the pigment and a 10% w/w of a wax binder, (BM-0002-1CEREOX® Licowax C Micropowder). To ensure the homogeneity of the samples the mixture of wax and pigment was shaken for 3 minutes with a frequency of 25 s^{-1} , and pellets were then formed using a hydraulic press, with 9 tonnes per surface pressure. The spectra collected do not show any signals attributable to the wax.

Results and discussion

A total of 32 pigments have been analysed using 5 different incident wavelength laser sources. The spectra are represented in Fig. 1–30 (in the ESI†). They are ordered by observed colour (red, purple, blue, yellow, white, green, black and inks). In Table 1 the positive identifications are summarized. Tables 2 to 6 list the wavenumber of the main peaks detected, with references from previous works,^{37–39,76–81} where spectra have been reported for similar conditions. Each table refers to a single wavelength laser source. The first column provides the highest observed peak in the measured spectral range for that wavelength. In the

Table 2 Characteristic peaks of Raman spectra of pigments acquired with a 488 nm excitation source. Broad bands are labelled with "br", shoulder bands are labelled with "sh"

$\lambda_0 = 488 \text{ nm}$	Spectral range			Power densit
Main band cm^{-1}	$(200-2500 \text{ cm}^{-1})$	Pigment name		mW μm^{-2}
200	458, 291, 524, 276, 550sh, 379, 305, 616, 595	Lead tin yellow I	Tin(11) sulfide, lead(11) stannate, Pb ₂ SnO ₄	3.90
225	291, 409	Caput Mortuum	Iron(III) oxide, Fe_2O_3	0.39
255	346	Cinnabar	Mercury(II) sulfide, HgS	0.39
255	346, 288sh	Vermillion	Synthetic mercury(II) sulfide, HgS	0.39
280br	391	Haematite	Iron(III) oxide, Fe_2O_3	1.02
289	386, 424,	Massicot	Lead(II) oxide, PbO	3.90
324	951, 233, 942sh, 1440, 1419, 216sh, 255, 1360, 688, 1668, 1055	Verdigris	Copper(π) acetate, Cu (CH ₃ COO) ₂ [Cu(OH) ₂] ₃ 2H ₂ O	3.90
354	311, 294, 382, 203,	Orpiment	Arsenic(m) sulfide, As ₂ S ₃	3.90
402	466, 1575, 1429, 1420, 1095, 249, 765, 838, ³⁹ 938, 1460, 1495, 541, 281, 267	Azurite	Basic copper(π) carbonate, Cu ₃ (CO ₃) ₂ (OH) ₂)	1.99
459	1497br, 1617br	Raw umber	Iron(III) oxide, Fe ₂ O ₃ + Manganese(IV) oxide, MnO ₂	3.90
461	283br, 695br	Red ochre	Iron(III) oxide, $Fe_3O_4 + clay + silica$	3.90
549	584, 1096, 1648, 256, 806, 2191, 1363	Ultramarine	$Na_6Ca_6(Al_6SI_6O_{24})$ (SO ₄ , S, S ₂ , S ₃ , cl, OH) ₂	3.90
550	480, 1094br, 390, ³⁷ 233	Minium/Red lead	Lead (II, IV) oxide, Pb_3O_4	0.39
1027	1008, 385, 1133, 491, 418sh, 297, 632.8, 547, 676, 241, 1084	Yellow ochre	Iron(III) oxide hydrate, $Fe_2O_3 H_2O$ + clay + silica	3.90
1052	1056sh, 1365br, 1297, 693, 422br, 1134	White lead	Basic lead(π) carbonate, 2PbCO ₃ Pb(OH) ₂	0.39
1306	1479, ⁸² 1702, ⁸² 1255	Cochineal	Carminic acid, C ₂₂ H ₂₀ O ₁₃	0.08
1321 (ref. 82)	1478, 1280sh, 1161 (ref. 82)	Purple madder	Alizarin C ₁₄ H ₈ O ₄ and purpurin C ₁₄ H ₈ O ₅	3.99
1322 (ref. 82)	1476, ⁸² 1297sh, 1271	Alizarin purple	Alizarin C ₁₄ H ₈ O ₄	3.90
1477 (ref. 82)	1325, ⁸² 1290, ⁸² 1163, 906, 839, 660, ⁸² 480	Alizarin crimson	Alizarin $C_{14}H_8O_4$	3.90
1495	434, 225, 272, 536, 1062, 1103, 1367, 356, 1297, 721, 755	Malachite	Basic copper(11) carbonate, Cu ₂ CO ₃ (OH) ₂	3.90
1573br	1347br	Lamp black	Carbon, C	3.90
1581br	1361 br	Carbon black	Carbon, C	3.90
1585	1703, 1363, 1252	Indigo	Indigo, $C_{16}H_{10}N_2O_2$	3.90
1592br		Bistre	Wood soot, carbon C	1.99
1594br	1362br, 469br	Ivory black	Calcium hydroxide phosphate, Ca ₅ (OH) (PO ₄) ₃ + carbon, C	3.90
1594	1632	Gamboge	Gambogic acid, C ₃₈ H ₄₄ O ₈	3.90
1642	1608, 1505br, 1278, 1192br, 489	Orcein	Natural red 28, C ₂₈ H ₂₄ N ₂ O ₇	3.90

second column the other peaks, in decreasing order of intensity, can be found; in the other columns the name and the compounds of the pigment. In the last column, the values of power density are recorded.

To use the library:

(1) Select the table pertinent to the laser wavelength used to carry out the measurements;

(2) Look for the highest intensity peak in the first column;

(3) Check the other peaks in the second column. The third column provides the name of the pigment.

Red pigments

For convenience, the pigments can be divided into two major groups. The first is the iron oxide compounds (haematite, red ochre and caput mortuum, which belongs also to the inks group

as well), and the others (cinnabar, vermillion, minium/red lead). All the iron oxide (Fig. 2-4) compounds show the main peaks at around 223 cm⁻¹, between 290 cm⁻¹, around 407 cm⁻¹, but only the spectra acquired with 532 nm and the 632.8 nm excitation beams show an intense peak between 1316 cm⁻¹ and 1323 cm⁻¹. The intensity enhancement is attributed to resonance effects since the absorption edge for Fe₂O₃ is at 580 nm. Spectra acquired with 488 nm all have poor signals, because of absorption by the pigment. Unfortunately, the red ochre (Fig. 3), which is a mixture of iron oxides, clays and silica, is the one that provided lowest signal to noise ratio with all the laser sources compared to haematite and caput mortuum. The red ochre spectra are indeed affected by fluorescence, likely related to the presence of a heterogeneous matrix. In the second group, minium (Fig. 1), or red lead, is a lead oxide, whose the main peak resulting from 632.8 nm

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A ₀ = 532 nm Spectral range Power d				
Main band cm ⁻¹	(100-2500 cm ^{-1})	Pigment name		Power densi mW μm ⁻²
130 (ref. 83)	96, ⁸³ 458, 293, 275, 252, 379, ⁷⁷ 304, 615 (ref. 77)	Lead tin yellow I	Tin(11) sulfide, lead(11) stannate, Pb ₂ SnO ₄	2.06
143 (ref. 77)	290, 386, 425, 169sh	Massicot	Lead(II) oxide, PbO	0.82
178 (ref. 76, 77 and 84)	151, ^{76,77} 169, ^{76,84} 430, ⁷⁷ 220, ⁷⁷ 266, ^{76,84} 1491, ^{76,84} 203sh, 532, ^{76,77} 535sh, ^{76,77,84} 350sh, ^{76,77} 509, ⁷⁷ 1092, ^{76,84} 1060, ^{76,77} 720, ^{76,84} 1459, ⁷⁷ 751, ^{77,84} 593	Malachite	Basic copper(π) carbonate, Cu ₂ CO ₃ (OH) ₂	2.06
254	290, 342	Cinnabar	Mercury(II) sulfide, HgS	0.08
254 (ref. 77)	343, ⁷⁷ 290	Vermillion	Synthetic mercury(II) sulfide, HgS	0.08
268	214, 582br	Raw umber	Iron(III) oxide, Fe ₂ O ₃ + Manganese(IV) oxide, MnO ₂	8.21
324	949, ^{76,84} 127, ^{76,84} 1439, ^{76,84} 297, 183, ⁷⁷ 234, ⁷⁶ 703, ^{76,77} 1296, 254, ⁸⁴ 1061, ⁷⁶ 1133, 1641br	Verdigris	Copper(n) acetate, Cu (CH ₃ COO) ₂ [Cu(OH) ₂] ₃ 2H ₂ O	2.06
354 (ref. 83)	311, 293, 382, 154, 203, ⁸³ 136, ⁸³ 180, ⁸³ 234, 490 br, 589br	Orpiment	Arsenic(m) sulfide, As_2S_3	0.82
356 (ref. 83)	343, ⁷⁷ 338, 192, ⁸³ 183, ⁷⁷ 221	Realgar	Arsenic(m) sulfide, As ₄ S ₄	0.01
400 (ref. 77)	247, ⁷⁷ 1096, ⁷⁷ 1578, ⁷⁷ 1430, ⁷⁷	Azurite	Basic copper(II) carbonate,	2.06
	1417, 132, 171, 281, ⁷⁷ 178, 138, ⁷⁷ 266, 154, 839, 764, ⁷⁷ 542, 737, ⁷⁷ 336		$Cu_3(CO_3)_2$ (OH) ₂)	
547	1089br, 479, 474, 389, 313, 227, 163(a)	Minium/Red lead	Lead (II, IV) oxide, Pb_3O_4	0.82
548 (ref. 77)	1097, ⁷⁷ 1647, ⁷⁷ 2191, ⁷⁷ 1644, ⁷⁷ 257, ⁷⁷ 864, 275, 1369, 1905 (ref. 77)	Ultramarine	$\begin{array}{l} Na_{6}Ca_{6}(Al_{6}SI_{6}O_{24}) \left(SO_{4},S,S_{2},S_{3},\\ cl,OH\right)_{2} \end{array}$	2.06
1007	388, 416, 299, ⁷⁷ 1138, 245, ⁸³ 549, 495, 675	Yellow ochre	Iron(III) oxide hydrate, Fe ₂ O ₃ H ₂ O + clay + silica	2.06
1050 (ref. 77)	131, 1370br, ⁷⁷ 151sh, 418sh, 1635	White lead	Basic lead(II) carbonate, 2Pb CO ₃ Pb(OH)2	0.82
1316	291, 227, 409, 613, 497, 247	Caput Mortuum	Iron(III) oxide, Fe_2O_3	0.82
1321	1479, 1159 (ref. 77)	Purple madder	Alizarin $C_{14}H_8O_4$ and purpurin $C_{14}H_8O_5$	0.82
1322	(b)	Cochineal	Carminic acid, C ₂₂ H ₂₀ O ₁₃	0.82
1322	1478	Alizarin purple	Alizarin C ₁₄ H ₈ O ₄	0.08
1323	290, 223, 407, 600, 238	Haematite	Iron(III) oxide, Fe_2O_3	0.82
1351	460, 299, 674, 629, 230, 414	Red ochre	$Iron(III)$ oxide, $Fe_2O_3 + clay + silica$	0.82
1480 (ref. 77)	1326	Alizarin crimson	Alizarin C ₁₄ H ₈ O ₄	0.08
1574br	1405br	Sepia	Melanin C ₁₈ H ₁₀ N ₂ O ₄	0.82
1572	1335	Lamp black	Carbon, C	8.21
1582	1361, 1701, 1251, 1628, 546, 598, 1461, 1485, 1312, 941, 756, 249, 1222	Indigo	Indigo, $C_{16}H_{10}N_2O_2$	8.21
1582	1346	Carbon black	Carbon, C	8.21
1587	1351 (ref. 77)	Ivory black	Calcium hydroxide phosphate, Ca ₅ (OH) (PO ₄) ₃ + carbon, C	8.21
1592br		Bistre	Wood soot	2.06
1600	1632.8	Gamboge	Gambogic acid, C ₃₈ H ₄₄ O ₈	2.06
1639	1499, 1335, 1275, 1191, 483	Orcein	Natural red 28, C ₂₈ H ₂₄ N ₂ O ₇	2.06
1644	1601, 1353, 560(c)	Iron Gall ink	Ferric Gallate C ₂₁ H ₁₅ FeO ₁₅	0.96

Table 3Characteristic peaks of Raman spectra of pigments acquired with a 532 nm excitation source (a) range between 100 and 2500 cm $^{-1}$, (b)range between 108 and 2500 cm $^{-1}$, (c) range between 150–2500 cm $^{-1}$. Broad bands are labelled with "br", shoulder bands are labelled with "sh"

excitation is at 121 cm⁻¹, due to the deformation of the O–Pb–O angle. This was not detectable with the other wavelengths because of the edge filters cutting off low wavenumbers or attenuating the signal, and in literature studies of manuscripts this band is often omitted for this reason. However, contrary to the report by Burgio *et al.*¹ who reported sample damage when

using 488 nm and 514.5 nm radiations, it was possible to detect the pigment thanks to a low power density at 0.39 mW μ m⁻². Cinnabar and vermillion (Fig. 5 and 6) are the mineral and synthetic forms of mercury sulphide, both show indeed the same very strong peak at 251–255 cm⁻¹. They are both detectable with all the five wavelengths, and only low laser power is Table 4 Characteristic peaks of Raman spectra of pigments acquired with a 632.8 nm excitation source. (a) Range between 80 and 2500 cm⁻¹, (b) range between 50 and 2500 cm⁻¹. Broad bands are labelled with "br", shoulder bands are labelled with "sh"

$\lambda_0 = 632.8 \text{ nm}$ Main band cm ⁻¹	Spectral range (50–2500 cm ⁻¹)	Pigment name		Power densit mW μm ⁻²
73	1438, 950, 183, 232, 1300, 315, 1133, 1647, 1064(b)	Verdigris	Copper(II) acetate, Cu $(CH_3COO)_2$	1.16
104	70, 1050, 1365, 1475br, 412, 963,	White lead	$[Cu(OH)_2]_3 2H_2O$ Basic lead(II) carbonate, 2Pb CO ₃	2.63
121 (ref. 37, 38	677, 325(a)	Minium/red lead	$Pb(OH)_2$	0.63
and 77)	549, ^{37,38,77} 149, ⁷⁷ 390, ^{37,38,77} 65, 223, ^{37,38,77} 313, ^{37,38,77} 480, ^{37,38} 84, ³⁸ 456sh, 290sh, 1094br ⁷⁷ (b)	Miniuni/red lead	Lead (II, IV) oxide, Pb ₃ O ₄	2.63
127 (ref. 77)	77, 193, ⁷⁷ 456, ⁷⁷ 273, ⁷⁷ 290, ⁷⁷	Lead tin yellow I	Tin(11) sulfide, lead(11) stannate, Pb ₂ SnO ₄	1.30
142 (ref. 37, 38, 77)	524, ⁷⁷ 377, ⁷⁷ 336br 288, ^{37,38,77} 86, ³⁸ 70, ³⁸ 384, ^{37,38,77} 215br, 423 (ref. 38)	Massicot	Lead(II) oxide, PbO	2.63
162	188, 89, 129, 226, 127, 439, 278, 359, 1501, 506, 1377	Malachite	Basic copper(II) carbonate, Cu ₂ CO ₃ (OH) ₂	2.63
251 (ref. 37 and 77)	342, ^{37,77} 280sh, ^{37,77} 86, 104sh	Vermillion	Synthetic mercury(II) sulfide, HgS	2.63
253	343, 283, 85, 102	Cinnabar	Mercury(II) sulfide, HgS	2.63
291 (ref. 78)	1316, 226, ⁷⁸ 408, ⁷⁸ 610, ⁷⁸ 822br, 658br, 243sh	Caput Mortuum	Iron(\mathbb{m}) oxide, Fe ₂ O ₃	2.63
293 (ref. 39)	1320, 409, ³⁹ 609, ^{38,39} 224, ³⁹ 1086br, 660, 243, ^{38,39} 495, ^{38,39} 820br	Haematite	Iron(III) oxide, Fe_2O_3	0.26
354 (ref. 37 and 77)	309, ^{37,77} 291, ³⁷ 153, ^{37,77} 381, ³⁷ 201, ³⁷ 66, 180, ³⁷ 104	Orpiment	Arsenic(\mathfrak{m}) sulfide, As ₂ S ₃	1.30
354 (ref. 37)	185, ⁷⁷ 191, ³⁷ 221, ^{37,77} 230sh, 368sh, ³⁷ 58, 168sh, 143, ^{37,77} 120	Realgar	Arsenic(III) sulfide, As ₄ S ₄	1.30
408	292, ³⁷ 224, 657br, 610br	Red ochre	$Iron(III)$ oxide, $Fe_3O_4 + clay + silica$	0.64
545 (ref. 77)	1096, ⁷⁷ 86, 1370, 1662, 255, 286sh ⁷⁷	Ultramarine	$Na_6Ca_6(Al_6SI_6O_{24})$ (SO ₄ , S, S ₂ , S ₃ , cl, OH) ₂	2.63
657 br (ref. 78)		Raw umber	Iron(III) oxide, $Fe_2O_3 + Manganese(IV)$ oxide, MnO_2	0.26
1005 (ref. 37)	385, ^{37,38} 1134, 1084, 670, 617, 547, 488, ³⁸ 296, ³⁷ 241, 92 (ref. 38)	Yellow ochre	Iron(III) oxide hydrate, $Fe_2O_3 H_2O$ + clay + silica	2.63
1095 (ref. 38)	835, 396, 761, ³⁸ 476br, 79, 240, ³⁸ 133, 171	Azurite	Basic copper(π) carbonate, Cu ₃ (CO ₃) ₂ (OH) ₂)	2.63
1324 (ref. 81)	1293, 1475, 1448, ⁸¹ 1157, ⁸¹ 825 (ref. 81)	Purple madder	Alizarin $C_{14}H_8O_4$ and purpurin $C_{14}H_8O_5$	2.63
1324	1594br	Carbon black	Carbon, C	4.68
1333 (ref. 77)	1585	Lamp black	Carbon, C	2.63
1477	1325, ⁸¹ 1291, 1187, ⁸¹ 1160, ⁸¹ 898, ⁸¹ 837	Alizarin crimson	Alizarin C ₁₄ H ₈ O ₄	0.26
1573br		Iron ink Gall	Ferric Gallate C ₂₁ H ₁₅ FeO ₁₅	2.63
1578	1253, 1223, 756, 672, 596, 545, 309, 250, 274	Indigo	Indigo, $C_{16}H_{10}N_2O_2$	2.63
1585	1373br	Bistre	Wood soot, carbon C	0.64
1592br	1392br	Sepia	Melanin C ₁₈ H ₁₀ N ₂ O ₄	0.33
1597	1349 (ref. 77)	Ivory black	Calcium hydroxide phosphate, Ca ₅ (OH) (PO ₄) ₃ + carbon, C	2.63
1644	1518, 1408, 1277, 1189, 629, 595, 821, 525, 479	Orcein	Natural red 28, $C_{28}H_{24}N_2O_7$	2.63

required when working with 532 nm and 488 nm lasers to prevent saturation of the detector even at short acquisition times. Realgar is a photosensitive mineral95 and its transformation to pararealgar occurred using the 532 nm and the 488 nm wavelength laser source. Unfortunately the minimum power level for both the configurations was sufficient for this transformation and good spectra were not obtained without damaging the sample (Fig. 7).

Purple pigments

Cochineal (Fig. 8), orcein (Fig. 9), brazil wood, kermes, purple madder (Fig. 10), alizarin crimson (Fig. 11) and alizarin purple (Fig. 12) are all organic compounds and, these spectra are prone to be affected by fluorescence depending on the wavelength of the excitation source. The spectrum collected with the 830 nm and 632.8 nm of cochineal does not show any Raman bands.

Paper

Table 5 Characteristic peaks of Raman spectra of pigments acquired with a 785 nm excitation source. (a) Range between 200–2500 cm⁻¹. Broad bands are labelled with "br", shoulder bands are labelled with "sh"

$\lambda_0 = 785 \text{ nm}$				Power
Main band cm- ¹	Spectral range (100–2500 cm^{-1})	Pigment name		density V μm ⁻²
549 (ref. 77)	391, 314, ⁷⁷ 224, ⁷⁷ 234sh, 456, 480(a)	Minium/red lead	Lead (II, IV) oxide, Pb_3O_4	3.58
107	1051, ⁷⁷ 1055sh, 410, 320br, 679, ⁷⁷ 1136, 1441, 1640	White lead	Basic lead(II) carbonate, 2Pb CO ₃ Pb(OH) ₂	3.58
129 (ref. 77)	196, ⁷⁷ 457, ⁷⁷ 291, ⁷⁷ 274, 111, 524, ⁷⁷ 379, ⁷⁷ 337, 508, 304, 337, 432sh	Lead tin yellow I	Tin(n) sulfide, lead(n) stannate, Pb ₂ SnO ₄	3.58
142 (ref. 77)	288, ⁷⁷ 384, ⁷⁷ 214, 427	Massicot	Lead(II) oxide, PbO	3.58
223 (ref. 77)	290, ⁷⁷ 406 (ref. 77)	Red ochre	Iron(III) oxide, $Fe_3O_4 + clay + silica$	3.58
253	343, 286, 107, 143	Vermillion	Synthetic mercury(II) sulfide, HgS	0.96
254	343, 287, 108, 143, 201	Cinnabar	Mercury(II) sulfide, HgS	0.32
290	408, 224, 244, 609, 498, 1323br	Haematite	Iron(III) oxide, Fe_2O_3	3.58
290	224, 407, 609, 244, 495, 1316	Caput mortuum	$Iron(III)$ oxide, Fe_2O_3	3.58
353 (ref. 77)	292, 202, ⁷⁷ 383, 325, 218, 472, 585, 653, 706	Orpiment	Arsenic(m) sulfide, As ₂ S ₃	3.58
354 (ref. 77)	192, ⁷⁷ 181, 220, 343, ⁷⁷ 142, ⁷⁷ 165, 171, ⁷⁷ 368, 374, 328, 212	Realgar	Arsenic(m) sulfide, As_4S_4	0.96
397	246, 464, 128, 1095, 1428, 762, 834, 1574	Azurite	Basic copper(II) carbonate, Cu ₃ (CO ₃) ₂ (OH) ₂)	3.58
482	1274, 1188, ⁹⁰ 592, ⁹⁰ 524, ⁹⁰ 809, 1463, 1492, 1640 (ref. 90)	Orcein	Natural red 28, $C_{28}H_{24}N_2O_7$	3.58
550 (ref. 77)	584sh, ⁷⁷ 256	Ultramarine	$Na_6Ca_6(Al_6SI_6O_{24})$ (SO ₄ , S, S ₂ , S ₃ , cl, OH)	3.58
980	1430, 1338, 573 br	Iron gall ink	Ferric Gallate C ₂₁ H ₁₅ FeO ₁₅	4.94
.007	1255, 1224, 384, 431, 1138, 495, 1410, 671	Yellow ochre	Iron(m) oxide hydrate, Fe ₂ O ₃ H ₂ O + clay + silica	3.58
1248br	1560br	Bistre	Wood soot, carbon C	4.94
1293	1318, ⁹⁰ 1442	Purple madder	Alizarin $C_{14}H_8O_4$ and purpurin $C_{14}H_8O_5$	3.58
1303 (ref. 90)	1321, ⁷⁷ 1473	Cochineal	Carminic acid, C ₂₂ H ₂₀ O ₁₃	3.58
1350br	1550br	Sepia	Melanin C ₁₈ H ₁₀ N ₂ O ₄	3.58
1430,	1136, 227	Verdigris	Copper(II) acetate, Cu (CH ₃ COO) ₂ [Cu(OH) ₂] ₃ 2H ₂ O	3.58
1431	1584, 1304	Lamp black	Carbon, C	3.58
1474	1302, 1323 (ref. 90)	Alizarin purple	Alizarin C ₁₄ H ₈ O ₄	3.58
1480 (ref. 90)	1328, 1292, ⁷⁷ 480, 1192, 1462sh, 1451sh, ⁹⁰ 841, 1163, ^{77,90} 659, 904 (ref. 77 and 90)	Alizarin crimson	Alizarin C ₁₄ H ₈ O ₄	3.58
1571	1580 sh, 250, 545, ⁷⁷ 597, ⁷⁷ 262, 674, ⁷⁷ 273, 756, ⁷⁷ 1223, 1308, 1459, 1362, 1015 (ref. 77)	Indigo	Indigo, $C_{16}H_{10}N_2O_2$	3.58
1576	1332	Carbon black	Carbon, C	3.58
1585 (ref. 77)	1329	Ivory black	Calcium hydroxide phosphate, Ca ₅ (OH) (PO ₄) ₃ + carbon, C	3.58
1592 (ref. 77)	1297, 1620sh, 1136, 1062	Gamboge	Gambogic acid, $C_{38}H_{44}O_8$	3.58

Orcein, purple madder and alizarin crimson were detectable with all the wavelengths, while alizarin purple did not provide any spectrum with the 532 nm (absorption at 530 nm) and 632.8 nm radiation. For kermes and brazil wood, no spectra could be obtained at any of the five laser wavelengths.

Blue pigments

Indigo (Fig. 14) provides better spectra with the NIR and IR laser since it possesses a broad absorption band in the visible

range.⁸⁵ However, it is still possible to recognise the pigment, thanks to the peak at 1580 cm⁻¹ circa and 545 cm⁻¹ also with lower wavelengths, which are superimposed upon the weak fluorescence from this material. Azurite (Fig. 15) yields a good spectrum with all the wavelengths except the 632.8 nm laser source, attributed to the strong absorption by the pigment at *ca.* 600 nm. When using the infrared sources at 785 and 830 nm, a very low energy (785 nm and 830 nm, 3.58 and 8.69 mW μ m⁻²) was used: at higher levels absorption of the radiation and localised heating of the sample results in its degradation. When

Analytical Methods

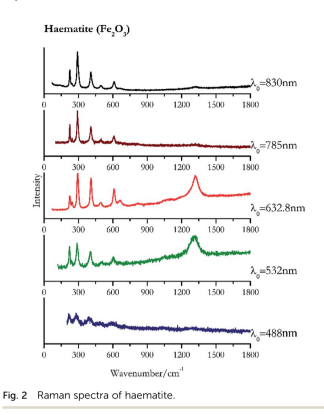
Table 6Characteristic peaks of Raman spectra of pigments acquired with a 532 nm excitation source. (a) Range between $100-2500 \text{ cm}^{-1}$.Broad bands are labelled with "br", shoulder bands are labelled with "sh"

$\frac{\lambda_0 = 830 \text{ nm}}{\text{Main band cm}^{-1}}$	Spectral Range (100–2500 cm ⁻¹)	Pigment name	Compound	Power densit mW μm ⁻²
	· · · · · ·			
92	315, 949, 179, 1296, 1436br	Verdigris	Copper(II) acetate, Cu (CH ₃ COO) ₂ [Cu(OH) ₂] ₃ 2H ₂ O	0.86
105	74, 1049, 1053, 416br, 679	White lead	Basic lead(II) carbonate, 2Pb CO ₃ Pb(OH) ₂	0.12
120	549, 390, 151, 314, 142sh, 230br, 63, 85, 454	Minium/red lead	Lead (II, IV) oxide, Pb_3O_4	0.86
128 (ref. 79)	79, 196, ⁷⁹ 457, ⁷⁹ 292, ⁷⁹ 273, ⁷⁹ 112, 524, ⁷⁹ 379, ⁷⁹ 96, 304, ⁷⁹ 339, 432	Lead tin yellow	Tin(11) sulfide, lead(11) stannate, Pb ₂ SnO ₄	0.37
142	87, 289, 70, 384, 217	Massicot	Lead(π) oxide, PbO	0.37
150	179, 77, 218, 269, 430, 1061br, 1495	Malachite	Basic copper(π) carbonate, Cu ₂ CO ₃ (OH) ₂	0.86
254 (ref. 79)	343, ⁷⁹ 283, ⁷⁹ 103, 85, 142	Vermillion	Synthetic mercury(II) sulfide, HgS	0.12
254 (ref. 79)	$343^{79}_{,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,$	Cinnabar	Mercury(n) sulfide, HgS	0.12
287	222, 406, 605, 489, 241sh	Caput Mortuum	Iron(\mathfrak{m}) oxide, Fe ₂ O ₃	2.85
291	225, 408, 612, 244, 498, 1321br	Haematite	Iron(III) oxide, Fe_2O_3	2.85
292	610br, 225, 390br, 725	Raw umber	Iron(III) oxide, $Fe_2O_3 +$	1.24
			Manganese(v) oxide, MnO ₂	
293	405, 398, 221, 608	Red ochre	Iron(III) oxide, $Fe_3O_4 + clay + silica$	2.85
353	191, 182, 220, 342, 166, 171, 142,	Realgar	Arsenic(II) sulfide, As_4S_4	0.37
000	367, 374, 328, 123	iteuigui	moeme(n) sumae, ms454	0.07
354	311, 293, 154, 202, 136, 382, 369sh, 179, 105, 69	Orpiment	Arsenic(\mathfrak{m}) sulfide, As ₂ S ₃	0.12
401	248, 1093, 171(a)	Azurite	Basic copper(n) carbonate, Cu ₃ (CO ₃) ₂ (OH) ₂)	0.86
488	1275, 1192, 600br, 1333br,887, 1080, 432, 528	Orcein	Natural red 28, $C_{28}H_{24}N_2O_7$	1.24
541		Ultramarine	$Na_6Ca_6(Al_6S_{I6}O_{24})$ (SO ₄ , S, S ₂ , S ₃ , cl, OH) ₂	8.69
979	(a)	Iron Gall ink	Ferric Gallate C ₂₁ H ₁₅ FeO ₁₅	8.69
1007	386, 415, 297, 92, 1134, 1087, 244, 496, 550, 617, 669, 145, 1297, 1434	Yellow ochre	Iron(m) oxide hydrate, $Fe_2O_3 H_2O$ + clay + silica	8.69
1037	1596br	Lamp black	Carbon, C	0.37
1298br	1550br	Bistre	Wood soot, carbon C	1.24
1300br	1554br	Sepia	Melanin $C_{18}H_{10}N_2O_4$	0.86
1307	1593br	Ivory black	Calcium hydroxide phosphate, Ca ₅ (OH) (PO ₄) ₃ + carbon, C	0.86
1309	1589br	Carbon black	Carbon, C	0.37
1318	1292, 1470br, 1448br	Purple madder	Alizarin $C_{14}H_8O_4$ and purpurin $C_{14}H_8O_5$	1.24
1321	1472, 1301br	Alizarin purple	Alizarin $C_{14}H_8O_4$	0.12
1472	1328, 1187	Alizarin crimson	Alizarin $C_{14}H_8O_4$	0.37
1571	252, 544, 1582, 133, 101, 265, 275, 599, 1224, 1310, 235, 674, 757, 182, 635, 1364, 1460, 172, 310, 73,	Indigo	Indigo, $C_{16}H_{10}N_2O_2$	1.24
1593	1015, 1246, 1625, 1146, 870, 1702 1435, 1632, 1453sh, 1331, 370, 1221, 1247, 1281	Gamboge	Gambogic acid, $C_{38}H_{44}O_8$	8.69

the particles of azurite do not disperse the heat efficiently and the rate of heat inside the single grain is higher than the rate of heat outside, they thermo-degrade.⁸⁶ Ultramarine (Fig. 16) can be identified by the strong band at 550 cm⁻¹ using all excitation wavelengths. The absorption band at around 610 nm means that the 532 nm and 632 nm excitation sources, close to the electronic absorption wavelength, benefit from strong resonance enhancement and yield a progression of bands due to the bending of S_3^{-} .⁸⁷ Indeed, in the spectrum collected with the 488 nm shows clearer a band at 584 cm⁻¹ result of the S_2^- vibration, which with the other sources appears only as a shoulder of the main peak at 550 cm⁻¹.

Brown pigment

The only brown pigment investigated was raw umber (Fig. 13). It is a mixture of iron oxides and manganese oxides. Investigation at 830 nm shows bands at 292 cm^{-1} , 610 cm^{-1} ,



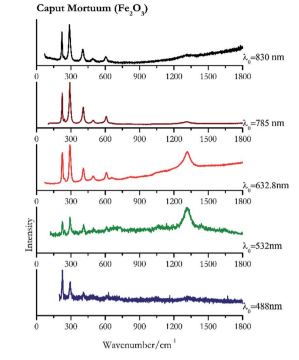


Fig. 4 Raman spectra of caput mortuum.

225 cm⁻¹, 390 cm⁻¹ and 725 cm⁻¹. Excitation with the green laser presents weak bands at 268 cm⁻¹, 214 cm⁻¹ and 582 cm⁻¹. No satisfactory Raman spectra were obtained with the other laser excitation wavelengths, 488 nm, 632.8 nm, 785 nm or 830 nm.

Yellow pigments

Orpiment (Fig. 17) is easily detectable with all the excitation wavelengths, requires only low laser powers to detect, and it may indeed cause saturation of the detector under some conditions. The main chromophore of yellow ochre is limonite, an iron oxy-hydroxide, and this pigment shows the same main

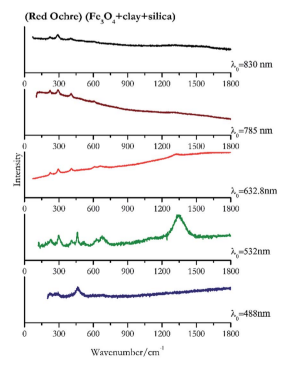


Fig. 3 Raman spectra of red ochre.

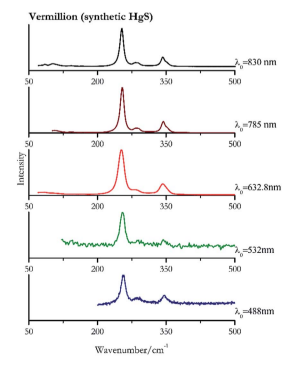
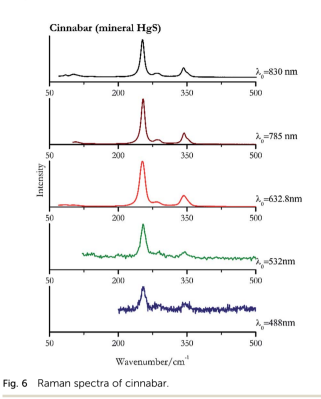
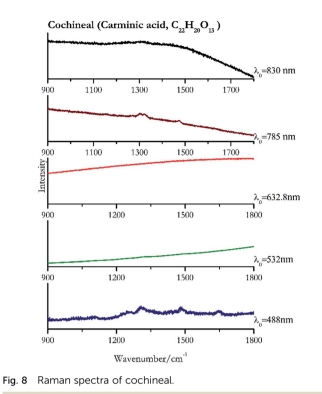


Fig. 5 Raman spectra of vermillion.





peaks of the mineral at all wavelengths. Ochres come in a range of compositions, due to natural variance of the mineral and the Raman spectrum (Fig. 18) can therefore reflect these differences when collecting spectra from real historical artefacts. Lead tin yellow type I (Fig. 19) and massicot (Fig. 20) are both easily identified with the different laser wavelengths, but extra care has to be taken since they are both lead compounds and careful management of the laser power is required to avoid photo degradation: Burgio records alteration at 10 mW with 514.5 nm excitation source.¹ In the field, this simply requires using a reliable power meter prior to measurement to ensure one knows precisely the light power being delivered at the sample

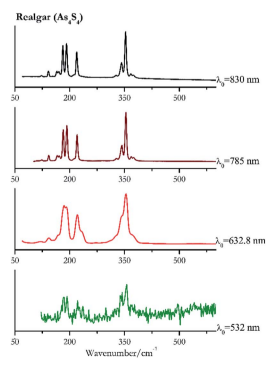


Fig. 7 Raman spectra of realgar.

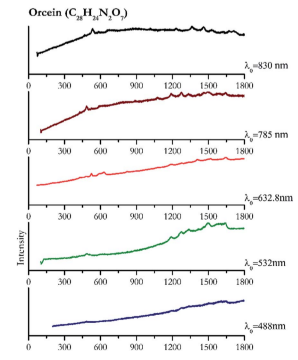
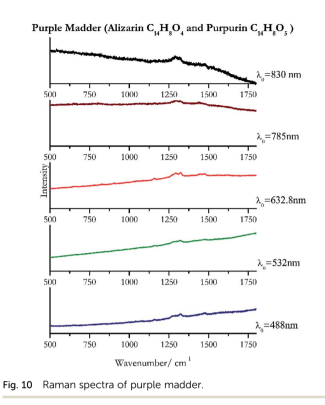
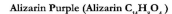
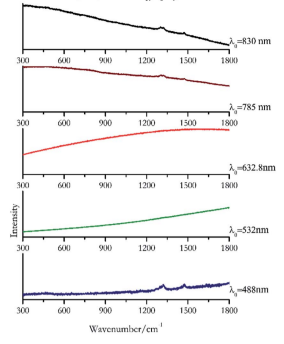


Fig. 9 Raman spectra of orcein.









surface and to work well below the damage thresholds. Lead tin yellow (Fig. 19) and massicot (Fig. 20) were detected with the 488 nm source at 3.9 mW μ m⁻² and with the 532 nm at 2.06 mW μ m⁻² for lead tin yellow type I and 0.82 mW μ m⁻² for massicot, without any degradation. No damage at the sample was noticed with the higher wavelengths and if low values of power (0.37

mW μ m⁻² with 830 nm laser) were used it was to avoid the detector saturation. The organic nature of gamboge (Fig. 21) causes fluorescence in the spectra, especially with higher frequency sources. The 830 nm laser is the one that yields the best spectrum, with an intense peak at 1593 cm⁻¹.

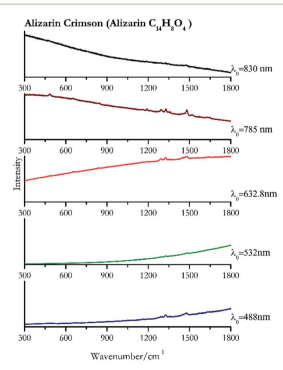


Fig. 11 Raman spectra of Alizarin crimson.

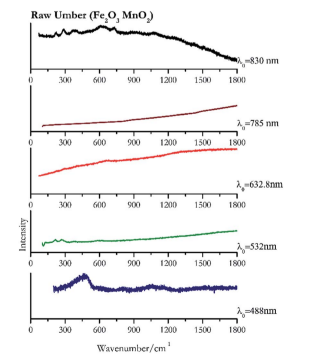


Fig. 13 Raman spectra of raw umber.

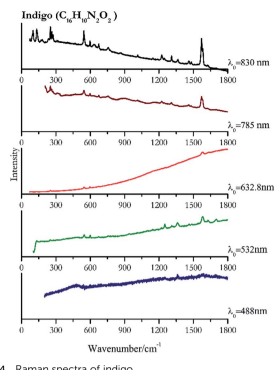
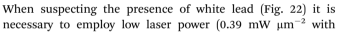


Fig. 14 Raman spectra of indigo.

White pigment



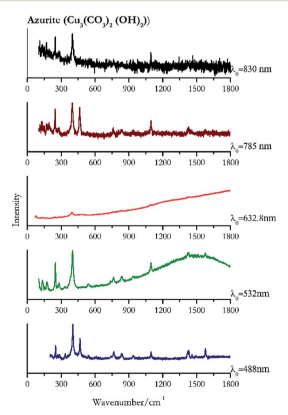


Fig. 15 Raman spectra of azurite.

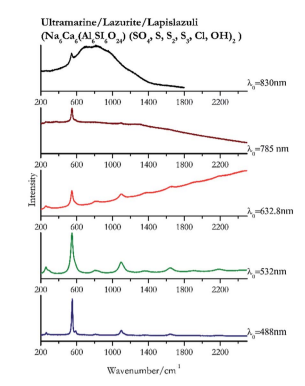


Fig. 16 Raman spectra of ultramarine.

488 nm and 0.12 mW μm^{-2} with 830 nm) because like other lead compounds it can easily be photodegraded by the laser. 88,89 The highest intensity peak is observed at 104 cm $^{-1}$ that requires a good laser light rejection cut off filter. As it can be seen from

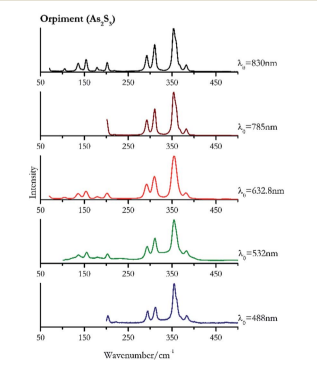
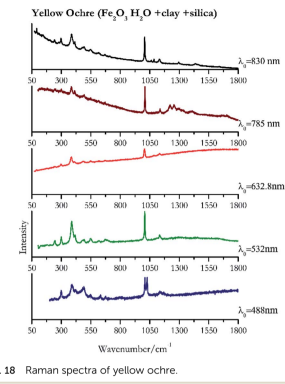
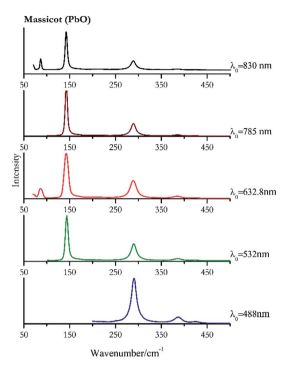
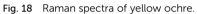


Fig. 17 Raman spectra of orpiment.

8









the spectra presented in Fig. 22 this band is clearly observed in our system when using the 632.8 nm excitation, where the notch filter has a shorter cut off. However, it remains observable using 830 nm excitation source, but lies on top of the sloping laser beam profile.

Green pigments

Verdigris (Fig. 23) and malachite (Fig. 24) both result in good Raman spectra for the shorter wavelengths and do not provide a spectrum with the 785 nm laser because of strong absorption of the light in this region.84 However, the spectra

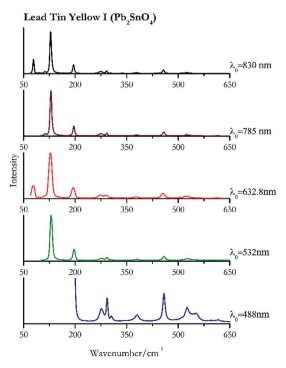


Fig. 19 Raman spectra of lead tin yellow I.

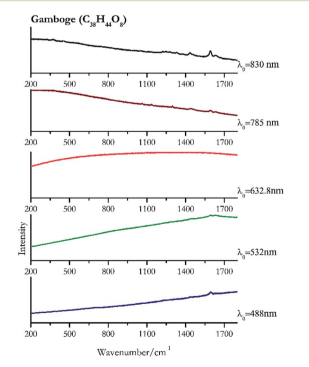


Fig. 21 Raman spectra of gamboge.

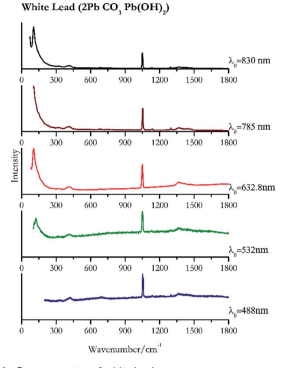


Fig. 22 Raman spectra of white lead.

obtained with the 830 nm do present peaks at 150 cm⁻¹, 179 cm⁻¹, 218 cm⁻¹, 269 cm⁻¹ to identify the pigments unambiguously.

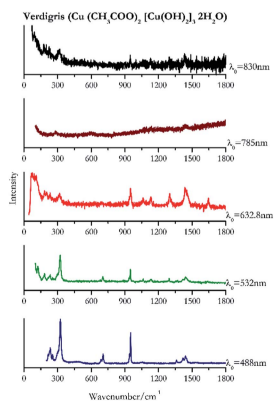
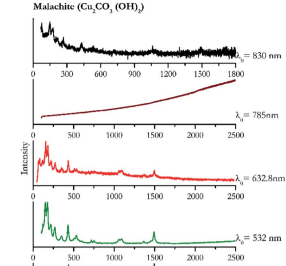


Fig. 23 Raman spectra of verdigris.



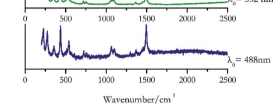


Fig. 24 Raman spectra of malachite.

Black pigments and inks

Carbon black, ivory black, lamp black and bistre (Fig. 25, 26, 27 and 28 respectively) are all characterized by broad bands between 1300 cm^{-1} and 1600 cm^{-1} due to the amorphous carbon. It is not possible to distinguish one from the others

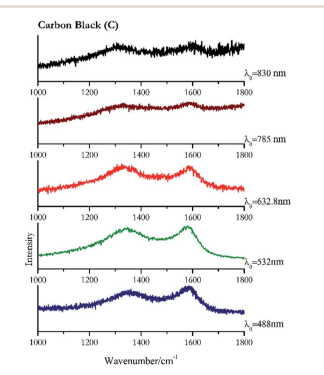
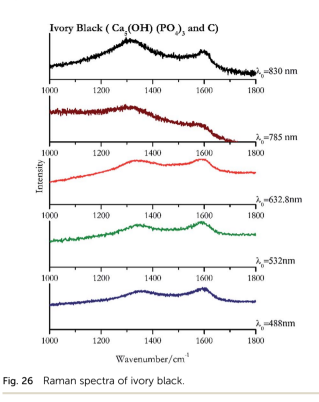
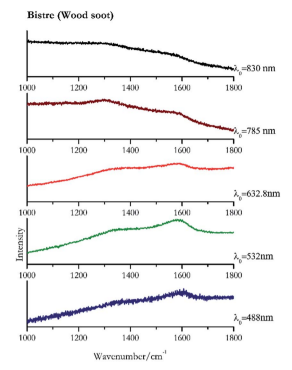
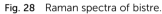


Fig. 25 Raman spectra of carbon black.







using a Raman spectroscopy.⁹¹ In a previous work a band at 965 cm⁻¹ is recorded for ivory black, but it is not present in these spectra. The 785 nm laser seemed to be the one that provided the less intense peaks for all these black pigments. The sepia also shows two broad bands (Fig. 29), but they are generated by melanin, the main constituent of the

pigment.^{92,93} Iron gall ink presents a peak at circa 980 cm^{-1} in the spectra collected (Fig. 30) with the two NIR sources, then a broad band in between 560 and 570 cm⁻¹ that can be observed in the 532 and 785 nm spectra, as well as one at about 1350 cm⁻¹. The spectrum acquired with the green laser

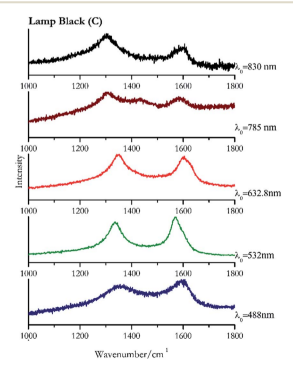


Fig. 27 Raman spectra of lamp black.

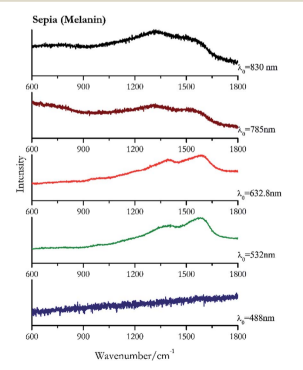
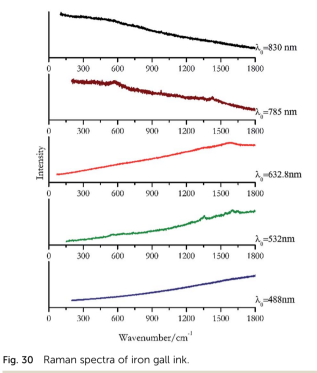


Fig. 29 Raman spectra of sepia.





has also peaks at 1601 cm⁻¹ and 1644 cm⁻¹. Historically better spectra have been obtained for this pigment,⁹⁴ however we were not able to reproduce these results at the lower power densities. Caput mortuum (Fig. 4) has been already considered among the iron oxide based red pigments.

Conclusions

This work seeks to provide an updated and useful reference handbook for identification by Raman spectroscopy of pigments of the middle ages meeting the needs of researchers working with portable equipment in situ and the increase in the availability of different wavelength lasers. Table 1 helps in choosing the best excitation wavelength to investigate expected pigments on a specific artefact. Indeed, the discussion of the spectra calls attention to the different phenomena that can occur according to the nature of the pigment and the wavelength used to investigate it, for example in considering the resonance Raman condition and how this effects relative peak intensities in comparison to normal Raman effect. The collection of spectra helps to compare the experimental data with references acquired with the same laser source, to unequivocally identify the pigment. The Tables 2-6 provide a practical guide, which in few steps allows the identification of pigments. Since the intensities of peaks vary according to the exciting source, the operator can search references recorded with the same wavelength used during the measurements. New pigment references collected with wavelengths not used in previous works are also provided. This work also concludes that it is evident that the Raman technique alone is not capable of fully characterizing all the pigments selected, especially the

fluorescent organic dyes that are presenting challenges to curators. This highlights the need to turn to other techniques that can provide complementary information.

Conflicts of interest

There are no conflicts to declare.

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