Silver Nanoflowers as an Interfacial Liquid-State Surface Enhanced Raman Spectroscopy (SERS) Sensor for Water Pollution

Zinah Salahuddin Shakir¹,²*, Ayad Abdul Razzak Dhaigham³, and Sameer Khudhur Yaseen⁴

¹Institute of Laser for Postgraduate Studies, University of Baghdad, Baghdad, Iraq.
²Department of Applied Sciences, University of Technology, Baghdad, Iraq.
³Directorate of Materials Research, Ministry of Science & Technology, Baghdad, Iraq.
⁴Department of Physics, College of Science for Women, University of Baghdad, Baghdad, Iraq.

*Email address of the Corresponding Author: zena.salalahdeen1101a@ilps.uobaghdad.edu.iq

Abstract: Water pollution has created a critical threat to the environment. A lot of research has been done recently to use surface-enhanced Raman spectroscopy (SERS) to detect multiple pollutants in water. This study aims to use Ag colloid nanoflowers as liquid SERS enhancer. Tri sodium phosphate (Na₃PO₄) was investigated as a pollutant using liquid SERS based on colloidal Ag nanoflowers. The chemical method was used to synthesize nanoflowers from silver ions. Atomic Force Microscope (AFM), Scanning Electron Microscope (SEM), and X-ray diffractometer (XRD) were employed to characterize the silver nanoflowers. This nanoflowers SERS action in detecting Na₃PO₄ was reported and analyzed concerning both shape and size using a 532 nm laser. We observed that the nanoflower's structure produced strong SERS signals. The increase in the SERS signal is related to the deposition of Na₃PO₄ molecules in the aggregated silver nanostructure in the solution. The concentration of Na₃PO₄ plays a main role in detection since the Raman signal becomes stronger as the concentration increases. The highest phosphate analytical enhancement factor obtained for SERS in colloidal nanoflowers was 1.7×10³ at 0.7×10⁻⁶ M which was the lowest concentration.

Keywords: AgNFs, Hotspot, Raman spectroscopy, Tri Sodium phosphate, Surface Plasmon resonance.

1. Introduction

Water pollution, one of the world's challenges, results from fast industrialization and poses a severe environmental danger. Organic and inorganic contaminants, like organic dyes, heavy metals, pesticides, sulfides, and so on, harm the aquatic system [1-3]. One of these pollutants that are dangerous to human health is fertilizers tri-sodium phosphate, which we will address in this research, as it can cause permanent
damage to human kidneys, leading to death [4]. Raman spectroscopy is a method that employs the vibrational spectrum to detect chemical interactions in a molecule and also being sensitive to surrounding changes [5]. As a consequence, Raman spectroscopy is a valuable tool for chemical and physical study.

The Raman spectrum offers information particular to a substance, allowing for molecular identification [6]. Furthermore, Raman spectroscopy necessitates a simple preparation of the sample [7]. Raman signals are typically weak, but they can be significantly increased by adsorbing molecules on a roughened surface of metal or metal nanoparticles. This technique is known as surface-enhanced Raman spectroscopy (SERS) [8]. SERS is considered an effective optical sensing process to detect various analytes. The SERS impact has been linked to an improvement in the effectiveness of Raman scattering for molecules that are present on or very close to the surface of specific metal nanostructures, particularly free-electron metals like gold, silver, and copper. As a result, chemically or physically adsorbed molecules on these metal nanostructures experience a considerable Raman signal increase. SERS is capable of detecting even a single molecule and achieving very large enhancement factors (more than $10^{10}$) with appropriate metal nanostructures and measurement settings [2,9]. The most frequently accepted explanations for SERS are the electromagnetic mechanism (EM) and the chemical mechanism (CM) [10,11]. EM has been reported to be more significant than CM [12]. EM is primarily based on surface plasmon in a metal nanostructure [13, 14]. In the EM process, electromagnetic waves interact with plasmonic metal nanostructures which leads to amplifying the Raman signal [15,16]. SERS has evolved into a valuable instrument for the quantitative study of hazardous compounds, even at low levels in food and the environment. SERS substrates are metal nanostructures that increase the Raman spectral signature of molecules deposited on them. SERS substrates can be of two different natures: colloidal nanoparticles in solution or nanostructures deposited on a surface [17]. It is commonly assumed that the significant Raman signal increase in SERS results from electromagnetic motivation at 'hot spots' on the SERS substrate [9,18,19]. These 'hot spots' are often nanogaps between neighboring metal nanostructures or narrow areas surrounding metal nanostructure tips [18,19]. Because of the number of 'hot spots,' experimental research and theoretical calculations have revealed that complex silver nanostructures including dendritic, flower-like, and star-like nanostructures can give an extremely high increase in the electromagnetic field [20-21].

This study will focus on the flower-like silver nanostructure as a liquid SERS substrate to demonstrate the detection of tri-sodium phosphate as a pollutant in water. Several research teams have created flower-like silver nanostructures known as silver nanoflowers-AgNFs to be used as SERS substrates [2].

2. Experimental work

2.1. Materials and methods

Tri-sodium phosphate (Na$_3$PO$_4$), and silver nitrate (AgNO$_3$) were purchased from CDH, India, and ascorbic acid (AA, C$_6$H$_8$O$_6$) tri-sodium citrate di-hydrate (TSC C$_6$H$_8$Na$_3$O$_7$ 2H$_2$O), polyvinylpyrrolidone (PVP) were purchased from SABIC, KSA, De-ionized (DI) water

2.2. Synthesis of silver nanoflower

First, an ice water bath arrangement was prepared by placing a 50 mL beaker holding 20 mL of distilled water in a 250 mL beaker having ice cubes. This configuration was exposed to 400 rpm magnetic stirring. After 10 minutes, a 0.5 M aqueous solution of AgNO$_3$ (2 mL) was added to the water. For another 10 minutes, the solution was treated with 0.3 M PVP aqueous solution (2 mL). The solution was then treated with 0.25 M tri-sodium citrate di-hydrate (0.2 mL) at 10-minute intervals. After 10 minutes, 0.5 M of AA (2 mL) was poured into the previously stated solution while stirring constantly [2]. A dark grey solution was obtained. A scanning electron microscope (SEM) AxiaChemiSEM by Thermo Scientific, the Netherlands, and an atomic force microscope (AFM) Model TT-2 AFM workshop, USA, were used to examine the surface morphologies and nanoparticle size distributions of the prepared Ag nanoflower. An
X-ray diffractometer (XRD) was applied to measure the crystallographic information of the Ag nanoflower in the range from 10 to 80. An ultraviolet-visible (UV-Vis) spectrophotometer was utilized to investigate the absorption spectrum of Na3PO4.

2.3. Preparation of Samples for SERS Spectra

Four samples of Na3PO4 were prepared at concentrations ($0.7 \times 10^{-3}$, $0.7 \times 10^{-4}$, $0.7 \times 10^{-5}$, and $0.7 \times 10^{-6}$) M in the Ag nanoflower colloidal. In order to make a comparison, we prepared a sample with a concentration of $0.7 \times 10^{-3}$ M of Na3PO4 in distilled water only. This bare sample will allow us to better understand the effects of the pollutant in question. SERS spectra samples were measured in a glass vial with the laser beam focused within. Raman scattering measurements were taken by using a Raman microscope (532 nm Preconfigured Raman Spectrometer System) by (Stellar Net, Inc. Florida, USA). Samples were stimulated by a 532 nm laser line generated by a laser with a power of 70 mW at the sample and an integration time of 9 ms. In all cases, the spectral resolution was adjusted to 2 cm$^{-1}$. SERS spectra were registered using a total acquisition time of 10s for each SERS spectrum and a single scan. All Raman measurements were taken in the spectral region (200 – 2000) cm$^{-1}$.

4. Results and Discussion

According to the SEM images in Fig.1, the size of silver nanoflower was approximately 400 nm. The nanoparticles were aggregated and formed hot spots which were thought to be the source of the substantial rise in signal strength required for single-molecule detection [22, 23].

According to the results obtained from the AFM, the average silver nanoparticle size was 13.61 nm; see Fig.2. The image shows that the nanosilver cluster formation in the topographic distribution is uniformly distributed. The test also revealed that the density of the silver nanoflower colloidal solution was 434 million particles/mm$^2$. The X-ray diffraction pattern of the AgNF structure exhibited unique diffraction peaks at (38.341°, 64.656°, and 77.625°), which are similar to pure silver crystal planes (111), (220), and (331); see Fig.3. These strong peaks in the planes suggest that the AgNFs are extremely crystalline. There were no further impurity peaks found, indicating that the samples were extremely pure. The crystalline size of Ag-NF was calculated from the Debye – Scherer equation and the average value of it was 43.6 nm [24].

$$L = \frac{k \lambda}{\beta \cos \theta_B}$$  \hspace{1cm} (1)
Where L, k, λ, β, and θ are the crystalline size, the shape factor value which equals 0.9, the wavelength of the X-ray in nm, full width at half maximum (FWHM) in radians, and the diffraction angle in radians respectively. It is clear from Fig.4 that the absorption spectrum of Na₃PO₄ is approximately 200 nm which makes the 532 nm laser used in the Raman measurement suitable hence it is not very close to the absorbance area that produces fluorescence.

**Fig.2:** AFM analysis: Granularity cumulation distribution histogram and AFM- 3 dimensions image of silver nanoflower.

**Fig. 3:** Typical XRD pattern of AgNFs that have been synthesized.
Fig. 4: The absorption spectrum of Na₃PO₄

Fig. 5: SERS intensity at the main peak (950 cm⁻¹) of Raman spectrum for Na₃PO₄ against the concentrations of Na₃PO₄ in Ag nanoflowers colloids.

The samples containing AgNFS showed greater Raman intensities at all Raman peaks than the bare ones; see Fig. 5. It has been linked to the massive hotspots generated by the sharp edges and nanogaps of AgNFS structure when subjected to laser light [2,25]. A significant local electromagnetic field is generated around the surface of the nanostructure which can be up to 100 times stronger than the incident one [26], resulting in a significant amplification of the Raman signal. This strong local electromagnetic field resulted from the excitation of localized surface plasmon (LSP), which correlates with collective oscillations of the cloud of electrons within nanostructures, or electromagnetic interaction between nanostructures. [27,28]. Based on previous research, it has been found that the greatest increase in hot spot enhancement occurs when the distance between nanoparticles is reduced and the particle diameter is smaller, especially when the surface curvature increases [29]. The electromagnetic coupling in the space between the nanoparticles is caused by...
the nanoflowers aggregated in the solution. Due to the adjacency effect, the enhancement with the aggregated nanoparticles is more intense than for individual ones owing to forming electromagnetic "hot spot" in the nanoparticle gap, which produces a strong SERS signal from the molecules around it [17].

![Figure 6: SERS intensity at the main peak (950 cm⁻¹) of Raman spectrum for Na₃PO₄ against the concentrations of Na₃PO₄ in Ag nanoflowers colloids.](Image)

The relationship between the concentrations of Na₃PO₄ and SERS intensity has shown in Fig.6. It is obvious that when Na₃PO₄ concentration increases, the SERS intensity increases too; this is attributed to the increase of the molecules of Na₃PO₄ attracted to AgNFs. Thus the Raman signal will be strengthened so the enhance Raman scattering strengthen which agree with articles [8, 30].

The analytical enhancement factor (AEF) was utilized to assess the actual SERS enhancement produced by EM and CM. AEF takes an analytical method for signal enhancement, combining signal strength with analyte concentration (C). This measure is useful when estimating the amount of analyte molecules present is difficult, especially for analytes that have no specific affinity for the plasmonic surfaces. AEF influences the vibration mode of a specific analyte in a normal Raman signal in the SERS technique, and it is directly proportional to the strength of the local electromagnetic field, referred to as a hot spot [31, 8].

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AEF = \frac{I_{\text{SERS}} \times C_{\text{NRS}}}{I_{\text{NRS}} \times C_{\text{SERS}}}
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Where; \(I_{\text{NRS}}\) and \(I_{\text{SERS}}\) refer to the counterpart intensities of normal Raman and SERS, respectively. \(C_{\text{NRS}}\) and \(C_{\text{SERS}}\) are the concentrations of the analyte in the normal Raman and SERS liquid substrates, respectively.

Based on our analysis, there appears to be a nonlinear increase of AEF at the highest peak (950 cm⁻¹) as the concentration of Na₃PO₄ decreases. This means that SERS is more efficient with lower concentrations. Overall, this information can help us better understand the effects of Na₃PO₄ on the samples we are analyzing; see Fig.7. Very low molecule concentrations improve the probability of specific molecule localization, which leads to better detection and hence raises the AEF by enhancing the SERS signal. At 0.7x10⁻⁶ M, the maximum phosphate AEF found for SERS in colloidal nanoflowers was 1.7x10⁵. The variation in (AEF) was mostly related to the variation in overlaps between SPR bands as a function of edge size and degree and the laser source. For ultrasensitive detection of phosphate molecules, the electromagnetic process of aggregated AgNF creates multiple hotspots from plentiful nanogaps on nanoflowers [29].
5. Conclusions

Tri sodium phosphate was examined as a pollutant utilizing liquid SERS based on colloidal silver nanoflowers. The structure of nanoflowers with sharp edges and corners produced strong SERS signals. The aggregated nanoflowers in the colloidal have generated hot spots. When Na$_3$PO$_4$ molecules are absorbed in solution and engage hot spots, the electromagnetic field increases, so the Raman signals are enhanced. The hot spot regions, which exist as minimal distances between nanoparticles, have an effect on the intense electromagnetic fields; SERS signals are greater within these microscopic hot spots. The AEF is directly proportional to the strength of the local electromagnetic field (a hot spot). The AEF influences the vibration mode of a specific analyte in a normal Raman signal in the SERS technique. As the Na$_3$PO$_4$ concentration increased, so did the Raman peaks. At 0.7x10$^{-6}$ M (the lowest concentration), the highest phosphate AEF for SERS in colloidal nanoflowers was 1.7x10$^3$ which means SERS is more efficient with lower concentrations.

References


حل محلل الفضة الغروي النانوي الشكل كمستشعر مطياف رامان المحسن للسطح للكشف عن ملوثات المياه

زيينة صلاح الدين شاكر1,2، إياد عبد الرزاق ضيغم3، سمير خضر ياسين4

1 معهد الليزر للدراسات العليا، جامعة بغداد، بغداد، العراق.
2 قسم العلوم التطبيقية، الجامعة التكنولوجية، بغداد، العراق.
3 دائرة بحوث المواد، وزارة العلوم والتكنولوجيا بغداد، العراق.
4 قسم الفيزياء، كلية العلوم للبنات، جامعة بغداد، بغداد، العراق.

البريد الإلكتروني للباحث: zena.salahaldeen1101a@ilps.uobaghdad.edu.iq

الخلاصة: قد خلق تلوث المياه تهديداً خطيراً للبيئة، وقد تم إجراء الكثير من الأبحاث مؤخراً لاستخدام مطيافية رامان المعززة (SERS) على السطح للكشف عن الملوثات المتعددة في المياه. تهدف هذه الدراسة إلى استخدام الفضة الغروية النانوية الزهرية ككاشف محسن السطح لتقنية رامان. تم فحص ثلاثي فوسفات الصوديوم (Na₃PO₄) باعتباره ملوثًا باستخدام SERS. تم بناء علي محلل الفضة الغروي الشكل بنفس الطريقة الكيميائية لتجميع الزهور النانوية من أيونات الفضة. تم استخدام المجهر الماسح الإلكتروني (SEM) والمجهر الماسح الذري (AFM) والمقياس حيود الأشعة السينية (XRD) لتصنيف الزهور النانوية الفضية. تم تقييم إجراء SERS في الكشف عن الفوسفات وتحليله فيما يتعلق بكل من الشكل والحجم باستخدام ليزر 532 نانومتر. لاحظنا أن البنية الزهرية للفضة النانوية أنتجت إشارات SERS قوية. ترتبط زيادة في إشارة SERS في البنية النانوية للفضة المتكتلة في المحلول بتركيز جزيئات Na₃PO₄ بشكل رئيسي. كان أعلى معامل تصحيح التحليلي للفوسفات الذي تم الحصول عليه لـ SERS في زهور النانو الغروية 1.7 × 10⁻⁶ مول. والذي كان أقل تركيز.