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A multi-analytical approach towards the investigation of Subarctic Athapaskan colouring of quillwork and its sensitivity to photo-degradation

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Abstract

Non-European dyed materials other than textiles [1] have received comparatively little systematic analysis, this is particularly true for objects made with dyed porcupine quills. This paper presents a comprehensive study of a group of Athapaskan porcupine quill specimens collected in 1862 which are held within the collections of National Museums Scotland, UK. Due to sampling limitations micro-destructive testing, or non-invasive analysis using PDA-UPLC, Raman Spectroscopy and PIXE were used to characterise the dye sources and metallic mordants. RBS was used to obtain additional information on the depth-profiling of the mordants in the keratin-based quill. The sensitivity of the quill specimens to photo-degradation was evaluated using Micro Fade Testing (MFT). The results from this multi-analytical study will be used to inform future display regimes of this unique collection.

1. Introduction

Non-European dyed materials other than textiles [1] have received little systematic study, particularly those made of porcupine quills, a material used by Native communities across North America and the Subarctic to decorate garments and basketry [2,3]. Since porcupine quills are made of a keratinous material similar to that of wool fibres [4,5], the dyeing processes might be expected to be related, but only limited information is available about the actual dye sources [6,7] and dyeing processes used by native North Americans [8]. Northern Athapaskan artefacts are particularly rare, but significant collections of these may be found at the Canadian Museum of Civilization and National Museums Scotland (NMS) [9,10]. It is known that quillwork decoration was an important element of Athapaskan clothing [11,12] and the Athapaskans used it to produce complex geometric coloured patterns [13]. Late 18th century European accounts of Athapaskan quillwork mention the use native species including Galium tinctorium L. Scop. and Helleborus trilfis L. to obtain red and yellow colours [8,14], while post 1850s Athapaskan work is characterised by the use of red, blue and white bands [13] but the dye sources for these are not well documented [8].

In this study, we analysed a unique group of dyed porcupine quills (Acc. N°: A.848.15) that were collected in 1862 from Northern Athapaskans (Fig. 1a) [9,15]. The range of colour is extraordinary and includes various shades of blue, green, orange, yellow, and red; together with very pale quills that could be either un-dyed or more faded than the rest of the materials. This range of colours reflects complex dying practices, with combinations of dyestuffs and even the use of additives (e.g., metallic mordants) to the dyebath.

The characterisation of the dye sources used in porcupine quillwork is particularly challenging, firstly due to sampling limitations, secondly as the dyestuff is only adsorbed on the very thin outer cuticle layer, giving a very small volume to be extracted for analysis (Fig. 18). To overcome these limitations, a combination of Photo Diode Array-Ultra Performance Liquid Chromatography (PDA-UPLC) and Raman Spectroscopy was used to characterise the dye sources either from micro samples, or completely non-invasively. This is the first application of PDA-UPLC to analyse quillwork, using a method recently developed for the study of historical textiles, [16] which provides a limit of...
2. Materials and methods

2.1. Dyeing processes and extraction

2.1.1. Reference specimens

Porcupine quills (Erethizon sp.) purchased from Native American artist Sarah Tronti were scoured and then dyed with either (i) cochineal (Dactylopius coccus C.) purchased from DBH Ltd Poole England, or (ii) turmeric (Curcuma longa L.) purchased from George Weil & Sons Ltd. Mordants, including alum, copper(II) sulfate, tin(II) chloride, iron(II) sulfate, chromium(IV) oxide, and cream of tartar were purchased from George Weil & Sons Ltd. Typically, dry quills (0.3 g) were dyed in a solution containing dyestuff (33 wt.%) and metallic mordants (25 wt.% alum, or 5–6 wt.% otherwise) for 1 h at 85–90 °C, as is traditionally reported for wool dyeing [24].

2.1.2. Extraction protocol for dye analysis

Reference and historical dyed porcupine quills (0.1–0.5 mg) were extracted with 37% HCl:H₂O:MeOH [200 μL, 2:1:1 (v/v/v)], at 100 °C for 10 min. The extract was centrifuged for 10 min at 10,000 rpm and then filtered directly into Waters UPLC vials® using a PTFE Phenomenex syringe filter (0.2 μm, 4 mm). The extract was then cooled with liquid nitrogen and dried under vacuum using a freeze drier system. The dry residue was then reconstituted with H₂O:MeOH [40 μL, 1:1 (v/v)] — allowing a single injection of 10 μL, see [16]. Samples containing turmeric were extracted with dimethyl sulfoxide (DMSO) [50 μL] at 100 °C for 60 min followed by filtration using a PTFE Phenomenex syringe filter (0.2 μm, 4 mm) allowing 10 μL injection.

2.1.3. Extraction protocol for ICP-OES analysis

A selection of historical dyed porcupine quills (1.0 ± 0.1 mg) were extracted with 37% HCl:H₂O:MeOH [200 μL, 2:1:1 (v/v/v)], at 100 °C for 10 min. The extract was filtered with a polyethylene filter (55 μm, 5 mm) from Crawford Scientific™. The frit was rinsed with methanol (200 μL) and the combined filtrates were then diluted using a solution of 37% HCl:H₂O [ca. 5 mL, 2:98 (v/v)] to give a final mass of 5.0000 g solution. The diluted extracts of the selected samples were analysed in triplicate and the mean concentration of Sn and Cu in each solution (per mg of quill) was calculated.

2.2. Dyestuff analysis

2.2.1. Reagents and calibration

All the standards were purchased from Sigma-Aldrich and ExtraSynth, France with a dye content ranging between 90 and 97%. Calibration was achieved using a selection of flavonoid and isoflavonoid dyes, see [16] and additionally: (a) a solution of carminic acid (1) (1.00 ± 0.01 mg) in H₂O:MeOH [25 mL, 1:1 (v/v); equivalent to 40 μg mL−1]; (b) a solution containing juglone (2) and curcumin (7) (0.50 ± 0.01 mg of each standard) in H₂O:MeOH [25 mL, 1:1 (v/v); equivalent to 20 μg mL−1]; (c) a solution containing alizarin (6) and purpurin (8) (0.20 ± 0.01 mg of each standard) in H₂O:MeOH [10 mL, 1:1 (v/v); equivalent to 20 μg mL−1]. Diluted solutions at concentrations of 20, 10, 5, 1, 0.5, 0.1, 0.05, 0.02 and 0.01 μg mL−1, were prepared using calibrated micro-pipettes.

2.2.2. Ultra performance liquid chromatography PDA-UPLC

A Waters Acquity UPLC® system was used with a Waters PDA detector (250 to 500 nm). Data were collected and processed by Waters Empower 2 software and Origin 8.5 (OriginLab, Northampton, MA, USA). The method used a PST BEH C18 (130 Å) reverse phase column, 1.7 μm particle size, 150 × 2.1 mm (length × i.d.), set-up with inline filter. The total run time was 37.33 min at a flow rate of 250 μL min−1 and the column was maintained at 55 °C. A binary solvent system, was used; A: 0.02% aqueous HCOOH (pH 3), B: MeOH. The elution program was isocratic for 3.33 min (77A: 23B) then a linear gradient from 3.33 min to 29.33 min (10A: 90B) before recovery of the initial conditions over 1 min and equilibration over 7 min. For details of the experimental
conditions used to extract reference and historical dyed porcupine quills, see [16].

2.2.3. Raman spectroscopy

Historical blue quills were further analysed with a Horiba XploRa Raman-Microscope equipped with three lasers working at 532, 638 and 785 nm. An edge filter prevents the laser wavelength entering the CCD detector. The 785 nm laser with 90 nW was used with its power reduced to 1% by filters, unless otherwise stated. A × 50 magnification objective was used to visualise the sample and to focus the laser on the object surface. A multi window acquisition mode was used; acquiring 3 × 2 spectral region for 600 s. Background subtraction and a smoothing filter were then applied to the spectra.

2.3. Mordant analysis

2.3.1. Particle induced X-ray emission and Rutherford backscattered spectrometry (PIXE/RBS)

Analyses were performed at the external beamline of the AGFA facility with a 3 MeV proton beam and a final analytical spot of 50 μm, two Si(Li) detectors for PIXE analysis (no filter for the low energy detector; 125 μm Be filter and 28.5 mm air for High Energy detector) and a PIPS charged particle detector in IBM geometry, with a scattering angle of 150° for RBS analysis. Measurements were undertaken in He-rich atmosphere. To reduce beam damage, a beam current of 3–6 pA was used and an area of 100 × 500 μm was scanned for an average of 4–5 min. Quantitative analysis was performed in trace mode using GUPIXWIN 2.1 [25] and TRAUPIXE [26] software, modelling a thick layer of keratin using the elements C, H, N, O, S in proportion to their reported relative wt% [27]. Because of the detector geometry and larger active area (30 mm²), the High Energy detector provided better statistical data, allowing the quantification of elements above 5 (Z = 16) present in the reference and historical porcupine quill specimens. Due to the lower statistical data obtained with the Low Energy Detector (10 nm²), the quantification of Na (Z = 11) and Al (Z = 13) are not discussed in this paper. A Dr-N certified standard was used to improve the quantification on the Low and High Energy detectors. RBS spectra were simulated with SIMNRA V6.5 software [28] and non-Rutherford reactions for the light elements in keratin (C, N, O) were added to the SIMNRA simulation [20].

2.3.2. Inductively coupled plasma optical emission spectroscopy (ICP-OES)

A selection of historical quills were analysed by ICP-OES using a Perkin Elmer Optima 5300 DV, employing an RF forward power of 1400 W, with argon gas flows, respectively. Using a peristaltic pump, sample solutions were taken up into a Gem Tip cross-Flow nebuliser based on the average value of the baseline noise (Hnoise) of several solutions. Calibration curves were obtained by pre-preparing standard solutions, one of copper and the other one tin, both at 1000 μg mL⁻¹. Calibration curves were prepared by preparing a range of concentrations with calibrated micro-pipettes, allowing the analysis of concentrations ranging between 200 and 0.02 μg mL⁻¹. Two wavelengths were selected: 327.393 nm for the analysis of copper(II) and 283.994 nm for the analysis of tin(II). Three replicates were run per sample.

2.4. Micro-fade testing (MFT)

An Oriel® Fading Test System (model 80190) was used with a modified probe head that was upgraded by an endoscope camera for better sample positioning. For irradiation of the sample under 0° (angle to the surface normal), a 75 W xenon arc lamp was used. The reflection spectra were taken at 45° and were recorded with a photodiode array detector (control development, model PDA-512). The light line is via fibre optics. The irradiated area has a diameter of ca. 0.4 mm. The intensity of light at the illuminated spot was approximately 3–4 Mlx. The reflected light was measured constantly with an integration time of 6 ms and 10 spectra were averaged. From these spectra colour values in the CIELAB colour space where calculated using an illuminant and observer combination of D65 and 2°, respectively. The colour difference to the initial (unfaded) measurement was given in colour difference value of CIE ΔE 2000 according to the recommendation of the International Commission on Illumination (Commission Internationale de l’Eclairage — CIE) [29].

The fading results were compared to ISO blue wool (BW) standard materials which have a known light fastness on a scale from BW 1 to BW 8, with BW 1 being the most light-sensitive. All samples were measured for 60 min in an environment of about 50–55% RH and 20–24 °C.

3. Results and discussion

Porcupine quills have a similar structure to other fibres, such as wool or hair, which are based on hard α-keratin [45]. X-ray diffraction studies have shown that porcupine quills have a highly crystalline structure, with the external layer of the quill (cuticle) being less ordered compared to the internal layer of the quill [30]. Although there are extensive studies on the properties of wool and hair fibres, or feathers, especially with regard to their uptake of dyes and metal ions [31–34], there are only a very limited number of studies on porcupine quills and their dyeing properties [6,7]. In contrast to wool fibres, where the dyestuff is evenly distributed through the fibre, generally only the thin cuticle layer of quills interacts with the dyebath (Fig. 1b) and adsorbs the dyestuffs and metal ions. Similarly to wool and hair fibres, this cuticle layer is reported to be slightly richer in sulphur [35] and can range from 30–100 μm depending on the size of the quills (ESI 3).

3.1. Dye analysis

Prior to the analysis of historical artefacts, a UPLC method for dyestuff identification in quillwork was developed, which would accommodate the limitations of sample size and volume. The UPLC method included dye sources which were anticipated to be found in these late 19th century specimens such as alizarin, purpurin, indigo and the tannin juglone, all characterised in Eastern Woodlands quillwork; [6,7] the dyes carminic acid and curcumin, which are included in the NMS purchase records for these specimens; [15] and the flavonoids luteolin, apigenin and genistein, which are present in a wide range of flavonoid dye sources [1,16,36].

For all the dye standards the average variation in retention time in 12 measurements by PDA-UPLC ranged between 0.04–0.06%. The limits of detection (LoD) and limits of quantification (LoQ) were calculated based on the average value of the baseline noise (Hnoise) of several solvent blanks, considering all data points. The baseline of the UV detector at 254 nm averaged (9 ± 1) × 10⁻⁶ AU, resulting in detection limits ranging from 0.5 ng for genistein (4) to 5.1 ng for purpurin (8) for an injection volume of 5 μL, while the LoD of curcumin (7) was calculated at 450 nm and averaged 4.0 ng. (Table 1)

Table 1

| Component   | Rₚ (min) | λ (nm) | LoD (ng) | λmax (MeOH:H2O) (nm) |
|-------------|----------|--------|----------|----------------------|
| Carminic acid (1) | 5.80 | 254 | 1.9 [5.9] | 276, 312 (sh), 493 |
| Juglone (2) | 12.10 | 254 | 2.2 [6.6] | 249, 330 (sh), 408 (sh), 424 |
| Luteolin (3) | 13.47 | 254 | 0.9 [2.6] | 252, 291 (sh), 349 |
| Genistein (4) | 14.03 | 254 | 0.5 [1.6] | 260, 332 (sh) |
| Apigenin (5) | 15.26 | 254 | 1.9 [5] | 267, 300 (sh), 338 |
| Alizarin (6) | 17.70 | 254 | 0.8 [2.3] | 230 (sh), 248, 280, 433 |
| Curcumin (7) | 20.41 | 450 | 4.0 [12.1] | 264, 429 |
| Purpurin (8) | 20.77 | 254 | 5.1 [15.3] | 256, 296, 456, 481, 515 (sh) |
A reference quill dyed with American cochineal (Dactylopius coccus Costa) and 18 red and orange coloured Athapaskan porcupine quills were acid hydrolysed for PDA-UPLC analysis. In all these samples carminic acid (1) was found associated with the dye components dcII, dcIV, dcVII and flavokermesic acid that are known to occur in cochineal species from South America [36–39]. An unknown red dye component (Fig. 2a) was additionally observed in most of the historical samples. A complementary Mass Spectrometric study would be necessary to suggest a structure for this component, but UV–Vis spectra indicate that it is related to flavokermesic acid. (ESI 1) The relative amounts of dcI and carminic acid were found to be quite variable in the historical samples, which could possibly relate to dyeing practices, as has been observed in the over-dyeing of textiles with flavonoid dye sources [16]. A reference quill dyed with turmeric (Curcuma longa L.) and 12 yellow, orange and green coloured Athapaskan porcupine quills were extracted with DMSO. The presence of bisdesmethoxycurcumin (7b), desmethoxycurcumin (7a) and curcumin (7) in these samples was characterised by PDA-UPLC (Fig. 2a and ESI 2); as with the acidic extraction conditions (HCl:H2O:MeOH), carminic acid (1) was also found in the orange samples. In all these samples bisdesmethoxycurcumin (7b) was observed to be the main dye component extracted. This varied from data published for turmeric rhizomes from India and China, [40, 41] which exhibit higher levels of curcumin (7). The differences observed could reflect a higher affinity of bisdesmethoxycurcumin for the quill substrate (Fig. 2c). The brighter shades of green and blue were obtained by an over-dyeing process with the sulfonated indigo carmine dye which was evidenced by Raman Spectroscopy (Fig. 2b).

The exclusive use of American cochineal instead of madder species, and of indigo carmine instead of indigo is in sharp contrast to results obtained in previous studies on pre-1850s Eastern Woodlands quillwork [6]. The semi-synthetic dye indigo carmine would have been traded from Europe, where it was available and used for textile dyeing from 1770 until the beginning of the 20th century [42]. Since indigo carmine does not require a vat dyeing process it would have been easier to use than indigo, allowing the production of more intense shades of blue. The highly variable diarylheptanoid content [40,41] combined with the unknown affinity of the dyestuff components for the quill substrate, meant that it was not possible to attribute the turmeric compositions to a specific source (ESI 2), although it is likely that the turmeric originated from India [1,24] and reached the Athapaskans via trade into Europe.

3.2. Metallic mordants

Applications of PIXE to organic based materials are limited and thus it was necessary to evaluate the thickness of keratin being analysed and the limit of detection (LoD) of various metallic mordants to be detected. The range of the beam in the matrix of keratin was determined as 136 μm using TRIM software (SRIM 2003 version), [44] and the effective depth values for S, K, Ca, Cr, Fe, Cu, Zn, Sn, As and Hg were calculated with GUCSA program in the GUPIXWIN 2.1 software package and vary

![Fig. 2.](image-url)
between a minimum of 52 μm for S (Kα) and a maximum of 80 μm for Hg (Lα) (Table 2). Since the average thickness of the cuticle was determined as 58 μm by SEM analysis (ESI 3), these effective depth values represent the contribution of both the cuticle and the cortex. The LoD of each element was averaged from 70 measurements using GUPIXWIN 2.1 software and ranged between 3 ppm for Fe (Kα), 110 ppm for the Sn (Lα) and 520 ppm for the Sn (Kα) lines. The quantification of Sn was therefore undertaken using the Sn(Lα) line, which presented lower LoD and better statistical data. The unfolding of Sn(Lα) and K(Kα) was achieved by GUPIXWIN 2.1 software.

A small set of modern quills prepared with dyebaths containing combinations of cream of tartar, alum, Cr, Fe, Cu and Sn were investigated by PIXE. Traces of Zn were detected in all the modern quills, including a scoured one, at an average concentration of 50 ppm, possibly corresponding to contamination. The levels of S were found to range between 2 and 4 wt.% as would be expected in keratin. K levels were found to be significantly increased (2000 to 4000 ppm) for the quills where cream of tartar was added to the dyebath, while traces of Cr, Fe, Cu and Sn were detected at concentrations ranging from ~100 ppm to ~1000 ppm, allowing differentiation between the individual dyebath processes.

Around 70 historical quills were analysed, covering all the different colours observed in the specimens (Fig 3a). The elemental data for a selection of quills is presented in Table 3. The S level was generally found to range between 1.5 and 2.5 wt.%. A few green and blue quills exhibit up to 4.0 wt.% of S (Fig. 3b), which might be related to the presence of the sulfonated indigo carmine dye or to the use of sulfur-containing mordants such as alum. Traces of Zn were also identified at a concentration below 100 ppm.

For the coloured historical quills, Fe, Cu and Sn were identified in varying amounts, while levels of K were found to be higher in blue, yellow and green hues reaching 4000 ppm, with levels as high as 8000 ppm for two of the darker green quills (Fig. 3b). These results are in sharp contrast to earlier Eastern Woodlands quillwork, where XRF analysis revealed the presence of only Fe (and sometimes Cu) residues in a few objects [7]. In the Athapascan samples, Cu was found predominantly in the darker shades of blue and green to a maximum concentration of 6000 ppm, while Fe was generally below 200 ppm with a few exceptions where it reached ~1000 ppm (Fig. 3c & d). The level of Sn was found to be significantly higher in the red and oranges hues and one bright blue quill, ranging from 4000 to 15000 ppm (Fig. 3d). These levels are significantly higher than the levels observed in modern quills prepared with 5–6 wt.% of mordant. In order to confirm these ranges of concentrations, a selection of samples were additionally analysed by ICP-OES, a technique routinely used for the characterisation of metals in textile materials [46]. The values obtained cannot be directly compared to PIXE values, as surface heterogeneity is an important consideration and a standard mass of quills (1 mg) was extracted for ICP analysis. However, the values showed some correlation for both Cu and Sn analysis, with the level of Sn reaching 2.9 × 10^4 μg g⁻¹ (or ppm) for the most concentrated samples.

**Table 2**

Effective depth values calculated with the GUCSA program of the GUPIXWIN 2.1 software package [25], representing the thickness in μm from which 95% of the detected X-rays are produced and associated limit of detection (LoD) calculated with GUPIXWIN for each element analysed in the quills, modelling a thick target of keratin [27,45]. PIXE conditions: 3 MeV protons, 125 μm Be filter and 28.5 mm air for High Energy Sn(L) detector.

| Element | Effective depth (μm) | LoD (ppm) |
|---------|---------------------|-----------|
| S       | 52                  | 32        |
| K       | 68                  | 23        |
| Ca      | 73                  | 31        |
| Cr      | 79                  | 6         |
| Fe      | 79                  | 3         |
| Cu      | 79                  | 5         |
| Zn      | 72                  | 5         |
| Sn      | 70                  | 520       |
| Hg      | 80                  | 110       |
| As      | 77                  | 13        |
| S       | 78                  | 6         |
| K       | 79                  | 5         |
| Ca      | 79                  | 3         |
| Cr      | 79                  | 5         |
| Fe      | 79                  | 5         |
| Cu      | 72                  | 5         |
| Zn      | 70                  | 520       |
| Sn      | 80                  | 110       |
| Hg      | 77                  | 13        |
| As      | 6                   | 5         |

**Table 3**

PIXE elemental data obtained with High Energy detector for a selection of modern quills and historical quills. In addition, the concentrations of Cu(II) and Sn(II) expressed in μg g⁻¹ (or ppm) obtained by ICP-OES are presented for comparison. PIXE analysis entries marked “-” correspond to a level below LoD; ICP-OES entries marked “-” were not analysed.

| Entry | PIXE | ICP-OES |
|-------|------|---------|
|       | S (wt%) | K (ppm) | Ca (ppm) | Cr (ppm) | Fe (ppm) | Zn (ppm) | As (ppm) | Hg (ppm) | Sn (ppm) | Cu (ppm) | Sn(II) (μg g⁻¹) | Cu(II) (μg g⁻¹) |
| Modern reference quills | | | | | | | | | | | | |
| Scoured quill | 2.8 | 350 | – | 10 | 68 | – | – | – | – | – | – | – |
| Al, K, Cr | 1.6 | 3944 | 379 | 1280 | 8 | 87 | – | – | – | – | 5 | – |
| Al, K, Sn | 2.3 | 4173 | 702 | 8 | 55 | – | – | – | – | 480 | 7 | – |
| Cu | 3.1 | 380 | 111 | – | – | 32 | – | – | – | – | 224 | – |
| K, Fe | 4.0 | 2460 | 109 | – | 101 | 18 | – | – | – | – | – | – |
| Athapascan quills (A.848.15) | | | | | | | | | | | | |
| B1 bright blue | 1.8 | 1335 | 458 | – | 39 | 25 | – | – | – | 592 | 859 | – |
| B1 spotted blue | 3.4 | 1138 | 140 | – | 66 | – | – | – | 122 | 580 | 404 | – |
| B2 red (1) | 1.6 | 69 | 125 | – | 688 | 16 | – | 7133 | 980 | – | – | 1.7 × 10⁴ | 3.0 × 10⁴ |
| B2 red (2) | 1.3 | 160 | 220 | – | 70 | 45 | – | – | 14000 | 200 | – | – | 2.9 × 10⁴ | 2.0 × 10⁴ |
| B1 light green | 1.3 | 305 | 230 | – | 115 | 31 | – | – | 1791 | 252 | – | – | – |
| B4 yellow (1) | 1.4 | 2040 | 480 | – | 815 | 37 | – | – | 826 | 1725 | – | – | 1.4 × 10⁴ | 3.0 × 10⁴ |
| B6 red (1) | 1.8 | 509 | 546 | – | 71 | 17 | – | – | 2492 | 288 | – | – | 9.0 × 10³ | 2.0 × 10³ |
| B6 red (2) | 1.7 | 254 | 653 | – | 68 | 41 | 10 | – | 42 | 1801 | 380 | – | 1.0 × 10⁴ | 1.0 × 10⁴ |
| B8 orange (1) | 1.6 | 365 | 137 | – | 59 | 52 | – | – | – | 4315 | 250 | – | 1.1 × 10⁴ | 2.0 × 10⁴ |
| B8 orange (2) | 1.3 | 230 | 224 | – | 69 | 40 | – | 71 | 7965 | 239 | – | 1.4 × 10⁴ | 1.0 × 10⁴ |
| B9A bright blue | 2.2 | 1388 | 508 | – | 222 | 16 | – | – | – | 1012 | 323 | – | 2.0 × 10³ | 4.0 × 10² |
| B11A colourless | 2.0 | 772 | 261 | – | 62 | 32 | – | – | – | 537 | 958 | – | – |
| B12A colourless | 1.9 | 419 | 372 | – | 90 | 61 | – | 30 | 990 | 432 | – | 5.0 × 10³ | 1.0 × 10³ |
| B17A dark green | 2.8 | 6779 | 1406 | – | 358 | 100 | – | 78 | 563 | 1551 | – | 5.0 × 10⁴ | 5.0 × 10³ |
| B17B dark green | 2.2 | 8038 | 942 | – | 444 | 150 | – | 171 | 14500 | 5780 | – | 1.2 × 10⁴ | 5.0 × 10³ |
The analysis of “colourless” quills also exhibited traces of Sn and Cu suggesting that either these quills were pre-mordanted or that these were previously dyed and are now extremely faded. Finally, As and Hg were found in only a few quills, with most analyses falling below the LoD, indicating that, in contrast to many organic collections in museums of this period, these specimens were unlikely to have been treated in the past with inorganic pesticides containing mercury and arsenic [47,48].

The quills were simultaneously analysed by RBS which provided additional information on the depth-profiling of the mordant in the cuticle layer. The RBS spectra were treated independently to the PIXE data. For the majority of the samples (modern and historical) a two-layered model was used for simulation with SIMNRA software comprising a very thin layer of keratin rich in S with traces of Sn and Cu (Layer 1) over an infinite layer of keratin (Layer 2) (Fig. 4a). The thickness of Layer 1 was evaluated to $80,000 \times 10^{15}$ Atoms cm$^{-2}$ which is equivalent to 9 $\mu$m considering the atomic composition and density of the keratin (ESI 4). This model showed a good correlation for the RBS spectra obtained from scoured (Fig. 4b) and mordanted modern quills, as well as dyed Athapaskan quills (Fig. 4c).

For all the samples, the composition of Layers 1 & 2 was found to closely match the expected atomic concentrations for keratin with the addition of traces of mordant (ESI 4). In all the samples analysed, the level of S and mordant was found to be higher in Layer 1 (4 to 5 wt.% for S) than the average values determined by PIXE (1.5 to 2.5 wt.% for S), see ESI 4. While a higher level of S is expected in the outer cuticle, [35] the higher levels of mordants could possibly correspond to a deposition on the surface of the cuticle. Finally, for a few blue and green quills a three-layered model was used (Fig. 4d), comprising of a very thin layer of keratin rich in S with traces of Sn and Cu (Layer 1, 1–2 $\mu$m), a second layer with a lower level of S and traces of Cu only (Layer 2, 3 $\mu$m) over an infinite layer of keratin (Layer 3), see ESI 4. These samples showed a clear increase in S level at the surface (up to 13 wt.% in Layer 1), confirming the observations made by PIXE and supporting the hypothesis that there is either indigo carmine dye or the deposition of a sulfur-containing mordant on the surface of the cuticle. The presence of Sn in Layer 1 only might reflect the use of multiple dyebaths with different mordants combinations, resulting in the formation of successive thin layers at the surface of the quill.

3.3. Micro-fade testing

A selection of coloured quills were tested for their photo-sensitivity by Micro-Fade Testing (MFT), as shown in Fig. 5a and Table 4. The MFT determines the photosensitivity of the measured spot as a colour change as perceived by the human eye (and not as a change of colourant concentration). The photo-sensitivity not only depends on the colourant and supporting materials but can also be influenced by mordant, surface particularities, chemical environment (air pollutants, oxygen level humidity) and colourant concentration. As a result MFT measurements are variable and require multiple repeats to obtain reproducible results which may be accurately interpreted. Table 4 summarises the MFT measurements; the results of one sample, B1 Spotted blue, have been omitted since the repeat variability was unusually high, probably due to the heterogeneity of the surface which was already visually observable by the speckled colour effect of the quill.

It can be seen that, with the exception of B2 red (2), all the quills fall into (or very close to) the Blue Wool class 1 to 3. Materials of this class are considered to be highly responsive to light following the categories given in CEN/TS 16163:2014 [49] and CIE 157:2004 [50]. It is advised that such materials should be illuminated at a maximum recommended-level of 50 lx and only for a limited time span not exceeding 15,000 lx h in a year.

It was observed that the concentration or type of mordant did not affect the MFT results as much as the type of dyestuffs used. As expected, indigo carmine and turmeric coloured quills [B1 blue, B3 blue, B4 yellow (1)] show the highest photo-sensitivity due to the low colourfastness of these colourants [22]. In mixtures the colours tend to be slightly more...
stable as can be seen for samples B3 light green and B8 orange (1). In B8 orange (1), the turmeric is mixed with the more stable red American cochineal which results in a colour that falls in the BW 3 class and may be categorised as a medium light responsive material. The photosensitivity of quill B8 orange (1) therefore lies between quills dyed with one of these colourants, i.e., American cochineal [B2 red (2) and B6 red (1)] and turmeric [B4 yellow (1)]. As the American cochineal is less responsive, a colour shift from orange to red results from the MFT process as indicated in Fig. 5b because the more responsive yellow turmeric fades first. The reason for the smaller colour change of the mixed dye is probably not a stabilising effect of the American cochineal on the turmeric, but the result of a smaller perceptual colour change due to the persisting colour of the American cochineal.

4. Conclusion

This study of Athapaskan quillwork, using a combination of techniques, has provided important new information on the dyeing processes used in the Subarctic in the 1860s. Firstly PDA-UPLC was successfully applied to micro-samples of porcupine quillwork allowing the dye...
sources to be characterised. This revealed the unexpected use of imported European dyes in the Athapaskan quillwork, in sharp contrast to the local dye sources found in Eastern Woodlands quillwork pre-dating 1856 [6,7]. Secondly the combined use of PIXE and RBS allowed the concentrations of various mordants (K, Ca, Fe, Cu and Sn) in the quills to be determined, providing an alternative non-destructive methodology for the study of keratin-based materials. RBS was applied for the first time to the investigation of quillwork and it allowed additional information on the depth-profiling of mordants at the surface of the quills to be obtained. Several quills presented a more complex layered system when analysed by RBS, which might be related to the use of multiple dyebaths with different mordant combinations, resulting in the formation of successive thin layers containing metallic residues at the surface of the quill. These results present a promising opportunity for the study of quillwork and related keratinous materials, and would need to be further investigated by analysing modern quills prepared with successive dyebaths.

In all the Athapaskan specimens investigated, two to three metallic mordants (Fe, Cu, Sn) were found to be combined, and by changing the concentrations of these mordants brighter and darker hues were obtained from the same dyestuff combination. The high levels of metallic mordants observed in several of the specimens suggest that the effect of the mordants was appreciated and that they were deliberately added to the dyebath to change the property of the dyestuffs. This transfer of European dyeing practices might not be surprising given that there is evidence that European dyes and mordants were traded to Nova Scotia from the 1830s, [51] suggesting that by 1862 these materials would also have been available in the Northern territories. Finally, the data obtained from the MFT confirmed previous findings which document the photosensitivity of Subarctic Athapaskan artefacts [21]. We expect that the combination of techniques presented in this paper will be adopted for the study of quillwork and related collections.

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Appendix A. Supplementary data

Electronic Supplementary Information (ESI) available: PDA-UPLC analysis of red and orange quills; PDA-UPLC analysis of yellow and green quills; natural dyes; SEM-EDX analysis; Rutherford Backscattering Spectrometry. Supplementary data associated with this article can be found in the online version, at http://dx.doi.org/10.1016/j.microc.2015.11.053.

Primary data files for this work are available at the Edinburgh DataShare.

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