**In Situ SERS Detection of Ferbam on Spinach Surfaces**

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**Abstract.** In this study an in situ surface enhanced Raman spectroscopic (SERS) method was developed to rapidly and sensitively detect Ferbam on spinach leave surfaces directly without extracting them out. Ferbam was used to contaminate the surfaces of spinach leaves. Then branched gold nanoparticles (GNPs) colloids were deposited on the leave surface contaminated with pesticides and then air-dried for Raman measurements. The results show that Ferbam can be detected and discriminated in situ using the developed SERS method. The limits of detection (LODs) of ferbam was 0.3 mg kg⁻¹ on fresh spinach leaves. The developed SERS method provides a simple, rapid, and sensitive way to monitor pesticides on plant surfaces for safe productions of commercial vegetables.

**Introduction**

Ferbam (Fer) is an bactericide widely applied in vegetables and fruits. Ferbam residue in food have traditionally been monitored by high-performance liquid chromatography (HPLC) based methods. However, before chromatography analysis, time consuming sample extraction, clean-up and pre-concentration of samples are always required for this technique. Therefore, it is difficult to detect Fer with one simple extraction method and instrument. Surface-enhanced Raman spectroscopy (SERS) is a highly sensitive technique utilizing Raman spectroscopy and nanotechnology, which can greatly enhance the weak Raman molecular "fingerprint" through the placement of the analyte on noble metal nanoscale-roughened surfaces. SERS has been widely explored to detect pesticides. Many studies compared with these SERS methods demonstrated the faster and more convenient sample prep of SERS as compared to HPLC.

Up to now, only a few studies were reported on the in situ detection of Organophosphates, pyrethroids and neonicotinoids residues on several plant surfaces using homemade SERS substrates. Therefore, further studies are urgently needed to investigate the in situ SERS methods for more pesticides residues on various plant surfaces. Here, we aim to develop a simple, rapid, and sensitive in situ SERS method to detect bactericide (Fer) on spinach leave surfaces using GNPs with different morphology. To the best of our knowledge, this is the first investigation of detecting Ferbam with no pretreatment on fresh spinach leaves using a branched GNPs substrate for in situ SERS method.

**Experimental section**

**Chemicals and reagents.** Ferbam (Fer) was obtained from Crescent Chemical Co. (Hauppauge, NY, USA). Hydrochloroauric acid trihydrate (HAuCl₄·3H₂O), SDS, sodium citrate, and polyvinylpyrrolidone (PVP) were purchased from China Medicine (Group) Shanghai Chemical Reagent Corporation. Peptide (NH₂-Leu-Aib-Trp-Ome) was synthesized by conventional solution-phase method by using a racemization-free fragmentation/condensation strategy.

**Synthesis of Gold Nanoparticles and Characterization.** Spherical gold nanoparticles were prepared using Frens’ method. Ginger-like gold nanoparticles were fabricated as our previous
method[11]. Branched gold nanoparticles were obtained as method[12]. All the products prepared above were separated by centrifugation (TGL-16G, Anting Scientific Instrument Factory, China), and washed with deionized water several times. Optical spectra were recorded using a SPECORD200 spectrophotometer (Analytik Jena AG, Germany). TEM images were recorded on a JEM-2010 microscopy operated at 200 kV.

**SERS Detection of Ferbam in water**[8]. In a typical preparation, 2 μL of working standard solution of Fer 10 μg mL⁻¹ was dropped on a piece of parafilm (1x1 cm). Pure solvent (no Fer) was used as a negative control. After that, 2 μL of 250 μg mL⁻¹ Au NPs was added into it and mixed by pipetting for 20 s, and then 2 μL mixture was transferred onto the gold coated microscope slide, then allowed to dry under room temperature for Raman measurement.

**SERS Detection of Fer on spinach leaf surface.** Fresh spinach leaves (Purity Organic Fuji) were bought form a local supermarket in Nanchang, China. Fer-exposed samples were made suing a spiking method as follows: 20 μL of each concentration (0.1, 0.5, 1.0, 5.0 μg mL⁻¹, in water) of different Fer samples was transferred on spinach leaves, and then allowed to dry at room conditions. The Fer-exposed plant surface was about 1 cm². 4 μL of Au NPs were dropped on the Fer-exposed plant surface, mixed briefly with Fer on it by gentle pipetting for 20 s, and then allowed to dry under room temperature. Raman spectra were collected and analyzed under a DXR Raman microscope(Thermo Fisher Scientific, Madison, Wis., U.S.A.). Tests were done in triplicates. OMNIC™ software version 9.1 was used to control the Raman instrument.

### Result and discussion

GNPs had the diameter of about 40 nm (Fig.1 a). Ginger-like GNPs shown in Fig.1b had several tips (3-5), which might act as "hot spots" for SERS detection. Branched GNPs (Fig.1c) seemed to form a fishnet structure. These GNPs sol indicated different UV-vis spectra (Fig.1d), respectively.

The SERS fingerprint patterns of Fer on Au NPs was shown in Fig. 2. Sspecial GNPs and ginger-like GNPs exhibited weak peaks due to their weak interactions between the groups and GNPs. Branched GNPs indicated distinct characteristic peaks due to its strong interactions between the chemical groups and GNPs. The highest peak of Fer was observed at 1380 cm⁻¹ which associated with the deformation of CH₃ and stretching of CN. In addition, there were other relatively stronger peaks, such as 934 cm⁻¹ associated with the stretching of CH₃N and C=S, 1146 cm⁻¹ with the rocking of CH₃ and stretching of CN. Obviously, branched GNPs substrate exhibited the most effective enhancement function and its main SERS peaks showed the strongest Raman intensity. The crystal powder Raman spectrum of Fer has characteristic peak at 1385cm⁻¹. Benjamin et al. found that Fer SERS spectra had characteristic peak at 1377cm⁻¹ arising from the ν(C–N) stretching mode coupled to the symmetric δ(CH₃) motion due to gold nanorods as surface enhanced Raman spectroscopy substrates[13, 14]. So the peak of 1380 cm⁻¹ can be used as finger peak for trace analysis of Fer investigation.

Five different concentrations of Fer in deionized water were analyzed to determine the limits of detection (LOD) and linear ranges using the optimized SERS method. The result (Fig. 3a) indicated that the concentration of Fer was positively correlated to peaks intensity at 1380 cm⁻¹ in the SERS spectra. PLS was used to evaluate the quantitative capacity of the method and PLS plots were shoen in Fig.3b. Fer suggested satisfactory quantitative response with high correlation coefficients (0.9715) and low Root Mean Square Error of Calibration (RMSEC: 0.957). This result indicated that it could be effective to detect Fer on/in food matrix using the SERS method because Fer with stronger binding with brached GNPs could compete with other weaker interactions with branched GNPs.

Fresh spinach leaves surfaces were spiked with 0.3 mg kg⁻¹ Fer. Without GNPs and Fer, there was no Fer signal detected. Without extracting Fer, we developed an in situ method to investigate Fer on the spinach leaves surfaces by transporting GNPs onto the leave surfaces. After dropping GNPs on the leave surface, there were "coffee rings" formed on leaves due to unevenly distributing of GNPs, which can be clearly observed under microscope (Fig.4b). We selected the border of the rings and found they have consistent good signal. There were no significant background intererence observed on spinach...
leaves. The characteristic peaks of Fer at around 1381 cm\(^{-1}\) and 1146 cm\(^{-1}\) were shown in Fig. 4a. Compared with other in situ SERS methods\(^{[1]}\), our method applied the branched GNPs which was more easily available and sensitive. The developed method have great potential for practical monitoring of pesticides on plant surface.

Fig. 1 TEM images of (a) spherical GNPs; (b) ginger-like GNPs; (c) branched GNPs. (d) UV-vis spectra of different gold hydrosols.

Fig. 2 SERS spectra of Fer from different gold nanoparticles substrates.

Fig. 3 (a) second-derivative SERS spectra of characteristic peaks of different concentrations targets enhanced by branched GNPs. (b) PLS.
Fig. 4 (a) Raman spectra of Fer-exposed spinach leaf control, spiked level: 0.3 mg kg\(^{-1}\) (\(\approx\) 30 ng cm\(^{-1}\)); (b) Optical images of selected scan point. exposed-Fer spinach leaf was dropped with branched GNPs and mixed.

Conclusions

In summary, a simple and sensitive in situ SERS method was developed to detect ferbam on spinach leave surface. The proposed method can detect at least at 0.05 ppm in water and 0.03 \(\mu\)g cm\(^{-1}\) on spinach leave surfaces. The established SERS method shows great capacity to analysis trace Fer residue on spinach surface. Further experiments are needed to explore the application of this method to other pesticides.

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References