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Non-invasive characterisation of molecular diffusion of agent into turbid matrix using micro-SORS

This st udy proposes a non-invasive analytical method to study the molecular diffusion of a chemical agent into a turbid matrix with an emerging analytical technique, micro-Spati ally Offse t Raman Spectroscopy (micro-SORS). Here, the micro-SORS concept has been extended from the analysis of chemically distinct stra tified layers to the studies and monitoring of the absorption and diffu sion processes,

1. Introduction

The stud y of the diffusion of an agent into a tutbid matrix is a fundamental topic in materiai science since the concentration profile and the penetration depth of the agent severely affect the final properties of the system. Plastics, papers, concretes and biologica! tissues can be affected by absorption or alteration processes with the consequent formation of concentration profiles, which have to be characterized for the in depth understanding of their performance . In Cultura! Heritage field, the knowledge of the depth distribution of a product applied on the historical buildings or art objects surfaces for conservation purposes is crucial for assessing the efficacy of consolidation and protection treatments [1]; another important issue concems the correct assessment of solvent diffusion and retention inside paint layers, for example, in evaluating new cleaning procedures. In fact, it is widely known that the use of not confined solvents for the

addr essing a key analytical need in a number of areas including polymer, pharma ceutical, forensic and biomedica! sciences.

In Cultura! Heritage the knowledge of the penet ration depth of a polymer use d to consolidate orto protect an object, or the absorption depth of solven ts used during a cleaning procedure is crucial for the performance evaluation of restoration methods and their safety towards the work of art. To date the most common protocol for obtaining this type of information comprises the application of stratigraphical analysis on cross-sections prepared after taking a small amount of sa mple from the work of art. This approach is destructive and may Ja ck of statistica! mean ing, since the analytical inform ation is limit ed to the micro area of samplin g. To overcome these d rawbacks, in this s tudy micro-SORS was successfull y use d, for the first time, to non-invasively characterise the penetration of a polymer and of a viscous solvent into a gypsum substrate, permi tting the reconstruction of the diffusion trends of the products into the matrix and the evaluation of their performances.

cleaning of paintings may produce negative effects such as swelling, soft ening and thus affecting the optical and mechanical properties of the paint layers [2]. Finally the diffusion of the decay products, as the solubl e salts, within the substrates (stone, plaster, stucco, etc.), brings essential information on the extent of the decay processes required, for instance, for planning more suitable restoration works [3].

To date, the main analytical techniques used in Cultura(Heritage field for investi gating the diffusion of chemical agents and decay products in the substrates includ e scanning elec tron microscopy coupled with energy dispersive spectrometry (SEM-EDS) [1,3], Fourier-transform infrared spectroscopy (FTIR) [4,5] and Raman spectroscopy [1,6]. These are currently destructive in nature due to their need of resorting to cross sectional analysis. Recently, alternative non-des tructive techniques such as neutron im aging [7], X-ray micro-computed tomography [8] or synchrotron diffraction imaging have been proposed for the same purpose; however, these are not easily accessible tools and the direct

retrieval of molecular composition of organic, inorganic, crystalline and amorphous compounds, which is extremely important for their unequivocai identification, is not fully accomplished by these techniques.

The nove! aspect of the present study is the development of a new non-invasive analytical method to study, at the molecular leve!, the diffusion of a chemical agent into a turbid matrix, using micro-Spatially Offset Raman Spectroscopy (micro-SORS) [9,10], an emerging, accessible and relatively easily deployable technique.

Micro-SORS is a conceptual evolution of its parent technique, (macro-scale) SORS [11,12], combining SORS with microscopy. This enables to non-invasively resolve thin, micrometre scale layers such as painted stratigraphy in the Cultura! Heritage field or turbid stratified systems in polymer, catalytic, biologica! and biomedica! sciences. Micro-SORS represents an important tool in situations where a nondestructive and/or non-invasive molecular analysis is required, e.g. when dealing with precious, unique samples or objects in art field and forensic science.

Micro-SORS principles and associated concepts have been discussed extensively in literature recently [13-23].

To date micro-SORS has been used to discriminate between the composition of different layers in stratified matrixes. Here, for the first time, micro-SORS has been optimized to monitor the diffusion of an agent into a turbid matrix, where there is no defined separation of compounds into chemically distinct Jayers . This proof of concept study demonstrates that defocusing, the most basic micro-SORS var iant , is capable to non-invasively distinguish different penetration depths of conservation products and solvents into a matrix, through the monitoring of their diffusion profiles. Two situations have been reproduced using Jab oratory mock-up samples which have a substrate made of gypsum (i.e. simulating a gypsum-based stucco matrix) but different applied products, namely an organic conservation product (Paraloid B72 - PB72), and DES, a deep eutectic solvent (Choline chloride-Urea 1:2). PB72 is an acrylic copolymer that, since the 1950, has been extensively applied as fixative and protective especially on wood, ceramic, glass, stone monuments and plasters [24-27]. Choline chloride-Urea 1:2 DES and its based gels are a new type of eco-friendly products [28] that are currently under evaluation for the removal of proteinaceous coatings from art objects.

2. Materials and methods

In the experiments, a matrix made of gypsum blocks (5 cm x 5 cm x 2 cm) was used for the application of the two products.

The first product was Paraloid B72 (PB72), an ethyl-methacrylate and methyl-acrylate copolymer commonly used as fixative, having good ageing proprieties and chemical stability [25,26]. The solution of Paraloid B72 (5% in acetone) was applied with a brush on the gypsum blocks surface. Two samples, S1 and S2, have been treated with an increasing amount of PB72 to obtain a different penetration depth of the product.

The second product was a deep eutectic solvent (DES), an innovative and eco-friendly product, having similar physical-chemical properties to common ionie liquids (!Ls) and severa! advantages over traditional ILs such as Jower price, easy storage and easy preparation [29]. Moreover, its efficacy for the extraction of proteins has been demonstrated [30], unlocking the possibility of using it as a solvent in new green gels for the removal of proteinaceous coating from paintings. Nonetheless, DES is a non-volatile solvent and thus it is retained in a (porous) materiai, possibly altering its physical-chemical and optical proprieties. It is therefore extremelyimportant to evaluate the extent of the diffusion of this product in an artwork.

DES solve nt was synthesized by mixing a hydrogen bond acceptor (Choline chloride) with a hydrogen bonding donor (Urea), at the mole ratio 1:2 in a round bottom flask. The mixture was stirred at 100 °C for 5 min unti!a homogeneous colorless liquid was obtained. A drop of DES was applied on the S3 gypsum surface and its diffusion was studied with

micro-SORS measurements.

A DES drop exceeded the quantity of solvent that could penetrate in a matrix during a cleaning procedure. Such relatively great amount was applied to enhance the DES Raman signal and thus to enable a better monitoring of its absorption process with micro-SORS. Further studies on a more realistic scenario are in progress, mimicking the standard cleaning procedures.

The experiments were carried out using a Senterra dispersive Raman microscope (Bruker Optik GmbH) equipped with a Peltier cooled charge coupled device (CCD) detector (1024 x 256 pixels) and a 785 nm excitation laser.

Defocusing micro-SORS measurements were carried out on the intact samples after their interaction with the products. The spectra were collected using a laser power of 100 mW and an acquisition time ranging from 100 to 300 s. The measurements were carried out at imaged and defocused positions moving the objective (20 x magnification,NA 0.4 and WD 1.3 mm) away from the sample in z direction ($\delta z = 100$, 200, 300, 400, 500, 700 and 1000 µm). The theoretical laser spot diameter at imaged position is 2.4 µm.

Three series of defocusing micro-SORS measurements were acquired on the samples S1 and S2. As for S3, the diffusion of DES was studied by performing a series of three micro-SORS acquisitions at t_0 (immediately after the application of the drop) and three additional series at t_1 (after 30 h) to evaluate the behavior of such an oily compound over time.

The progressive decrease of PB72 and DES signals with respect to gypsum was evaluated calculating the intensity ratio between two selected Raman bands of the compounds. The decreasing trends observed for the three series of each sample are reproducible; slight differences observed among them have to be ascribed to the heterogeneity of the samples. To provide an easy data visualization, the intensity values obtained for the three series were averaged to geta representative trend for each sample (the single series are reported in the supplementary materiai).

Moreover, the cross-sections of Sl and S2 were prepared cutting and polishing a fragment of the samples. The cross-sections were conventionally analysed with vertical linear scans to check the penetration depth of PB72 and the outcomes were compared with the results obtained with defocusing micro-SOR.SThe spectra were acquired using a 20 x magnification objective, a laser power of 100 mW at the sample and an acquisition time of 100 s per spectrurn. Three line scans were performed, starting from the surface and progressively moving to the internal part of the samples, with a step size of 50- 100 μ m. Por each depths, the PB72/ gypsum intensity ratio was calculated and averaged for the three line scans (in the supplementary materiai, the single lines scans are reported).

The S3 actual penetration depth was not checked with the samples cross-section, since DES remains in a liquid state in the gypsum matrix, and therefore can move from its original position during the sample cutting and polishing.

Calculations and plot construction were carried out using OPUS and ORIGN software.

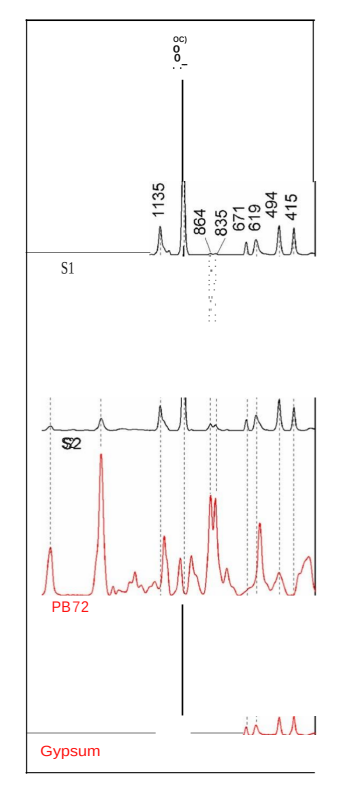
3. Results and discussion

3.1. Paraloid B72 on gypsum (S1 and S2)

The conventional Raman spectra acquired on Sl and S2 surfaces show the presence of both gypsum and PB72 (Fig. 1). Gypsum characteristic bands are visible at 1135, 1008, 670, 620, 495 and 415 cm⁻¹ PB72 bands, Jocated at 1725, 1450, 864 and 835 cm⁺¹ are much Jess intense compared with the gypsum, and this is consistent with the relatively Jow amount of PB72 in the samples.

As expected in S2, the PB72 Raman bands intensity is higher than in S1 because more product was applied on S2 surface (Fig.1).

The penetration depth of PB72 and its diffusion profile were studied with both conventional destructive Raman analyses on the sample



1800 1600 1400 1200 1000 800 600 400

Wavenumbers (cm·¹)

Fig. 1. Raman spectra (black) collected on SI and S2 surfaces at imaged position (corresponding to conventional Raman measurements), normalized to the most intense gypsum band at 1008 cm⁻¹. The reference spectra of PB72 and gypsum are also reported (red). (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

cross-sections and with non-invasive defocusing micro-SORS measurements directly on the mock-ups surfaces. With the conventional measurements on cross-section, PB72 Raman signal was detected up to 250 µm of depth in S1 and up to 500 r1m of depth in S2. To visualize the decrease of the PB72 signal with depth, the PB72/ gypsum ratios (intensity ratio of two representative bands) were calculated and plotted against the defocusing dista nce. The two bands chosen for ratio calculation are the 1450 cm⁻¹ band of PB72 (the most intense of the compound, assigned to one of the C-H bending modes [31]) and the 415 cm band of gypsum (SO₄ symmetr ic bending [32]). The latter was selected for its intensity, as it is the most comparable to the one of PB72 ensuring easie r monitoring of the relative intensity changes with micro-SORS defocusing distances. The averaged ratio values obtained in S2 cross section are substantially higher than in S1 along the vertical linear scans, and we can therefore assume that in S2 there is considerably more PB72 compared to S1 at all measured depths (Fig. 2a). To compare the diffusion profiles of the two samples, the averaged ratio were normalized (Fig. 2b; see supplementary materiai, Figure S1 and Figure S2, for the single series): two different intensit ies ratio decay rates are observed, and the decrease of the PB72/ gypsum ratio is faster in Sl.

In the defocusing series, a progressive decrease of PB72 bands is observed in both S1 and S2 relative to those of gypsum. In Fig. 3, a defocusing series collected on S2 sample is reported to illustrate this trend. It can be observed that the Raman bands position slightly shifts to higher wavenumbers with the increasing of the defocused distance and this effect is likely to be an artefact of the instrumental setup, since the shift affects the entire spectrum. The PB72/gypsum ratios (intensity ratio of two representative bands, namely bands at 1450 and 415 cm⁻¹ for PB72 and gypsum, respectively) were calculated, normalized and plotted against the defocusing distance to highlight the trends obtained with the two samples.

The averaged ratio plot obtained with defocusing measurements in the intact samples (Fig. 4a) shows higher ratio values for S2, similarly

to the cross-section plot. The information obtained with the two methods is consistent and points out to a higher quantity of PB72 in S2. The plot in Fig. 4b shows the normalization of the averaged ratio of

SI and S2. In the supplementary materiai, the plots of the three single series are reported (Figure S3 and Figure S4).

There is a distinct difference in the decay rates for Sl and S2 samples signals plotted as normalized intensity ratios: Sl shows a faster ratio decay rate, that is attributed to the more rapid lowering of the product concentration with depth. Therefore, the different decay rates can be related to the different penetration depth of the product in the two samples. In this case, the faster ratio decay rate in S1 suggests a more superficial penetration of PB72.

Defocusing and conventional measurements on cross-sections give consistent results, as with both methods the two samplesshow different ratio decay rates: the ratio values obtained on the cross-section decrease more rapidly in S1 than in S2, as was observed in the defocusing measurements. Nonetheless, there is a differe nce in the slope of the curve obtained with the two methods. This happens because in defocusing Raman spectra the contribution of the surface is always present even at large defocusing distances: the photon generated at (and dose to) the surface are collected together with the Raman photons generated from the inner portion of the materiai. In contrast, in conventional backscattering Raman measurements only the Raman photons generated at (and dose to) the surface are predominantly collected. The persistence of the surface Raman photons in the defocusingspectra is also the reason why the defocusing surface/ subsurface ratio decay rate is not expected to fai! to zero (unless there is a drop of the overall signal intensity).

Moreover, in S2 cross-section, anomalous constant intensities ratio values are observed between 25 r1m and 100 μ m (Fig. 2), p ro vidi ng

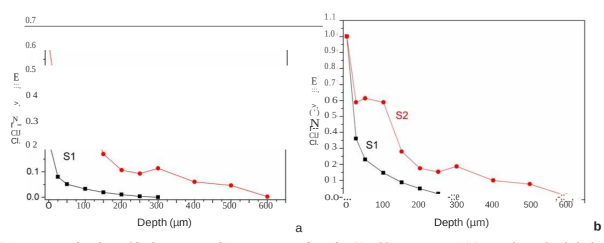


Fig. 2. PB72/gypsum ratio plots obtained for the conventional Raman spectra performed on SI and S2 cross-sections. (a) Averaged ratio plot: the higher ratio values observed in S2 are related to a more abundance of the produci within the sample; (b) averaged and normalized ratio plot: two different intensities ratio decay rates are observed.

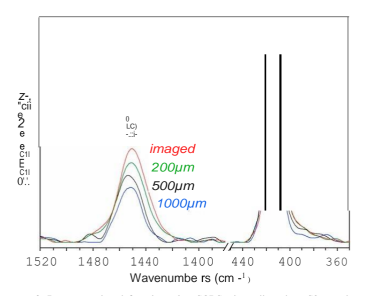


Fig. 3. Represen tative defocusing micro-SORSseries collected on S2 sample and normalized to 415 cm-¹ band of gypsum. The micro-SORS defocusing steps are indicated (values typed in different colours). A progressive relative decrease of PB72 Rama n band intensity is observed with increasing defocusingdistance. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

interesting information on the diffusion of the product which is probably affected by the heterogeneous porosity of the matrix. None theless, in line with expectations, this trend does not emerge in the defocusing plot (Fig. 4), because in defocusing measurements the laser spot and collection zones are enlarged, and the Raman signal is collected from a volume that is larger than the conventional measurements. Therefore, defocusing intrinsically averages the Rama n signal and some heterogeneities under the surface can be missed.

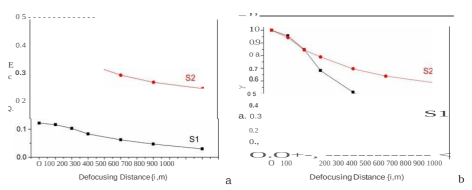
3.2. DES on gypsum (S3)

In this sample, the micro-SORS measurements aim at monitoring the diffusion of DES into gypsurn matrix in time resolved manner. Therefore, two sets of defocusing micro-SORS measurements were performed. The first set was performed immediately after DES application on sample surface (t_0). Afterwards, a series of conventional spectra were repeated on the sample surface: a progressive lowering of DES bands was detected, suggesting that DES was diffusing into the gypsum matrix. In the range between 20 and 30 h after the application DES band intensity was constant, implying the end of the absorption process. The second set of measurements was then performed at 30 h (ti).

Twomaged spectra collected at to and t_1 are shown in Fig. 5. The spectra are dominated by gypsum Raman bands, but some of the bands ascribable to DES are also visible: the band at 716 cm $_i$ ass igned to the C-N symmetric stretching of choline and the C-N asymmetric stretching band of urea located at 1450 cm 1 [33]. At t_1 , the imaged spectrum shows less intense DES Raman bands because DES has leac hed away from the surface. Upon performing micro-SORS experiments, it became evident that DES bands intensities progressively decreased compared with gypsum ones as is illustrated in Fig. 6.

For this sample, the 716 cm $_{i}$ band of DES and the 670 cm 1 Raman band of gypsum (S0 $_{4}$ asymmetric bending [32]) were used for the calculation of the solvent-substrate ratio. These Raman bands were

E PB/2/ gypsum ratio plots obtained for the defocusing micro-SORSspectra performed on SI and S2 surface. (a) A veraged ratio plot: the higher ratio values observed in S2 are related to a more abundance of the produci within the sample; (b) averaged and normalized ratio plot: the two penetration depths can be distinguished through the different intensities ratio decay rates of the samples.



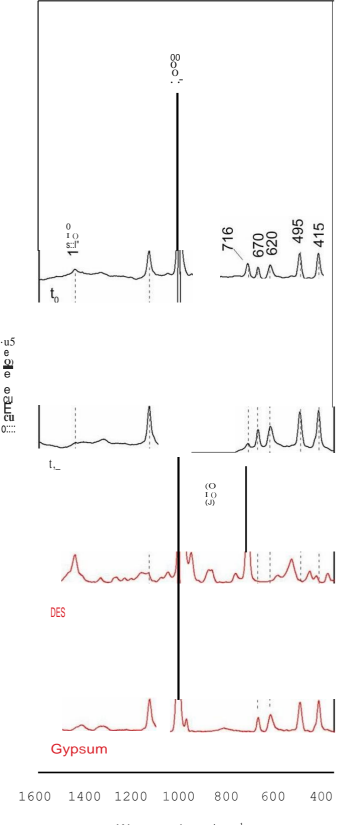




Fig. 5. Raman spectra obtained at imaged position on S3 surface at t,, and t_1 (black) and reference spectra of DES and gypsum (red). (For interpr etation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

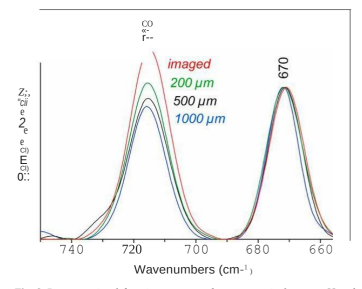


Fig. 6. Representative defocusing sequence of spectra acquired at t, on S3 and normalized to gypsurn band at 670 cm⁻¹. The micro-SORS steps are indicated (values typed in different colours). A progressive relative decrease of DES Raman band intensity is observed by increasing the defocusing distance. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

selected because of their intensities being comparable. The averaged ratio plots are shown in Fig. 7, whereas the ratio plots relative to the single series collected at to a nd t_1 are report ed in the supp le mentary materiai, Figure SS and Figure S6. The normalized Raman intensity ratio plots obtained at t_0 and t_1 show different decay rates, as expected (Fig. 7a): it is possible to observe that at t_1 the decrease of DES signals is less pronounced than at to. The rapid decrease of the ratio values at t_0 is associated to a higher concentration of the product dose to the surface, whereas the more constant ratio values obtained at t_1 are related to the deeper diffusion of the product inside the matrix.

It is noted that the ratio values of t_1 show a certain degree of fluctuation. This is ascribed to the Raman peak intensity of DESat t_1 being very low resulting in very small ratio values (see the not normalized ratio plot in Fig. 7b) and noisier outcome. For S3, a sample cross-section could not be analysed since DES, which remains liquid, can easily move from its original position during the sample cutting and polishing, rendering ineffect ive the monitoring of its distribution.

4. Conclusions

This study demonstrates the possibility of non-invasively assessing the concentration profile of a product within an absorbent matrix using micro-SORS. In micro-SORS experiments the samples with different penetration depths of a product could be differentiated based on the decay rates of their agent-matr ix Raman intensity ratio. The study is particularly beneficiai to the Cultura! Heritage field, as a non-invasive characterisation ensures avoiding damage to art materials. Using this method, a wide range of situat ions could be non-invasively studied, that include the penetration depth of a fixative agent or a consolidant in a stucco, a plaster or a stone, and monitoring of the absorption process of a liquid, such as a cleaning solvent, under the exposed surface.

The measurements were performed using the most basic variant of micro-SORS,name ly defocusing micro-SORS.This variant has the advantage of being compatible with existing micro-Raman instruments and its implementation does not require any instrumental nor software modifications. This work is therefore important by pointing out the

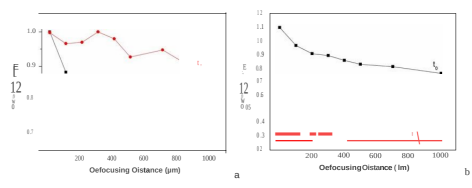


Fig. 7. (a) Normalized and (b) not normalized DES/gypsum averaged ratios of micro-SORS series of S3. The penetration depths at lo and t_1 are evidenced by different intensities ratio decay rates (a).

potential of this ready-to-use method for monitoring of absorption processes.

At this stage of the development, defocusing micro-SORS does not allow the determination of the absolute depth penetration reached by a product, instead it establishes if a product is more or is less deeply penetrated in a sample compared to another sample. None theless, these results suggest the possibility of estimating the absolute penetration depth of an agent into matrix after a calibration set is prepared and quantitative calibration established. The key requirement is, however, that the calibration set is built up having the same matrix and the same agent combination as that for the measured sample.