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Surface-Enhanced Raman Spectroscopy for the Investigation of Chromogenic Motion Picture Films: A Preliminary Study

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Abstract: In the present work, surface-enhanced Raman spectroscopy (SERS) is proposed for the identification of dyes in chromogenic films. These substances, which are generated within the film itself during the colour development process, are first studied on reference molecules synthesised for the purpose and, subsequently, on dyes extracted from a chromogenic film. SERS spectroscopy proved to be an efficient technique for their investigation, providing a proof of concept for its applicability for further studies on this complex topic. SERS spectra of the reference molecules were also compared with normal Raman spectra, acquired with a spectrometer based on SSE™ technology. Furthermore, the excellent SERS properties of anisotropic nanomaterials, such as silver nanostars, have been confirmed.

Keywords: SERS; chromogenic motion picture films; dyes; silver nanostars; SSE™ Raman

1. Introduction

In the last century, photographic and motion picture film materials have enriched our cultural heritage. At the beginning, colours were applied on every single frame of the film by hand or with special stencils. Subtractive synthesis was then developed: the chromatic tone was initially obtained with the dye-transfer process, i.e., adding from one to three layers of colour on the film. In this regard, it is worth mentioning the Technicolor process n. IV which provides that three black-and-white negative films coloured by imbibition with magenta, cyan and yellow dyes, are printed on a film coated with a substance capable of absorbing the dyes. Finally, chromogenic Monopacks were introduced, where the colour-forming substances are divided into several layers of the emulsion, and are still in use nowadays [1]. These motion picture films, despite being quite recent, tend to degrade due to environmental conditions (i.e., humidity, temperature, exposure to light) and the main problem concerns the loss of colour due to the degradation of the dye. Currently digital restoration is the only means to restore the original appearance of films, but it requires the formulation of hypotheses on the evolution of the degradation processes as a function of time, with the aim of obtaining information on the initial colorimetric coordinates. In this scenario, a very limited number of studies have been reported on the identification of colouring substances in film materials [2,3]. In this regard, it is worth noting that motion picture films have a very complex structure, and the chemical mechanism underlying colour formation is also complex. For simplicity, the general stratigraphy of a film can be summarised as follows: a layer of polymeric support (the thickest one), typically cellulose triacetate or polyester; an image layer consisting of a photosensitive gelatin emulsion, which contains the substances responsible for the colours; an anti-halo undercoat and a final coat for protective purposes. In the case of black-and-white images, the hues are given by silver halide crystals which turn into silver during processing. Referring instead to the colour images, the emulsion is divided into three layers (each 5–10 µm thick), alternating with as many filtering layers and each containing magenta, yellow, or cyan dye. Together,
they determine the final colouring. In such systems, the three subtractive primary dyes are generated during processing. Prior to this process, each colour record stores information in the form of a ‘latent’ silver image that catalyses the formation of the visible dyes. Reddish light at 600–700 nm gives a cyan dye, greenish light at 500–600 nm gives a magenta dye and bluish light at 400–500 nm gives a yellow dye [4].

The colour formation process is quite complex, as the dyes are formed “in situ” during the processing itself. In particular, the colour is given by the reaction between an oxidized developer, which is generally a derivative of $p$-phenylenediamine, and a dye-forming material or “coupler” to obtain magenta, cyan, or yellow colouring compounds.

A wide variety of couplers can be used, although manufacturers provide very little information on their precise nature. Generally, in the chromogenic films the magenta and yellow dyes belong to the family of azomethines, while the cyan dyes belong to that of indoanilines. In particular, the most important classes of yellow-forming couplers are $\alpha$-benzoylacetanilides and $\alpha$-pivaloylacetanilides. For magenta-forming couplers the main classes are 5-pyrazolone, indazolone, pyrazolobenzimidazole, and pyrazolotriazole. Finally, all the couplers that form cyan are of the families of phenol or naphthol, subsequently joined by 2-aryl-5-(arylamido)phenols due to their higher stability and superior colour properties [2,5] and even more recently by pyrrolotriazoles and pyrazolotriazoles [4].

Although extensive studies have reported on the performances and the degradation processes affecting these dyes [2,5], as mentioned above, the analytical detection of the molecules responsible for colour in a film is still an open challenge. In this work three dyes, representative of the most common classes in chromogenic films, were synthesised and characterised. In particular, analytical methods based on Raman spectroscopy were investigated as a means of identifying such dyes. Raman spectroscopy is a common tool in the field of diagnostics of cultural heritage materials, thanks to its specificity and potential non-invasiveness. However, when applied for the detection of organic compounds, it suffers from the problem of fluorescence emission associated with these molecules. Recently, the development of the innovative SSE™ (Sequentially Shifted Excitation) strategy has made it possible to solve, or at least limit, this problem [6]. On the other hand, surface-enhanced Raman spectroscopy (SERS) allows one to take advantage of the increased Raman signal of an analyte, when it is absorbed, or in proximity of, a rough metal surface. Thanks to this close contact, Raman scattering excited by visible light can experience an enhancement of up to a factor of $10^8$ [7] allowing the detection of analytes even at very low concentration. Furthermore, metal nanoparticles absorb most of the fluorescence photons emitted by molecules, acting as a quencher of the fluorescence emission often associated with organic substances. Therefore, SERS spectroscopy has been extensively exploited in the field of cultural heritage diagnostics for the detection of natural and synthetic dyes in fabrics and lakes. [8–12] The most typical SERS substrates are colloidal suspensions of metal nanoparticles, as they can be prepared quickly and easily used. This traditional approach involves the analysis in solution of the target analyte. Typically, the materials chosen for this purpose are spherical silver nanoparticles, prepared as a colloidal dispersion by the Lee–Meisel method [13]. More recently, it has been demonstrated that nanomaterials with anisotropic structure have shown greater efficiency in producing significant SERS enhancements [14–17]. For this reason, a silver nanostar colloid was used as SERS substrate, in the present work.

In the present work, three dyes have been synthesised as references, one for each of the three colours magenta, yellow, and cyan. In particular, a benzoylacetanilide, a pyrazolone, and a substituted phenol were selected as couplers for the yellow, magenta, and cyan dye, respectively. $N,N$-diethylbenzene-1,4-diamine was used to reproduce the role of the developer [18]. The synthesised compounds have, to the best of our knowledge, been successfully studied for the first time using Raman and SERS techniques and the results have been exploited to characterise dyes extracted from a real motion picture film. Preliminary results were promising, particularly those obtained by SERS spectroscopy, revealing a new potential application of this technique in the field of conservation science.
2. Materials and Methods

2.1. Synthesis of Dyes

The dyes used as references for yellow (1), magenta (2), and cyan (3) colour (Figure 1) were prepared according to the procedures reported in the literature [18–22], as described in the Supporting Information (see the Word file “Synthesis and NMR characterisation of the reference dyes”). The molecular structures of the synthesized compounds were checked by recording their $^1$H and $^{13}$C NMR spectra (Word file “Synthesis and NMR characterisation of the reference dyes” Supporting Information).

![Figure 1. Structure of the dyes used as reference for yellow (1), magenta (2) and cyan (3) colour.](image)

2.2. Synthesis of Silver Nanostars

Silver nanostars were synthesised according to the procedure reported by Garcia-Leis [16]. Briefly, 500 µL of a 0.05 M NaOH solution was stirred at 670 rpm for one minute with 500 µL of 50% w/v hydroxylamine solution. A total 9 mL of a $10^{-3}$ M AgNO$_3$ solution was then added and the mixture was stirred for 5 min. Finally, 100 µL of a 1% trisodium citrate solution (0.045 M) was dropped into the flask and stirring continued for 15 min. The resulting dark green colloid was stored overnight in the dark before use to allow the nanoparticles to develop their final branched shape. The successful synthesis of star-shaped nanoparticles was verified by UV-Visible spectroscopy. An absorption band at 380 nm, caused by the central core, was followed by a progressive increase in the NIR region (from about 700 nm) produced by the several tips (Figure S1 Supporting Information) [17].

2.3. Sample Preparation

As a real case study, a 35 mm Fujicolor Positive Film Eterna-CP 3513DI was examined, containing the trailer for the animated film Happy Feet (Warner Bros, Burbank, CA, USA, 2006). In short, this film consists of a polyethylene terephthalate (PET) support, on which there are the three layers of light-sensitive emulsion, containing from top to bottom the magenta, cyan, and yellow dyes, isolated by suitable separatory layers and covered with a final protective coating [23].

For the analyses, the surface coating was removed, and the underlying part of the film was scratched with a scalpel in order to separate from the support all the layers of emulsion containing the colourants. The sample obtained was dispersed in a 3:1 of ethanol/water solution and the dyes were extracted under magnetic stirring for 2 days. The suspension was then centrifuged for 20 min at 6000 rpm and the supernatant was collected, leaving the solid residue at the bottom of the centrifuge tube. The dyes were separated by thin layer chromatography (TLC). An ALUGRAM(R) Xtra SIL G/UV plate with a silica stationary phase was used, while the mobile phase was a mixture of diethyl ether and hexane (ratio 1:1). After separation, the spots corresponding to each dye (Figure S2 Supporting Information) were scratched away from the plate and the analytes were extracted from silica by sonication in ethanol. The extraction procedure was used because in a previous attempt of a TLC-SERS analysis, carried out by depositing few microlitres of pre-concentrated nanostars on the spots [24], the signals due to the colloid were the predominant ones. It should be noted that only one spot was obtained for magenta and yellow dyes respectively, while two cyan-coloured spots were observed. In principle, more than one dye for a given colour may be contained in the film and degradation products can also be observed in an aged film [2]. However, the latter hypothesis can probably be
discarded as cyan dyes are the most stable and the degradation products formed by their residual couplers are generally colourless [2]. Of the two cyan spots, the extract obtained from the one better separated from those due to the other dyes was examined by SERS.

2.4. SERS and Raman Analysis

SERS analyses were performed on the reference dyes in ethanolic solution (10 mM) or on the extracts obtained from the film. A total 200 µL of silver nanostars were stirred on a magnetic plate, then 10 µL of the dye solution was added. Although the activation by aggregation is not necessary for these nanoparticles due to the large enhancement provided by their special morphology, it was found that more intense SERS spectra are obtained if 5 µL of sodium perchlorate (1.8 M) is added to the nanoparticles before the analyte. A micro-drop of the final suspension was deposited on a glass slide and SERS analyses were performed using a JASCO RMP 100 portable Raman micro-probe. The micro-probe is equipped with an 1800 lines/mm grating, a notch filter, an Olympus 50× microscope objective and a Peltier-cooled CCD detector. A Nd:YAG laser delivered the exciting radiation at 532 nm, with an output laser power of approximately 100 mW. All SERS spectra were recorded between 2000 and 200 cm$^{-1}$ as a sum of 30 scans with an exposure time of 2 s.

Normal Raman analyses were performed directly on the powdered dye for the reference compounds. A Bruker BRAVO handheld spectrometer was used for the measurements. This instrument is based on the patented SSE™ technology, which involves the excitation of spectra by means of two diode lasers operating at different temperature and emitting respectively at 785 and 850 nm. A suitable algorithm, therefore, allows to extract the final Raman spectral data. The spectra are collected in two sequential steps, from 300 cm$^{-1}$ to 2000 cm$^{-1}$ and from 2000 cm$^{-1}$ to 3200 cm$^{-1}$. The average spectral resolution is approximately 11 cm$^{-1}$ and the applied laser power is always less than 100 mW for both lasers. The acquisition time varies from 500 ms to 2 s and the number of accumulations from 5 to 300 (these parameters are automatically set by the instrument).

3. Results and Discussion

The dyes synthesised as references were initially studied. The Raman spectra show peculiar pattern for each dye, characterised by intense and well-defined signals (Figure 2). The SSE™ technology, in fact, allows to strongly reduce the fluorescence background associated with these organic molecules.

Satisfying results were obtained also by SERS spectroscopy upon visible excitation at 532 nm (Figure 3). The interaction between the dye molecules and the silver nanostars, in fact, leads to the enhancement of the Raman scattering and to the quenching of the fluorescence emission. As expected, a correspondence was found between SERS and Raman spectra of the same dyes, despite small shifts of band positions and variations in the intensity ratios (Figure S3 Supporting Information). Tentative assignment of Raman and SERS bands are reported in Table 1. The SSE™ technology, in fact, allows to strongly reduce the fluorescence background associated with these organic molecules.

The azomethine yellow dyes belonging to the class of the compound synthesized in the present work are known to have generally less planar molecular structures [4]. In particular, for a molecule similar to 1, except for a methyl group on the p-diethylaminophenyl ring and a chlorine atom on the phenyl ring of the carbamoyl moiety, crystallographic studies have shown that the p-aminophenyl fragment is coplanar with the azomethine-carbamoyl fragment, but is perpendicular to the acyl one [27]. Consistently, in the SERS spectrum
of the synthesized yellow dye the strongest bands are again due to vibrational modes of C=C bonds, but a $\beta$(C-H) band with medium intensity is still observed at 1172 cm$^{-1}$ (Table 1). This fact, coupled with the unshifted position of the carbonyl band, suggests that the interaction still takes place through the $\pi$ system, but with a tilted orientation of the molecule relative to the silver nanoparticles. A similar situation probably takes place for the indigoaniline cyan dye 3, for whose molecule a dihedral angle of 47$^\circ$ has been calculated between the quinone and the aniline rings [28]. In the corresponding SERS spectrum, the band due to the C=O group of indigoaniline at 1630 cm$^{-1}$ is still intense, but does not show any shift possibly associated with an interaction with the silver nanoparticles through the lone pairs of the oxygen atom. Furthermore, two particularly strong bands are located at 791 and 433 cm$^{-1}$ and have been tentatively assigned to the 10a and 16a modes of the diethylaniline ring (Table 1), both corresponding to out-of-plane vibrational modes in the Wilson notation [29]. On the other hand, a band possibly associated to the in-plane 9a mode also appears with medium intensity at 1191 cm$^{-1}$. In summary, an interaction through the double-bond system, with a tilted orientation of the molecule, seems the most probable also for the cyan dye, despite the fact that the $\pi$-conjugation is limited due to the lack of perfect planarity.

Figure 2. SSE™ Raman spectra of pure powdered (a) cyan, (b) yellow and (c) magenta dyes.
Figure 3. SERS spectra of (a) cyan, (b) yellow, (c) magenta, and (d) a mixture 1:1:1 of the three dyes. Spectra were obtained from 10 mM ethanolic solutions.

Table 1. Tentative assignment of Raman and SERS bands for the reference dyes magenta, yellow, and cyan.

| Raman       | SERS       | Tentative Assignment                      |
|-------------|------------|-------------------------------------------|
| Magenta [30–33] |            |                                           |
| 1699 w      | -          | ν (C=O) amide                               |
| 1667 w      | 1667 m     | ν(C=O) pyrazolone + ν(C=N)                  |
| 1624 m      | 1624 s     | ν(C=C) + ν(C=N)                             |
| 1473 vs     | 1484 mw    | ν(C=O)                                     |
| 1411 m      | -          | β(C-H)                                     |
| 1325 m      | 1325 ms    | pyrazolone ring stretching + β(C-H)        |
| 1230 m      | -          | β(C-H)                                     |
| -           | 1194 mw    | β(C-H) ring + ν(C=C) + β(CNN)              |
| 1110 m      | -          | β(C-H)                                     |
| 1076 m      | -          | β(C-H)                                     |
| 883 mw      | -          | γ(C-H)                                     |
| -           | 840 mw     | γ(C-H)                                     |
| 765 m       | 765 mw     | ring deformation                            |
| 638 w       | 638 mw     | aromatic and heterocyclic ring deformation  |
| 595 mw      | 595 mw     | aromatic and heterocyclic ring deformation  |

Yellow [30,34]
Table 1. Cont.

| Raman       | SERS       | Tentative Assignment                  |
|-------------|------------|---------------------------------------|
| 1670, 1660 mw | 1670, 1660 sh | ν(C=O) amide, ν(C=N)                   |
| 1619 w      | 1620 m     | ν(C=C)                                |
| 1555 vs     | 1550 ms, sh | ν(C=C) + β(N-H)                       |
| 1513 vs     | 1515 vs    | ν(C=C)                                |
| 1440 mw     | 1440 w     | ν(C=C)                                |
| 1311 mw     | 1316 w     | ν(C=C)                                |
| -           | 1284 mw     | -                                     |
| 1266 mw     | 1269 sh    | -                                     |
| 1214 m      | 1214 mw    | -                                     |
| 1170 s      | 1172 m     | β(C-H)                                |
| 1150 s      | 1151 sh    | β(C-H)                                |
| -           | 1110 w     | -                                     |
| 1075 w      | 1073 w     | β(C-H)                                |
| 950 mw      | 950 mw     | ν(C=C)                                |
| 828 w       | 828 w      | β(C=O) anilide?                       |
| -           | 712 m      | γ(N-H)?                               |

Cyan [30,35,36]

| Raman       | SERS       | Tentative Assignment                  |
|-------------|------------|---------------------------------------|
| 1683 w      | -          | ν(C=O) amide                           |
| 1628 m      | 1630 s     | ν(C=O) indoaniline                     |
| 1594 mw     | 1601 sh    | 8a a diethylaniline                    |
| -           | 1567 sh    | ν(C=C)                                |
| 1511 sh     | 1514 w     | ν(C=N)                                |
| 1464 vs     | 1464 m     | ν(C=C)?                               |
| 1401 m      | 1401 vw    | δ(C-H) ethyl?                         |
| 1352 mw     | -          | δ(C-H) ethyl?                         |
| -           | 1340 mw    | ν(C-N)                                |
| 1321 m      | 1327 mw    | -                                     |
| -           | 1191 m     | 9a a diethylaniline?                   |
| 1150 s      | 1153 m     | ν(C-N)                                |
| -           | 791 ms     | 10 a diethylaniline?                   |
| 428 w       | 433 vs     | 16a a diethylaniline?                  |

* Numbers refer to the Wilson notation.

It is worth noting that, on the other hand, normal Raman spectroscopy with visible excitation (532 nm) did not lead to significant results as almost no signals due to the any of the three dyes could be recognised, and only those due to the glass slide were observed (Figure S4 Supporting Information).

In addition, a 1:1:1 mixture of the three dyes was analysed. The acquired SERS spectrum (Figure 3d), although the relative concentration is the same for each dye, is dominated by the signals due to magenta, and only minor features are attributable to yellow and cyan dyes. This is probably due to the fact that the red hue of this dye fulfils the resonance conditions with the green laser beam.

SERS analysis was then applied to the dyes extracted from the film, before and after TLC separation. It should be noted that the low amount of dye extracted for each colour prevented the acquisition of SSE™ spectra, as the instrumentation used did not allow a microanalysis. On the other hand, SSE™ measurements were also performed directly on differently coloured frames of the film (results not shown), but of course the obtained spectra were complicated by the bands due to the polymer and by the superposition of bands due to the different dyes. As shown in Figure 4, good-quality SERS spectra were obtained from each dye and were compared with those of the reference compounds.
The similarity between the spectral patterns suggests that the dyes in the film belong to the same chemical class as the reference ones, at least as regards the core of the molecules. On the other hand, it should be remembered that the true dyes in the chromogenic films are much larger molecules than the reference ones. This is due to the presence of substituents that have the role of “ballast”, i.e., useful to retain the couplers in their coating layer [4,5] and to avoid their migration in other layers. Such substituents can be for example hydrophobic groups or polymeric chains [2].

Thanks to its selectivity towards the conjugated core of the molecules, SERS spectroscopy can be more effective in the classification of these dyes compared to other molecular analysis techniques, such as infrared spectroscopy. This information is relevant because, in any case, most of the degradation processes that must be considered in the conservation and restoration of the films are mainly related to the molecular classes to which the couplers belong.

4. Conclusions

The present work aimed at a preliminary investigation on the possibility of applying SERS spectroscopy to the classification of dyes in chromogenic films, to obtain information that may be used for the purposes of conservation and digital restoration.
Reference dyes deriving from three of the most common classes of couplers used in such films were synthesized and characterized for the first time by normal Raman and SERS spectroscopy. Interpretation of SERS spectra suggested that the main interaction between the molecules and the silver nanoparticles occurred through the conjugated π system.

The comparison between the SERS spectra of the reference compounds and those of dyes extracted from a motion picture film and separated by TLC indicated that such dyes most likely belong to the same chemical classes as the reference ones. SERS spectroscopy can therefore be proposed as a promising method to individuate these classes, each having peculiar features from the point of view of degradation mechanisms.

Naturally, this preliminary study should be extended to a wider number of reference compounds, representative of other types of couplers, and of dyes extracted from films of different brands and datable to different periods.

**Supplementary Materials:** The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/chemosensors10030101/s1, Synthesis and NMR characterisation of the reference dyes; Figure S1: UV-visible spectrum of the silver nanostar colloid; Figure S2: TLC separation of the extract from the emulsion obtained from the Fujicolour positive film; Figure S3: Comparison between SERS (black line) and SSE™ Raman (grey line) spectra of (a) cyan, (b) yellow and (c) magenta reference dyes. The SERS spectra were obtained from 10 mM ethanolic solutions, the Raman ones from the pure powder dyes. Figure S4: Normal Raman spectra (λexc = 532 nm) obtained from ethanolic solution (10 mM) of the (a) cyan, (b) yellow and (c) magenta reference dyes. * = signals due to glass.

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