

Contents lists available at ScienceDirect

Dyes and Pigments



journal homepage: www.elsevier.com/locate/dyepig

Shedding light on the composition and degradation mechanism of dyes in historical ink's collection (19th-20th century)

Adele Ferretti^a, Ilaria Degano^{a,b,*}, Stefano Legnaioli^c, Beatrice Campanella^c, Aurora Sainati^a, Maria Perla Colombini^a

^a Dipartimento di Chimica e Chimica Industriale, Università di Pisa, Via G. Moruzzi 13, 56124, Pisa, Italy

^b Center for Instrument Sharing of the University of Pisa (CISUP), University of Pisa, Italy

^c Consiglio Nazionale Delle Ricerche (CNR), Istituto di Chimica Dei Composti Organometallici, SS di Pisa, Via Moruzzi 1, 56124, Pisa, Italy

ARTICLE INFO

Keywords: Historical red inks Raman spectroscopy TLC-SERS HPLC-ESI-Q-TOF Synthetic organic pigments

ABSTRACT

Man has always used writing to be able to communicate, express, and disseminate his thoughts. In time, many different coloured extracts of plant and animal origin have been used to produce inks; after the development of synthetic chemistry, artificial and synthetic dyes started to be widely exploited. The end of the 19th century marked great technological and industrial innovations in commercial production of artists' materials. To reveal ink formulations and build a database of red inks by different producers, we developed a multi-analytical approach and investigated a collection of writing inks produced in France in the late 19^{th} – early 20^{th} century. The materials used as binders, additives, dyes, and pigments have been investigated by high performance liquid chromatography coupled with diode array and tandem mass spectrometric detectors (HPLC-DAD-MS²), in situ derivatisation pyrolysis coupled with gas chromatography-mass spectrometry (Py(HMDS)-GC/MS), surface enhanced Raman (SERS, TLC-SERS) and X-Ray Fluorescence spectroscopies. Several dyes and pigments were detected, showing that the French ink's formulations of the early days of synthetic dye industry were based on rhodamine B and 6G, eosin Y, rose Bengal and methyl or crystal violet. Instead, as binder and additives only gum Arabic and shellac resin have been identified, respectively. Mass spectrometry also allowed us to detect possible by-products of the synthesis of ink's dyes and even early degradation products, that can be used for ink identification in historical writings and drawings. Our studies can pave the way to investigate inks in historical samples by introducing ultra-sensitive chromatographic and mass spectrometric methods in the array of analytical tools available to the chemist.

1. Introduction

The investigation of historical and archaeological artefacts, such as pottery, writing instruments, inks, and textiles is influential in improving society's understanding of the past, present and future [1,2]. The technique and the art of writing and drawing has progressed over the centuries, along with the supports and materials necessary for its expression [3]. Although the composition of inks may differ based on the desired colour and specific products, they are all composed of three main components [4,5]: the *liquid vehicle*, or the ink's portion which suspends and transports the dye to the paper support; *colourants*, that impart a well-defined colour to the writing material; *additives*, or rather substances that are specifically added to enhance writing performances. The early development of inks involved the use of natural or artificial

pigments [6–8], such as *lampblack* [9], used to prepare carbon-based inks, or organic pigments like *tannins, hematein* and *sepia*, employed in iron gall [10,11], logwood [12] and sepia ink [13] formulations, respectively. The 19th century was a period of great technological and industrial innovations in this field. The increase in the global literacy rate caused a dramatic change in the ink manufacture, leading to industrial production. Furthermore, in 1856 Perkin discovered Mauveine [14,15], marking the beginning of the synthetic organic pigment (SOPs) industry. In this context, ink formulations began to change. Thus, if in the 19th century both natural and synthetic pigments played a fundamental role in the production of inks, starting from the early 20th century the use of SOPs became a trend in art and in ink formulations [16]. SOPs allowed a wide range of different colours to be obtained in a simpler and more cost effective way, and writing materials to perform

https://doi.org/10.1016/j.dyepig.2023.111672

Received 14 April 2023; Received in revised form 31 August 2023; Accepted 1 September 2023 Available online 4 September 2023

^{*} Corresponding author. Dipartimento di Chimica e Chimica Industriale, Università di Pisa, Via G. Moruzzi 13, 56124, Pisa, Italy. *E-mail address: ilaria.degano@unipi.it* (I. Degano).

^{0143-7208/© 2023} The Authors. Published by Elsevier Ltd. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

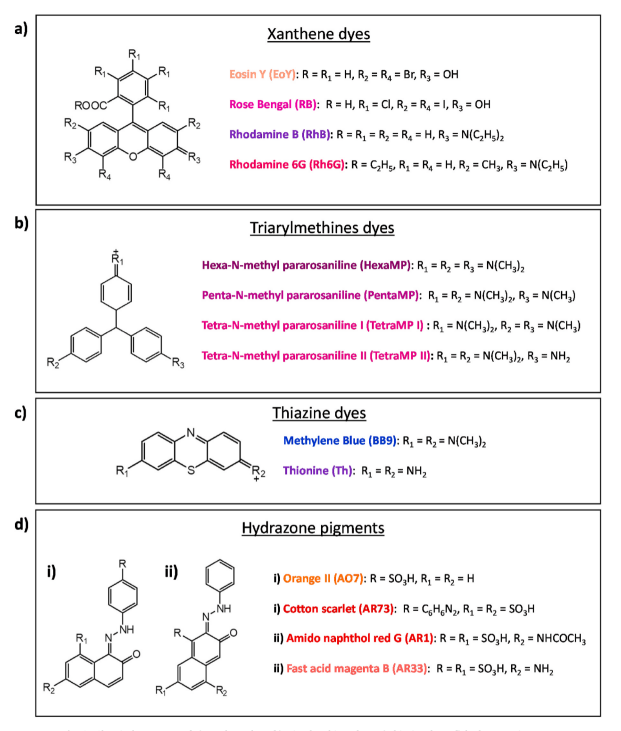


Fig. 1. Chemical structures of a) xanthene dyes; b) triarylmethines dyes; c) thiazine dyes; d) hydrazone pigments.

better than inks prepared from natural dyes, opening up a new and fascinating landscape for ink's manufactures. Among the most important classes of SOPs introduced on the market in the late 19th century, *triarylmethines, xanthenes, thiazine* and *hydrazone dyes* (Fig. 1) were widespread in ink's recipes. Triarylmethines dyes are nowadays known as the most widely used early synthetic dyes. All the SOPs belonging to this class are characterized by bright colours, given by three conjugated aromatic rings linked to a central atom of carbon and various aux-ochromic substituents [17]. Thanks to their versatility, triarylmethines dyes have been widely used up to now in textiles, artworks, ballpoint pens and writing inks [18–20]. Xanthenes dyes, obtained through an intramolecular cyclization of triarylmethines bridged with an oxygen atom [17], are fluorescent dyes that have been very attractive for their brilliant hues [21,22]. Consequently, xanthenes have been broadly exploited in the artistic field, for the printing industry, but also in the cosmetic, pharmaceutical and food fields [18,23,24]. However, both of these classes exhibit a strong tendency to fade upon light exposure, and the photostability of some representative xanthene dyes has been discussed in the literature [18,25,26]. Hydrazone pigments, characterized by 1-arylhydrazone-2-naphthol system [27], constitute the largest group of synthetic organic pigments ever produced [28]. On account of their brilliant shades, excellent lightfastness and high tinting power, these compounds have been employed for several applications, such as leather, plastic, medicine, food and ink's industry [28–30]. Thiazine

A. Ferretti et al.

Table 1

Historical ink's collection from $19^{\text{th}}-20^{\text{th}}$ century. The following information is reported for each ink: sample code used during the analysis (samples code), name of the ink and/or manufacturer reported on the bottle labels (label), photo of the ink bottles (photo), and colours of the ink mock-ups obtained in accordance with section 3.3 (colour).

| Samples code | Label | Photo | Colour |
|--------------|---|---------------------------|---------------------------------------|
| 1 | Encre Écarlate; New Red Ink; Spécialement préparée pour les plumes métalliques; N. Antoine & Fils; Paris | | |
| 2 | Encre Rouge Écarlate; Edouard Devillers; Paris | | |
| 3 | Encre a tampon; sans huile; rouge; cette encre sèche rapidement et donne de très belles empreintes; Bagnolet Paris. | | Ink's 3 bottle was found empty. |
| 4 | Vermillon; Indélébile; Yang Tse; J.M. Paillard; Paris | | |
| 5 | Olympic; Encre De Chine | DLY MPIC | |
| 6 | Bayard; Encre Supérieure | BAYARD SUDERIEURE | |
| 7 | Manufacture Parisienne; Au Chat Noir; Encre Rouge; Paris | | |
| 8 | Vermillon; Indélébile; Yang Tse; J.M. Paillard; Paris | | Ink 8 was a replica of ink 4. |
| 9 | Encre a Tampon; Tiflex; Supérieure; Sans Huile | Tories Tarico Tribe | |
| 10 | Encre Écarlate; New Red Ink; Spécialement préparée pour les plumes métalliques; Encre Antoine; Paris | | |
| 11 | Encre Rouge Supérieure | CHICRE RADIE | |
| 12 | Encre De Couleur; Parfumée; Carminée; Médailles D'Or; Paris 1889 & 1900; Adrien Maurin Paris | | |

Dyes and Pigments 220 (2023) 111672

| Samples code | Label | Photo | Colour |
|-----------------|---|--------------------------|--------|
| 13 | Papeterie Generale Des Ecoles; Encre Couleur Superieure; Boudignon Sartiaux; Paris | | |
| 14 | Encre Gauloise; Carminée Supérieure; V. Chantraine Paris | | |
| 15 | Encre Modèle Rouge; R. Barth | CICRE MODEU P. BAIRTH | |
| 16 | Pourpre; Couleurs Liquides Indélébiles; Tien-Tsin; Encres Antoine | TOMP | |

Table 1 (continued)

dyes are tricyclic and heterocyclic compounds with one atom of sulphur and one nitrogen in the central ring [31]. The high luminescence and photoconductivity of these SOPs have promoted their spreading as dyes in paper, writing inks, lake-pigments, textiles, or in more recent applications for microscope cell staining, pharmaceuticals, and photo galvanic cells [31–34].

From an analytical point of view, the identification of inks in drawing or manuscript samples is challenging, to a large extent due to the numerous recipes used over the centuries. Furthermore, the ageing and fading processes promote a variation in the chemical profile of the inks, providing a different molecular composition over time compared to the original one. This phenomenon, reported in the literature for crystal violet-based inks [25] and for a model iron gall ink [35], is particularly evident when SOPs, amongst the most fragile materials in works of art and documents, are the main dye component of the ink's formulations. Hence, the analysis of historical artistic materials is of utmost importance to improve our understanding of manufacturing techniques, as already showed in the literature for the study of tube paints, felt-tip pens, crayons, historical pigments and ink's collection [16,18,36]. At the same time, the study of historical collections could allow to detect dyes/pigments degradation products, enabling heritage science to gain information on synthetic pathways and degradation processes, and help planning sustainable conservation strategies. Several techniques have been exploited for the detection of synthetics organic pigments in the literature, such as Raman or surface enhanced Raman spectroscopy (SERS) [37-39], thin layer chromatography (TLC) coupled to SERS analysis [40,41], pyrolysis coupled to gas chromatography - mass spectrometry [18,20] and capillary electrophoresis [42]. In the last decade a general consensus identified high performance liquid chromatography (HPLC) coupled to several detectors as the technique par excellence for the analysis of SOPs and their by-products [15,18,43-45] since its high sensitivity and selectivity allow both the determination of trace compounds and of mixture of dyes.

2. Research aim

This work aims to reveal trends in historical ink formulations, to develop a multi-analytical approach able to fully characterise the ink composition, and to highlight degradation markers enabling the identification of the colourants in aged and possibly faded historical ink samples. Thus, a multi-analytical approach was developed for the

Table 2

Summary of the results of the analyses of French inks (if the results did not significantly differ from the analytical blank, the corresponding cell is marked with an x).

| | Ink | SERS | TLC- SERS | HPLC-DAD | HPLC | -ESI-Q-ToF | Py-GC/ MS | Summary | |
|--|-----|--------------|----------------------|---|------------------|--|---------------------------|------------------------------|--|
| Methyl or crystal violet- based inks | 1 | Mv or Cv | AR73 | AR73 | (+) (-) | Cv AR73 | X | Dyes Binder, Additives | Cotton scarlet (AR73), crystal violet (Cv x |
| | 6 | Mv or Cv | x | AR33, AR1 | (+) | Mv or degraded Cv, BB9, BB9–Me | Arabic gum (?), AR1 | Dyes | Amido naphthol red G (AR1), fast acid magenta B (AR33), methylene blue (BB9) methylene blue photo-oxidation product (BB9–Me), degraded crystal violet (Cv) o methyl violet (Mv) |
| | | | | | (–) | AR33, AR1 | | Binder, Additives | Arabic gum? |
| | 9 | Mv or Cv | AR73 | AR73 | (+) (-) | Cv AR73 | AR73 | Dyes Binder, Additives | Cotton scarlet (AR73), crystal violet (Cv x |
| | 14 | Mv or Cv | AR73 | AR73 | (+) (-) | Cv AR73 | x | Dyes Binder, Additives | Cotton scarlet (AR73), crystal violet (Cv x |
| Eosin Y or rose Bengal- based | 2 | EoY | EoY | EoY, DBEo | (+) | x | х | Dyes | Eosin Y, eosin Y by-products (DBEo, pseudo EoY) |
| inks | | | | | (–) | EoY, DBEo, pseudo EoY | | Binder, Additives | x |
| | 7 | EoY | EoY | EoY, DBEo | (+) | x | x | Dyes | Eosin Y, eosin Y by-products (DBEo, pseudo EoY) |
| | 11 | EoY | EoY | EoY, DBEo | (-) (+) | EoY, DBEo, pseudo EoY x | ЕоҮ | Binder, Additives Dyes | x Eosin Y, eosin Y by-products (DBEo, |
| | 11 | LOI | LUI | E01, DDE0 | (+) | A EoY, DBEo, pseudo EoY | LUI | Binder, | pseudo EoY) x |
| | 12 | RB | RB | EoY, DBEo | (+) | x | x | Additives Dyes | Eosin Y (EoY), eosin Y by-products (DBE |
| | | | | , | | | | _ , | 2DBEo, pseudo EoY), rose Bengal (RB), rose Bengal by-products or degradation markers (DCRB, 2DCRB) |
| | | | | | (–) | EoY, DBEo, 2DBEo, pseudo EoY, RB, DCRB, 2DCRB | | Binder, Additives | x |
| | 13 | ЕоҮ | EoY, RB | EoY, DBEo, AR33, AR1 | (+) | x | ЕоҮ | Dyes | Eosin Y (EoY), eosin Y by-products (DBE 2DBEo, pseudo EoY), rose Bengal (RB), rose Bengal by-product or degradation marker (DIRB), fast acid magenta B (AR33), amido naphthol red G (AR1) |
| | | | | | (–) | EoY, DBEo, 2DBEo, pseudo EoY, RB, DIRB, AR33, AR1 | | Binder, Additives | x |
| | 15 | EoY | EoY | EoY, DBEo | (+) | x | ЕоҮ | Dyes | Eosin Y (EoY), eosin Y by-products (DBE 2DBEo, pseudo EoY) |
| | | | | | (–) | EoY, DBEo, 2DBEo, pseudo EoY | | Binder, Additives | x |
| Rhodamine- based inks | 4 | RhB, Rh6G | RhB, Rh6G | RhB, Rh6G, DERhB, bisDERhB | (+) | RhB, Rh6G, DERhB, DERh6G, bisDERhB, bisDERh6G, pseudo RhB | Shellac, RhB | Dyes | Rhodamine B (RhB), rhodamine 6G (Rh6G), rhodamine photo-oxidation products (DERhB, DERh6G, bisDERhB, bisDERh6G), rhodamine B by-product (pseudo RhB) |
| | | | | | (–) | Shellac | | Binder, Additives | Shellac resin |
| | 5 | RhB | RhB | RhB, AO7, DERhB, bisDERhB | (+) | RhB, DERhB, bisDERhB, pseudo RhB | Shellac, RhB, AO7 | Dyes | Rhodamine B (RhB), rhodamine B photo oxidation products (DERhB, bisDERhB), rhodamine B by-product (pseudo RhB), orange II (AO7), orange 2R (AO8) |
| | | | | | (–) | Shellac, AO7, AO8 (?) | | Binder, Additives | Shellac resin |
| | 10 | Rh6G | RhB, Rh6G, EoY | RhB, Rh6G, EoY, AR1, AR33, DERhB, bisDERhB | (+) | RhB, Rh6G, DERhB, DERh6G, bisDERhB, bisDERh6G, trisDERhB, trisDERh6G, Rh, | x | Dyes | Rhodamine B (RhB), rhodamine 6G (Rh6G), rhodamine photo-oxidation products (DERhB, DERh6G, bisDERh6G, Rh eosin Y (EoY), eosin Y by-products (DBE 2DBEo, pseudo EoY), fast acid magenta (AR33), amido naphthol red G (AR1) |
| | | | | | (–) | EoY, AR1, AR33 | | Binder, Additives | x |
| | 16 | RhB | RhB | RhB, DERhB, bisDERhB | (+) | RhB, DERhB, bisDERhB, pseudo RhB | Shellac, RhB | Dyes | Rhodamine B (RhB), rhodamine B photo oxidation products (DERhB, bisDERhB), rhodamine B by-product (pseudo RhB) |
| | | | | | (–) | Shellac | | Binder, Additives | Shellac resin |

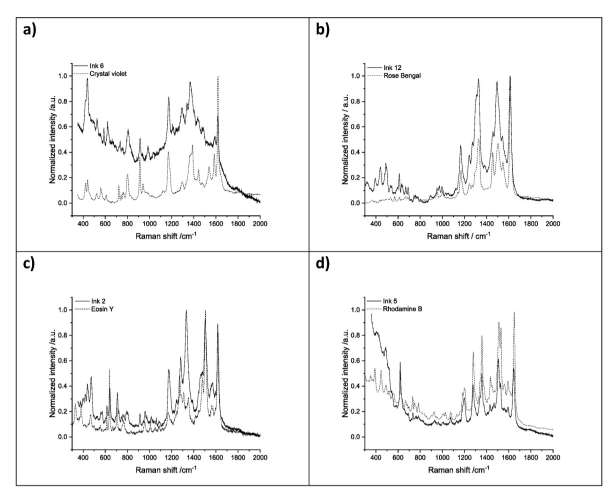


Fig. 2. SERS spectra of a) French Ink 6 (crystal violet-based); b) French Ink 12 (rose Bengal-based); c) French Ink 2 (eosin Y-based); d) French Ink 5 (rhodamine-based). Reference spectra are reported as dotted lines.

analysis of historical inks and applied to a collection of writing materials produced in France between the late 19th and the early 20th century. The array of analytical techniques used comprises surface enhanced Raman scattering spectroscopy (SERS), thin layer chromatography coupled to SERS (TLC-SERS), X-Ray fluorescence spectroscopy (XRF), high performance liquid chromatography coupled to diode array detection and tandem mass spectrometry (HPLC-DAD, HPLC-ESI-Q-ToF), and in situ derivatisation pyrolysis coupled to gas chromatography-MS (Py(HMDS)-GC/MS). Furthermore, HPLC-ESI-Q-ToF analysis also allowed us to perform studies on the degradation products of ink dyes. The collected results provided valuable information for art technological studies, dating and authentication processes.

3. Materials and methods

3.1. 19th-20th century inks

In the present work an historical red ink's collection from the late 19th and the early 20th century was analysed. The collection is composed by sixteen inks produced in France from different ink's manufacturers, and they were collected from an antique dealer (Vintage Waterman Pens, USA). The inks were contained in their original glass bottles, and their chromatic shades range from pale pink to bright purple and from peach orange to scarlet red. The inks are today in the form of a powder. To assess whether our procedure would be able to detect the dye/ pigment components in an historical sample, the powders were collected, and several sample pre-treatments were applied. The inks analysed were classified and the samples, with the associated samples

code, label, photos, colours, and results of the analyses are listed in Table 1.

3.2. Raman, surface enhanced Raman scattering (SERS) and thin layer chromatography- surface enhanced Raman scattering (TLC-SERS)

The synthesis of the silver nanoparticles (AgNPs) was performed according to Lee & Meisel procedure [46]. SERS measurements were made on a Renishaw InVia instrument coupled with an optical Leica DLML microscope, equipped with a NPLAN objective $20 \times$ and $50 \times$. Prior to SERS analysis, 1–2 mg of ink's powder were extracted in an ultrasonic bath (60 °C, 15 min) with 50 µL of ethanol. SERS measurements were made by mixing at room temperature on a glass slide 5 µL of AgNPs, 1 µL of KNO₃ (aggregating agent) and 1 µL of sample extract in ethanol. Spectra were acquired immediately by focusing on the top of the drop.

For TLC-SERS, the separation took place on Silica gel 60 plates (Supelco) as stationary phase and a isopropyl alcohol:acetone:ammonia (11:1:2) as eluent. 1–2 mg of ink's powder was suspended in 50 μ L of acetone and ~2 μ L of each extract was deposited onto the TLC plate by means of a glass capillary and eluted in a glass developing chamber. The separated spots were visualized under a UV lamp at 254 and 365 nm. The separated components were analysed directly on the TLC plate by placing 1 μ L of AgNPs on top of each spot [40].

More details can be found in [40] and in Supplementary Information S1.1-4.

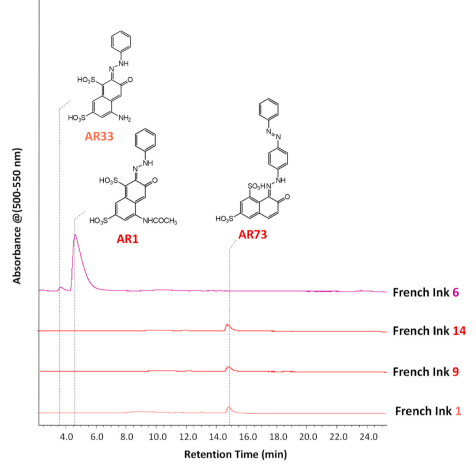


Fig. 3. HPLC-DAD chromatograms (500–550 nm) of French Ink 1 (light red profile), 9 (red profile), 14 (dark red profile), and 6 (violet profile). All chromatograms are presented in the same scale and are stacked for purpose of clarity.

3.3. X-Ray Fluorescence (XRF)

XRF experiments were performed using the Elio portable XRF Analyzer (XGLab, Italy), equipped with a 10–40 keV/200 μ A X-ray tube (Rh electrode, 1 mm collimated beam on the sample) and a large-area Energy Dispersive Si-Drift detector (130 eV FWHM at Mn K α). All the XRF spectra were acquired for 90 s with 40 kV of voltage and 80 μ A of current. Prior to XRF measurements, the ink's powder (1–2 mg) was dispersed in methanol. XRF analysis were performed on ink's mock-ups prepared by spreading the ink's dispersion on Whatman filter paper (USA, grade 42, diameter 110 mm, pure cellulose).

3.4. High performance liquid chromatography (HPLC-DAD-MS²)

The HPLC-DAD system consists of a PU-2089 quaternary pump equipped with a degasser, an AS-950 autosampler and an MD-2010 spectrophotometric diode array detector (DAD) (all modules Jasco International Co., Japan). The HPLC-ESI-Q-ToF consists of an HPLC 1200 Infinity, coupled to a Jet Stream ESI-Q-ToF 6530 Infinity detector and equipped with an Agilent Infinity autosampler (Agilent Technologies, Palo Alto, CA, USA) was used. Prior to HPLC analysis, 100–200 µg of ink's powder were suspended in 300–400 µL of 0.1% EDTA aqueous solution/DMF (EDTA-DMF, 1:1, v/v), and sonicated for 1 h at 60 °C. The supernatant was purified with PTFE syringe filters (4 mm thickness and 0.45 µm pore diameter, Agilent), and then injected in the chromatographic system.

Further details can be found in [47] or in the Supplementary Information file S1.5.

3.5. Pyrolysis coupled to gas chromatography-mass spectrometry (Py-GC/MS)

Py-GC/MS analysis was performed using a multi-shot pyrolyzer EGA/PY-3030D (Frontier Lab, Japan) coupled to a 8890 gas chromatograph, combined with a 5977B mass selective single quadrupole mass spectrometer detector (Agilent Technologies, USA). To characterize the ink's binder, pyrolysis with in-situ silylation using 4 μ L of hexamethyldisilazane (HMDS) as derivatizing agents was carried out. Further details can be found in [48] and in Supplementary Information file S1.6.

4. Results and discussion

Our studies lead to a broad overview of trends in ink's composition in $19^{th} - 20^{th}$ century in France. The results are summarized in Table 2 and will be presented in detail in the following paragraphs. The identification of both the dyes/pigments and binders was performed by comparison with standard and reference materials or based on matching data found in the literature. Instead, degradation products related to dye components, and never described before in the literature, were characterized through MS and MS² spectra interpretation. All the molecular markers determined by SERS, TLC-SERS, HPLC-DAD-MS² and Py (HMDS)-GC/MS are summarized in Tables 3–6, respectively (Supplementary Information file).

SERS spectroscopy was first carried out to obtain a preliminary screening of the ink's dyes and allowed us to split the historical inks into three groups (as presented in Table 2): the first, encompassing the inks

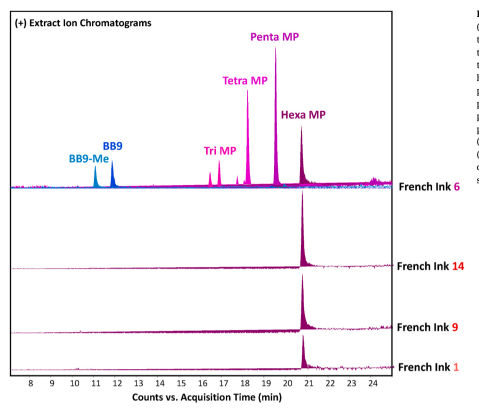


Fig. 4. HPLC-ESI-Q-ToF Extract Ion Chromatograms (EIC) obtained by plotting the counts in function of time of the ions corresponding to the molecular ions of the single species detected in the EDTA-DMF extracts of the French Inks 1, 9, 14 and 6. The detected species, highlighted in the chromatograms, are hexa-N-methyl pararosaniline (Hexa MP, $C_{25}H_{30}N_3^+$), penta-N-methyl pararosaniline (Penta MP, $C_{22}H_{26}N_3^+$), terta-N-methyl pararosaniline (Tetra MP, $C_{22}H_{26}N_3^+$), tri-N-methyl pararosaniline (Tri MP, $C_{22}H_{26}N_3^+$), tri-N-methyl pararosaniline (Tri MP, $C_{22}H_{24}N_3^+$), methylene blue (BB9, $C_{16}H_{18}N_3S^+$) and demethylated methylene blue (BB9–Me, $C_{15}H_{16}N_3S^+$). Positive acquisition mode. All chromatograms are presented in the same scale and are stacked for purpose of clarity.

containing triarylmethine pigments (methyl or crystal violet); two subgroups, comprising the inks based on xanthene pigments with halogen substituents, namely rose Bengal and eosin Y; the third one, containing the amino-substituted xanthene pigment rhodamine. The specific wavenumbers obtained for each standard pigment and French ink are reported in Tables S2-S4 (Supplementary Information file). In detail.

- French Inks 1, 6, 9 and 14 were preliminary classified as **methyl** or **crystal violet-based inks** (Fig. 2a). Methyl (Mv) and crystal violet (Cv) belong to the class of *triarylmethine pigments*: Mv (C.I. 42535), introduced on the market in 1861, consists of a mixture of tetra-, penta- and hexa-*N*-methyl pararosaniline, while Cv (C.I. 42555), commercialized in 1883 by Kern and Caro, contains hexa-N-methyl pararosaniline only [49]. For their differentiation, a separative technique is necessary;
- French Ink 12 was identified as rose Bengal-based ink (Fig. 2b). Rose Bengal (Acid Red 94, C.I. 45440), first synthesised by Gnehm in 1882, belongs to the class of xanthenes, imparts a pink colour and is a halogen-containing dye-pigment (chlorine and iodine atoms) [50]; French Inks 2, 7, 11, 13 and 15 were characterized as eosin Y-based inks (Fig. 2c). Eosin Y (Acid Red 87, C.I. 45380) is an acid dye produced since 1873 thanks to Heinrich Caro. It is a brominated fluorescein which is normally orange in its insoluble acid form [51, 52];
- French Inks 4, 5, 10 and 16 were determined as **rhodamine-based inks** (Fig. 2d). Specifically, rhodamine B (Basic Violet 10, C.I. 45170) was detected in ink's formulations 4, 5, and 16, while rhodamine 6G (Basic Red 1, C.I. 45160) was determined in French Ink 4 and 10. Both dyes are basic xanthenes: rhodamine 6G is characterized by a scarlet tone and was patented in 1887, while rhodamine B is a highly fluorescent violet dye commercialized in 1887 by Cérésole [53,54].

To obtain additional information regarding the trend's composition of 19th - 20th century ink's formulations, further analyses were carried out with liquid chromatography techniques (TLC-SERS, HPLC-DAD, HPLC-ESI-Q-ToF), since a separation step is fundamental for the analysis of complex mixtures such as writing inks.

4.1. Methyl/crystal violet-based inks

Concerning the inks determined as methyl violet (Mv) or crystal violet (Cv) based, further information was achieved by liquid chromatography techniques. TLC-SERS provided the identification of cotton scarlet (AR73) in French Inks 1, 9, and 14. Thus, the technique did not confirm SERS results, and this could be due to the non-optimal chromatographic conditions adopted for basic triarylmethine dyes. The presence of AR73 in French Inks 1, 9, and 14 was further confirmed by HPLC-DAD (Fig. 3), which allowed us to detect also fast acid magenta B (AR33, 3.5 min) and amido naphthol red G (AR1, 4.8 min) in French Ink 6. These three acid dyes (AR73, AR1, AR33) were also identified by HPLC-ESI-Q-ToF (negative ionisation mode) results. According to the current knowledge, AR33 was identified as a synthetic intermediate of AR1 [55], since the two dyes only differ in the presence of an acetyl substituent. Alternatively, the formation of AR33 could be the result of the deacetylation of AR1, since ageing phenomenon involving a deacetylation step are known in the literature [56]. Further information was achieved by liquid chromatography-tandem mass spectrometry in positive ionisation mode (Fig. 4), which finally provided the identification of French Inks 1,9 and 14 as Cv-based formulations, confirming the results of SERS spectroscopy. Instead, in French Ink 6 the recognition of the demethylated forms of hexa-N-methyl pararosaniline (Hexa MP, 20.7 min; Penta MP, 19.5 min; Tetra MP, 18.2 and 17.7 min; Tri MP, 16.9 and 16.4 min) allowed us to hypothesize the use of Mv in the formulation, or that the originally present Cv underwent the photodegradation pattern described in the literature [18]. Moreover, a blue dye belonging to thiazine dye class, methylene blue (BB9, 11.8 min), was determined in French Ink 6. The high selectivity and sensitivity of liquid chromatography-tandem mass spectrometry also provided the identification of an unknown compound, which was characterized as the demethylated form of BB9 (BB9-Me, 11.1 min) through the

a)

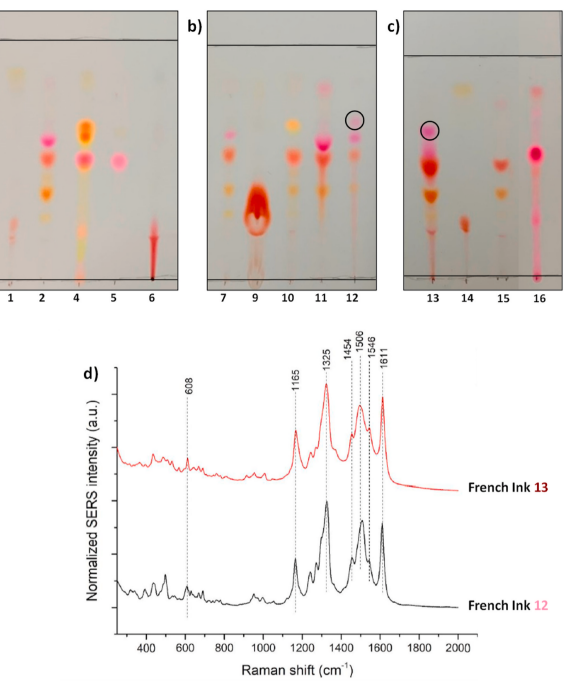


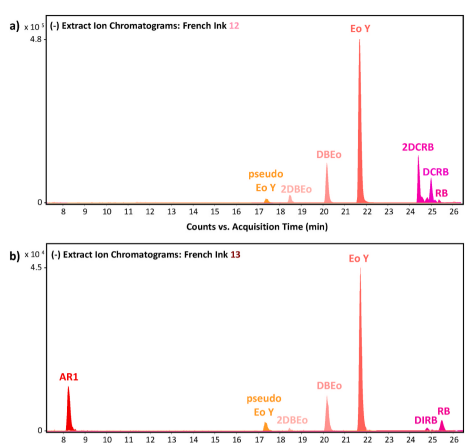
Fig. 5. TLC of the French Inks under visible light. a) TLC of French Inks 1, 2, 4, 5 and 6; b) TLC of French Inks 7, 9, 10, 11 and 12; c) TLC of French Inks 13, 14, 15 and 16; d) SERS spectra acquired on rose Bengal stain (highlighted with black circle).

interpretation of its tandem mass spectrum (Figure S1). BB9–Me could be a synthetic by-product, or be due to a degradation process, as typical photo-oxidation pathways entailing the loss of methyl groups are known for dyes featuring substituted amino functionalities [18,31].

Finally, Py(HMDS)-GC/MS analysis allowed us to determine levoglucosan (in its derivatized form) in French Ink 6 (Figure S2). Levoglucosan is the pyrolytic marker of several materials, and in the heritage field it could be ascribed to the presence of cellulose or polysaccharidic gums, such as gum Arabic, mesquite gum etc. Since gum Arabic was the most used binder in historical inks [6], and given the context of the analysis, the identification of levoglucosan suggests the presence of this binder in French Ink 6 formulation. In addition, pyrolytic markers related to *amido naphthol red G* and *cotton scarlet* were identified in French Ink 6 and 9 [57,58], respectively, while for French Ink 1 and 14 no additional information was obtained with respect to the analytical blank.

4.2. Eosin Y and Rose Bengal-based inks

From SERS analysis *eosin* Y (Eo Y) was determined in French Inks 2, 7, 11, 13 and 15, while *rose Bengal* (RB) was detected in French Ink 12 only. The results acquired with SERS spectroscopy were all confirmed by TLC-SERS, which also additionally provided the identification of rose Bengal in French Ink 13 (Fig. 5). The presence of the tetra-brominated dye (Eo Y) has also been demonstrated by XRF spectroscopy (Figure S3), which allow to identify bromine in all the Eo Y-based inks



Counts vs. Acquisition Time (min)

Fig. 6. HPLC-ESI-Q-ToF Extract Ion Chromatograms (EIC) obtained by plotting the counts in function of time of the ions corresponding to the molecular ions of the single species detected in the EDTA-DMF extracts of French Ink 12 (Figure a) and French Ink 13 (Figure b). The detected species, highlighted in the chromatograms, are rose Bengal (RB, $C_{20}H_5Cl_4I_4O_5$), dechlorinated rose Bengal (DCRB, $C_{20}H_6Cl_3I_4O_5$), deidoinated rose Bengal (DCRB, $C_{20}H_6Cl_3I_4O_5$), deidoinated rose Bengal (DIRB, $C_{20}H_6Cl_4I_3O_5$), comparison Y (Eo, $C_{20}H_8Br_4O_5$), debrominated eosin Y (DBEo, $C_{20}H_8Br_4O_5$), bis-debrominated eosin Y (2DBEo, $C_{20}H_9Br_2O_5$) and amido naphthol red G (AR1, $C_{18}H_14N_3O_8S_2$). Negative acquisition mode.

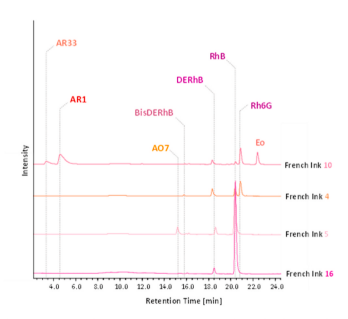


Fig. 7. HPLC-DAD chromatograms (500–550 nm) of French Ink 16 (violet profile), 5 (light pink profile), 4 (orange profile) and 6 (pink profile). All chromatograms are presented in the same scale and are stacked for purpose of clarity.

spectra (French Inks 2, 7, 11, 13 and 15). Instead, XRF analysis did not provide any information related to chlorine and iodine atoms, typical of rose Bengal dye. Further information were obtained by high performance liquid chromatography analysis: HPLC-DAD (Figure S4) and

HPLC-ESI-O-ToF in negative ionisation mode (Fig. 6) consistently detected eosin Y and its debrominated molecular markers [59] in all the Eo Y and RB- based inks, allowing us to further determine eosin Y in French Ink 12. Moreover, amido naphthol red G (AR1) and fast acid magenta B (AR33) were identified in French Ink 13 through HPLC techniques. Besides, HPLC-ESI-O-ToF provided us with the identification of further degradation products or synthetic by-products. The examination of the exact masses, isotopic patterns, and tandem mass spectra of the precursor ions allowed us to hypothesize the molecular structure of several secondary components of the dye mixture. Specifically, chlorine and iodine containing species were assigned as Rose Bengal's synthesis by-products or degradation markers. The dechlorinated (DCRB, 25.0 min), bis-dechlorinated (2DCRB, 24.4 min) and deiodinated (DIRB, 24.7 min) dye molecules were all detected. Considering the literature studies on eosin Y [59], the hypothesis of subsequent dechlorination or deiodination from the Rose Bengal structure is a plausible degradation pathway. Furthermore, dechlorinated derivatives of rose Bengal were already detected by Tamburini et al. in a textile sample book dated 1893 [60]. Interestingly, in all the Eo Y-based inks (French ink 2, 7, 11, 12, 13 and 15) a further unknow species characterized by the isotopic pattern of a dye containing four bromine atoms was detected. The exact mass of the molecular ion $([M - H]^{-} = 716.736)$ could be attributed to the molecular formula $C_{24}H_{14}Br_4O_6,\ proving a relation with eosin dyes.$ However, further studies are needed to determine its molecular structure. All the data acquired on this unknow marker are reported in the Supplementary Information file (Figure S5 and S6).

Finally, Py(HMDS)-GC/MS results ensured the identification of pyrolytic markers of eosin Y (benzoic acid TMS) in French Ink 11, 13 and 15, and of amido naphthol red G in French Ink 13 [57,58]. Instead, the pyrograms obtained for French Ink 2, 7 and 12 were at blank level. No binder or additive was detected for this group of samples.

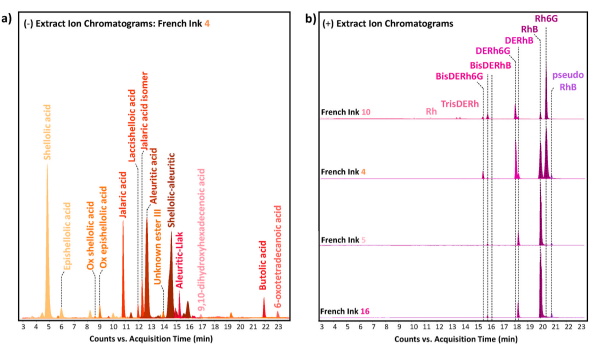


Fig. 8. Chromatograms obtained for Rh-based inks. **a)** HPLC-ESI-Q-ToF Extract Ion Chromatograms (EICs) obtained by plotting the counts in function of time of the ions corresponding to the molecular ions of the single species detected in the EDTA-DMF extract of French Ink 4 in negative ionisation mode. The molecular markers highlighted are related to shellac resin. **b)** HPLC-ESI-Q-ToF Extract Ion Chromatograms (EICs) obtained by plotting the counts in function of time of the ions corresponding to the molecular ions of the single species detected in the EDTA-DMF extracts of the French Inks 16, 5, 4, and 10 in positive ionisation mode. The detected species, highlighted in the chromatograms, are rhodamine 6G (Rh 6G, $C_{28}H_{31}N_2O_3^+$), Rhodamine B (Rh B, $C_{28}H_{31}N_2O_3^+$) and its degradation products (DERhB, $C_{26}H_{27}N_2O_3^+$; BisDERhB, $C_{24}H_{23}N_2O_3^+$; TrisDERhB, $C_{22}H_{19}N_2O_3^+$; Rh, $C_{20}H_{15}N_2O_3^+$). All chromatograms are presented in the same scale and are stacked for purpose of clarity.

4.3. Rhodamine-based inks

The final subgroup is related to Rhodamine dyes. TLC-SERS experiments confirmed the SERS results and contributed to the identification of rhodamine B (RhB) and eosin Y (EoY) in French Ink 10, besides the already detected rhodamine 6G. HPLC-DAD chromatograms obtained for the EDTA-DMF extracts of French Inks 4,5, 10, and 16 are reported in Fig. 7. RhB (19.5 min) and its photo-oxidation products DERhB (17.8 min) and bisDERhB (16.3 min) [53] were detected in all the rhodamine-based inks. HPLC-DAD analysis also enabled us to verify the presence of both rhodamine dyes (RhB and Rh6G) in French Inks 4 and 10, proving that the combination of rhodamine B and 6G was widespread, besides in textiles [44] and paint tubes manufacturing, also in historical ink's formulations. Moreover, orange II (AO7, 15.2 min) was determined in French Ink 5, while EoY, AR1, and AR33 were detected in French Ink 10. Both TLC-SERS and HPLC-DAD results were confirmed by HPLC-ESI-Q-ToF analysis, which allowed us to further investigate the composition of these historical ink's formulations. Positive ionisation mode enabled us to identify first trisDERhB and trisDERh6G (13.7 and 13.5 min, respectively) and Rh (11.5 min) in French Ink 10 (Fig. 8b), highlighting an advanced degradation compared to the other rhodamine-based inks [53]. Furthermore, a product related to rhodamine B was detected in French Ink 4, 5 and 16. This unknown molecular marker is characterized by a molecular ion $[M]^+$ at 729.452 m/z, which gives MS^2 fragments similar to those of rhodamine B (m/z = 443.235, 399.169) [53]. All the data acquired on this unknow marker are reported in the Supplementary Information file (Figure S7). Interestingly, the chromatogram acquired in negative ionisation mode features the free acid and esters profile typical of shellac resin [61] in French Inks 4, 5 and 16, as shown in Fig. 8 a. Moreover, negative ionisation mode allowed us to identify a methylated analogous of AO7 in French Ink 5 (Figure S8). The MS^2 fragmentation of this species seems to follow the same pathway observed for AO7 (Figure S9). This suggests the presence of an additional methyl group on the sulfonated ring. The detected molecular marker could be related to Orange 2R (AO8); however, as the standard is not commercially available, this hypothesis was not confirmed.

The presence of shellac resin in French Inks 4 and 5 was further confirmed by Py(HMDS)-GC/MS (Figure S2), which allowed us to detect butolic acid in its derivatized form [62]. Shellac resin, introduced into Europe as lac dye source in the 12th century and as varnish in the late 16th century [61,63,64], was widespread in ink's recipes to increase the viscosity of the writing materials [65]. Moreover, pyrolytic markers that may be related to rhodamine dyes were detected in French Ink 4, 5 and 16, while the identification of β -naphthol TMS markers in French Ink 5 allow to confirm the presence of orange II [57,58]. Concerning the French Ink 10, the resulting pyrogram was at blank level.

5. Conclusions

To discover the secrets behind the industrial production of writing materials, in the present work a multi-analytical approach was applied for the investigation of an historical ink's collection. The results obtained show that the combination between spectroscopic and chromatographic techniques is a valuable approach to gain information on complex mixture such as writing inks. Trends in historical ink's formulations have been outlined, leading to an overview of the composition of French inks in the late 19th - early 20th century. Shellac resin, used as an additive to increase the viscosity of writing materials, was detected in three formulations, while the presence of gum Arabic, the binder the most used in historical ink's recipes, was hypothesized only in one ink. Concerning the other inks, no information was obtained for the binder component. This result is in agreement with historical recipes reported in the manuscripts, where the use of binders is not mentioned, e.g. in the details on alizarine ink produced by Leonhardi industry, and Bottger's recipe of logwood ink. The dyes/pigments analysis enabled us to

determine that ink's formulations were mainly based on crystal violet, eosin Y, rose Bengal and rhodamine B and 6G. In particular, mixtures of cotton scarlet and crystal violet were quite characteristic in 19th-20th century ink's formulations. Moreover, eosin Y was widespread in ink formulations of the selected historical period, either as a single component or mixed with other dyes such as amido naphthol red G and rose Bengal. Between all the formulations identified, the one related to rhodamine dves was the most peculiar. Rhodamine B was widely used in $19^{th} \ 20^{th}$ century inks. However, each manufacturing industry had different secrets on how to tune the composition to obtain specific colours and more performing writing materials. This reflects in ink's formulations in which *rhodamine B* was used alone and in others in which it was combined with complex mixtures of xanthenes and/or hydrazone pigments. Furthermore, new molecular markers for the identification of methylene blue, rose Bengal, eosin Y and rhodamine B by liquid chromatography analysis were detected and partially characterized. In addition, TMS-derivatized pyrolytic markers that could be associated to amido naphthol red G, cotton scarlet, orange II, eosin Y and rhodamine dyes were identified in several French ink formulations. Thus, this paper highlights the suitability of Pv(HMDS)-GC/MS methods for the identification of synthetic dves. Our investigation provides useful information for art technological studies of historical ink collection, expanding the datasetavailable in the field of dye degradation products for ink analysis of manuscripts and drawings. Further studies on the historical synthetic routes or possible degradation pathways will be performed, to deepen our understanding of the early days of organic chemistry.

CRediT authorship contribution statement

Adele Ferretti: Methodology, Validation, Investigation, Visualization, Writing – original draft, Writing – review & editing. Ilaria Degano: Conceptualization, Methodology, Writing – review & editing, Supervision. Stefano Legnaioli: Methodology, Investigation, Writing – review & editing. Beatrice Campanella: Methodology, Validation, Investigation. Aurora Sainati: Investigation, Visualization. Maria Perla Colombini: Writing – review & editing, Supervision, Project administration.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

Acknowledgements

Anne-Laure Carrega and Franck Baille, Hôtel des Ventes de Monte-Carlo, are acknowledged for providing the samples under study. Ilaria Bonaduce, Università di Pisa, is acknowledged for the fruitful discussion on shellac resin. The Centre for Instrument Sharing of the University of Pisa (CISUP) is acknowledged for providing the pyrolysis instrumentation.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.dyepig.2023.111672.

References

 Nardella F, Landi N, Degano I, Colombo M, Serradimigni M, Tozzi C, et al. Chemical investigations of bitumen from Neolithic archaeological excavations in Italy by GC/MS combined with principal component analysis. Anal Methods 2019; 11:1449–59. https://doi.org/10.1039/c8ay02429d.

- [2] Sabatini F, Bacigalupo M, Degano I, Javér A, Hacke M. Revealing the organic dye and mordant composition of Paracas textiles by a combined analytical approach. Herit Sci 2020;8:1–17. https://doi.org/10.1186/s40494-020-00461-5.
- [3] Sharma N, Agarwal A, Negi Y, Bhardwaj H, Jaiswal J. History and chemistry of ink-A review. World J Pharmaceut Res 2014;3:2096–105.
- [4] Feraru DL, Meghea A. Possibilities to differentiate ballpoint pen inks by spectroscopic and chromatographic techniques. UPB Sci Bull Ser B Chem Mater Sci 2014;76:113–26.
- [5] Drury N, Ramotowski R, Moini M. A comparison between DART-MS and DSA-MS in the forensic analysis of writing inks. Forensic Sci Int 2018;289:27–32. https://doi. org/10.1016/j.forsciint.2018.05.009.
- [6] Mitchell CA, Hepworth TC. Inks: their composition and manufacture. London: C. Griffin & Company LtD; 1904.
- [7] Carvalho D N. Forty centuries of ink or A chronological narrative concerning ink and its backgrounds. London: C.Griffin & Company LtD; 1904.
- [8] Pines CC. The story of ink. Am J Police Sci 1931;2:290–301. https://doi.org/ 10.2307/1147356.
- [9] Wei S, Fang X, Cao X, Schreiner M. Characterization of the materials used in Chinese ink sticks by pyrolysis-gas chromatography-mass spectrometry. J Anal Appl Pyrolysis 2011;91:147–53. https://doi.org/10.1016/j.jaap.2011.02.001.
- [10] Teixeira N, Nabais P, de Freitas V, Lopes JA, Melo MJ. In-depth phenolic characterization of iron gall inks by deconstructing representative Iberian recipes. Sci Rep 2021;11:1–11. https://doi.org/10.1038/s41598-021-87969-3.
- [11] Kolar J, Strlič M. Iron gall inks: on manufacture characterisation, degradation and stabilisation. 2006.
- [12] Centeno SA, Ropret P, del Federico E, Shamir J, Itin B, Jerschow A. Characterization of Al(III) complexes with hematein in artistic alum logwood inks. J Raman Spectrosc 2010;41:445–51. https://doi.org/10.1002/jrs.2455.
- [13] Roldán ML, Centeno SA, Rizzo A. An improved methodology for the characterization and identification of sepia in works of art by normal Raman and SERS, complemented by FTIR, Py-GC/MS, and XRF. J Raman Spectrosc 2014;45: 1160–71. https://doi.org/10.1002/jrs.4620.
- [14] Slama H Ben, Bouket AC, Pourhassan Z, Alenezi FN, Silini A, Cherif-Silini H, et al. Diversity of synthetic dyes from textile industries, discharge impacts and treatment methods. Appl Sci 2021;11:1–21. https://doi.org/10.3390/app11146255.
- [15] Lech K, Wilicka E, Witowska-Jarosz J, Jarosz M. Early synthetic dyes a challenge for tandem mass spectrometry. J Mass Spectrom 2013;48:141–7. https://doi.org/ 10.1002/jms.3090.
- [16] Reissland B, Gaibor AN Proaño, Ligterink Jgn F. Exploring the late 19th-century landscape of ink manufacturing via a collection of 90 bottles. ICOM-CC 18th Trienn Conf; 2017.
- [17] Gessner T, Mayer U. Triarylmethane and diarylmethane dyes. Ullmann's Encycl. Ind. Chem. 2000 10.1002/14356007.a27_179.
- [18] Colombini MP, Degano I, Nevin A. Analytical chemistry for the study of paintings and the detection of forgeries. 2022.
- [19] Moretti P, Germinario G, Doherty B, van der Werf ID, Sabbatini L, Mirabile A, et al. Disclosing the composition of historical commercial felt-tip pens used in art by integrated vibrational spectroscopy and pyrolysis-gas chromatography/mass spectrometry. J Cult Herit 2019;35:242–53. https://doi.org/10.1016/j. culher.2018.03.018.
- [20] Germinario G, Rigante ECL, van der Werf ID, Sabbatini L. Pyrolysis gas chromatography-mass spectrometry of triarylmethane dyes. J Anal Appl Pyrolysis 2017;127:229–39. https://doi.org/10.1016/j.jaap.2017.08.001.
- [21] Fox MA. In: Fox MA, editor. Dyes and pigments Glossary for the worldwide transportation of dangerous Goods and hazardous materials. Berlin, Heidelberg: Springer Berlin Heidelberg; 1999. p. 63–4. https://doi.org/10.1007/978-3-662-11890-0_24.
- [22] Wright P, Staff Ub. Xanthene Dyes. In: Kirk-Othmer Encyclopedia of Chemical Technology. John Wiley & Sons, Inc; 2014. https://doi.org/10.1002/0471238 961.2401142023090708.a01.pub2.
- [23] Fox MA. Glossary for the worldwide transportation of dangerous goods and hazardous materials. 1999. p. 63–4. https://doi.org/10.1007/978-3-662-11890-0_ 24.
- [24] Gagliardi L, Orsi D De, Cavazzutti G, Multari G, Tonelli D. HPLC determination of rhodamine B (C. I. 45170) in cosmetic products. Chromatographia 1996;43:76–8.
- [25] Confortin D, Neevel H, Brustolon M, Franco L, Kettelarij AJ, Williams RM, et al. Crystal violet: study of the photo-fading of an early synthetic dye in aqueous solution and on paper with HPLC-PDA, LC-MS and FORS. J Phys Conf Ser 2010; 231. https://doi.org/10.1088/1742-6596/231/1/012011.
- [26] den Uijl MJ, Lokker A, van Dooren B, Schoenmakers PJ, Pirok BWJ, van Bommel MR. Comparing different light-degradation approaches for the degradation of crystal violet and eosin Y. Dyes Pigments 2022;197:109882. https://doi.org/10.1016/j.dyepig.2021.109882.
- [27] Angelin EM, Oliveira MC, Nevin A, Picollo M, Melo MJ. To be or not to be an azo pigment: chemistry for the preservation of historical β-naphthol reds in cultural heritage. Dyes Pigments 2021;190. https://doi.org/10.1016/j. dyepig.2021.109244.
- [28] Bafana A, Devi SS, Chakrabarti T. Azo dyes: past, present and the future. Environ Rev 2011;19:350-70. https://doi.org/10.1139/a11-018.
- [29] Stolz A. Basic and applied aspects in the microbial degradation of azo dyes. Appl Microbiol Biotechnol 2001;56:69–80. https://doi.org/10.1007/s002530100686.
- [30] Tanaka K, Padermpole K, Hisanaga T. Photocatalytic degradation of commercial azo dyes. Water Res 2000;34:327–33. https://doi.org/10.1016/S0043-1354(99) 00093-7.

A. Ferretti et al.

- [31] Berneth H. Azine Dyes. Ullmann's Encycl Ind Chem 2008. https://doi.org/ 10.1002/14356007.a03_213.pub3.
- [32] Vara J, Ortiz CS. Thiazine dyes: evaluation of monomeric and aggregate forms. Spectrochim Acta Part A Mol Biomol Spectrosc 2016;166:112–20. https://doi.org/ 10.1016/j.saa.2016.05.005.
- [33] Li D, Lv L, Sun P, Zhou W, Wang P, Wu J, et al. The facile synthesis of novel phenothiazine derivatives for blue, yellow-green, and red light emission. Dyes Pigments 2009;83:180–6. https://doi.org/10.1016/j.dyepig.2009.04.006.
- [34] Mall C, Tiwari S, Solanki PP. Comparison of dye (oxazine and thiazine) materials as a photosensitizer for use in photogalvanic cells based on molecular interaction with sodium dodecyl sulphate by spectral study. J Saudi Chem Soc 2019;23:83–91. https://doi.org/10.1016/j.jscs.2018.04.007.
- [35] Ferretti A, Sabatini F, Degano I. A model iron gall ink : an in-depth study of ageing processes involving gallic acid. Molecules 2022;27.
- [36] Sundberg BN, Pause R, van der Werf ID, Astefanei A, van den Berg KJ, van Bommel MR. Analytical approaches for the characterization of early synthetic organic pigments for artists' paints. Microchem J 2021;170. https://doi.org/ 10.1016/j.microc.2021.106708.
- [37] Rosi F, Grazia C, Fontana R, Gabrieli F, Pensabene Buemi L, Pampaloni E, et al. Disclosing Jackson Pollock's palette in Alchemy (1947) by non-invasive spectroscopies. Herit Sci 2016;4:1–13. https://doi.org/10.1186/s40494-016-0089-
- [38] Burgio L, Clark RJH. Library of FT-Raman spectra of pigments, minerals, pigment media and varnishes, and supplement to existing library of Raman spectra of pigments with visible excitation, vol. 57; 2001. https://doi.org/10.1016/S1386-1425(00)00495-9.
- [39] Pause R, van derWerf ID, van den Berg KJ. Identification of pre-1950 synthetic organic pigments in artists' paints. A non-invasive approach using handheld Raman spectroscopy. Heritage 2021;4:1348–65. https://doi.org/10.3390/ heritage4030073.
- [40] Campanella B, Botti J, Cavaleri T, Cicogna F, Legnaioli S, Pagnotta S, et al. The shining brightness of daylight fluorescent pigments: Raman and SERS study of a modern class of painting materials. Microchem J 2020;152:104292. https://doi. org/10.1016/j.microc.2019.104292.
- [41] Cañamares MV, Reagan DA, Lombardi JR, Leona M. TLC-SERS of mauve, the first synthetic dye. J Raman Spectrosc 2014;45:1147–52. https://doi.org/10.1002/ jrs.4508.
- [42] López-Montes AM, Dupont AL, Desmazières B, Lavédrine B. Identification of synthetic dyes in early colour photographs using capillary electrophoresis and electrospray ionisation-mass spectrometry. Talanta 2013;114:217–26. https://doi. org/10.1016/j.talanta.2013.04.020.
- [43] Tamburini D, Shimada CM, McCarthy B. The molecular characterization of early synthetic dyes in E. Knecht et al's textile sample book "A Manual of Dyeing" (1893) by high performance liquid chromatography - diode array detector - mass spectrometry (HPLC-DAD-MS). Dyes Pigments 2021;190:109286. https://doi.org/ 10.1016/j.dyepig.2021.109286.
- [44] Tamburini D, Breitung E, Mori C, Kotajima T, Clarke ML, McCarthy B. Exploring the transition from natural to synthetic dyes in the production of 19th-century Central Asian ikat textiles. Herit Sci 2020;8:1–27. https://doi.org/10.1186/ s40494-020-00441-9.
- [45] van Bommel MR, Berghe I Vanden, Wallert AM, Boitelle R, Wouters J. Highperformance liquid chromatography and non-destructive three-dimensional fluorescence analysis of early synthetic dyes. J Chromatogr A 2007;1157:260–72. https://doi.org/10.1016/j.chroma.2007.05.017.
- [46] Lee PC, Meisel D. Adsorption and surface-enhanced Raman of dyes on silver and gold sols. J Phys Chem 1982;86:3391–5. https://doi.org/10.1021/j100214a025.
- [47] Sabatini F, Degano I. Investigating the fragmentation pathways of β-naphthol pigments using liquid chromatography/electrospray ionization quadrupole timeof-flight mass spectrometry. Rapid Commun Mass Spectrom 2020;34:1–10. https:// doi.org/10.1002/rcm.8789.

- [48] La Nasa J, Nodari L, Nardella F, Sabatini F, Degano I, Modugno F, et al. Chemistry of modern paint media: the strained and collapsed painting by Alexis Harding. Microchem J 2020;155:104659. https://doi.org/10.1016/J. MICROC.2020.104659.
- [49] Degano I, Sabatini F, Braccini C, Colombini MP. Triarylmethine dyes: characterization of isomers using integrated mass spectrometry. Dyes Pigments 2019;160:587–96. https://doi.org/10.1016/j.dyepig.2018.08.046.
- [50] Neckers DC. Rose bengal. J Photochem Photobiol Chem 1989;47:1–29. https://doi. org/10.1016/1010-6030(89)85002-6.
- [51] Sabatini F, Eis E, Degano I, Thoury M, Bonaduce I, Lluveras-Tenorio A. The issue of eosin fading: a combined spectroscopic and mass spectrometric approach applied to historical lakes. Dyes Pigments 2020;180:108436. https://doi.org/10.1016/j. dyepig.2020.108436.
- [51] Chieli A, Romani A, Degano I, Sabatini F, Tognotti P, Miliani C. New insights into the fading mechanism of Geranium lake in painting matrix. Dyes Pigments 2020; 181:108600. https://doi.org/10.1016/j.dyepig.2020.108600.
- [53] Sabatini F, Giugliano R, Degano I. Photo-oxidation processes of Rhodamine B: a chromatographic and mass spectrometric approach. Microchem J 2018;140: 114–22. https://doi.org/10.1016/j.microc.2018.04.018.
- [54] Ferreira BRV, Correa DN, Eberlin MN, Vendramini PH. Fragmentation reactions of rhodamine B and 6G as revealed by high accuracy orbitrap tandem mass spectrometry. J Braz Chem Soc 2017;28:136–42. https://doi.org/10.5935/0103-5053.20160156.
- [55] Souto C. Analysis of Early Synthetic Dyes with HPLC-DAD-MS: an important database for analysis of colorants used in cultural heritage, 2010. 0–45.
- [56] Blanton TN, Kaduk JA, Johnson Q. X-ray diffraction characterization of a distorted Debye-Scherrer film strip - the effect of deacetylation on cellulose triacetate and an improved structural model for cellulose II. Powder Diffr 2014;29:108–12. https:// doi.org/10.1017/S0885715614000141.
- [57] Ghelardi E, Degano I, Colombini MP, Mazurek J, Schilling M, Learner T. Py-GC/MS applied to the analysis of synthetic organic pigments: characterization and identification in paint samples. Anal Bioanal Chem 2015;407:1415–31. https://doi. org/10.1007/s00216-014-8370-y.
- [58] Russell J, Singer BW, Perry JJ, Bacon A. The identification of synthetic organic pigments in modern paints and modern paintings using pyrolysis-gas chromatography-mass spectrometry. Anal Bioanal Chem 2011;400:1473–91. https://doi.org/10.1007/s00216-011-4822-9.
- [59] Sabatini F, Degano I, Colombini MP. Development of a method based on highperformance liquid chromatography coupled with diode array, fluorescence, and mass spectrometric detectors for the analysis of eosin at trace levels. Sep Sci Plus 2020;3:207–15. https://doi.org/10.1002/sscp.202000002.
- [60] Tamburini D, Shimada CM, McCarthy B. The molecular characterization of early synthetic dyes in E. Knecht et al's textile sample book "A Manual of Dyeing" (1893) by high performance liquid chromatography - diode array detector - mass spectrometry (HPLC-DAD-MS). Dyes Pigments 2021;190:109286. https://doi.org/ 10.1016/j.dyepig.2021.109286.
- [61] Tamburini D, Dyer J, Bonaduce I. The characterisation of shellac resin by flow injection and liquid chromatography coupled with electrospray ionisation and mass spectrometry. Sci Rep 2017;7:1–15. https://doi.org/10.1038/s41598-017-14907-7.
- [62] Colombini MP, Bonaduce I, Gautier G. Molecular pattern recognition of fresh and aged shellac. Chromatographia 2003;58:357–64. https://doi.org/10.1365/s10337-003-0037-3.
- [63] Castro R, Pozzi F, Leona M, Melo MJ. Combining SERS and microspectrofluorimetry with historically accurate reconstructions for the characterization of lac dye paints in medieval manuscript illuminations. J Raman Spectrosc 2014;45:1172–9. https://doi.org/10.1002/jrs.4608.
- [64] Masschelein-Kleiner L. Ancient binding media. Varnishes and Adhesives; 1995.
- [65] Centeno SA, Lladò Buisan V, Ropret P. Raman study of synthetic organic pigments and dyes in early lithographic inks (1890–1920). J Raman Spectrosc 2006;37: 1111–8. https://doi.org/10.1002/jrs1594.