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Abstract. Surface-enhanced Raman spectroscopy (SERS) was employed to detect deoxyribose nucleic acid (DNA) variations associated with the development of nasopharyngeal carcinoma (NPC). Significant SERS spectral differences between the DNA extracted from early NPC, advanced NPC, and normal nasopharyngeal tissue specimens were observed at 678, 729, 788, 1337, 1421, 1506, and 1573 cm$^{-1}$, which reflects the genetic variations in NPC. Principal component analysis combined with discriminant function analysis for early NPC discrimination yielded a diagnostic accuracy of 86.8%, 92.3%, and 87.9% for early NPC, advanced NPC, and normal nasopharyngeal tissue DNA, respectively. In this exploratory study, we demonstrated the potential of SERS for early detection of NPC based on the DNA molecular study of biopsy tissues. © 2016 Society of Photo-Optical Instrumentation Engineers (SPIE) [DOI: 10.1117/1.JBO.21.12.125003]

Keywords: surface-enhanced Raman spectroscopy; nasopharyngeal carcinoma; deoxyribose nucleic acid; principal component analysis-discriminant function analysis.

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1 Introduction

Nasopharyngeal carcinoma (NPC) is one of the most commonly diagnosed head and neck cancers in southern China and Southeast Asia, with a reported incidence of 15–40 per 100,000 population in endemic areas.1,2 With advances in radiotherapy, patients with early stage (stages I and II) NPC tend to achieve satisfactory treatment outcomes with an associated 5-year survival of up to 94% compared with 50% to 60% in patients diagnosed at advanced stages (stages III and IV).3,4 Therefore, early diagnosis plays a vital role in the treatment of patients with NPC. The symptoms of NPC in early stages are not obvious;5 pathological examination of the biopsy specimen is the gold standard for diagnosis. However, early stage NPC often occurs in the pharyngeal recess where it is hard to detect. Moreover, the size of the tumor in the early stages is often small; some of these are submucosal, which makes it challenging to obtain a biopsy on pharyngorhinoscopic.6 An inadequate specimen may also obscure the diagnosis.

Deoxyribose nucleic acid (DNA) is subject to modifications and is a single molecule tool for bioanalytics. The Raman spectrum, referred to as the “molecular fingerprint,”7 can detect scattered photons with a spectrum of Raman peaks, each of which is characterized by a specific molecular bond. Collectively, these peaks provide intrinsic information of the sample and may yield a wealth of information about the chemical bonds associated with DNA and ribonucleic acid. Therefore, Raman spectroscopy technology can help in the early diagnosis of NPC by discriminating from the NPC tissues’ DNA. Furthermore, the weak Raman signals in conventional Raman spectroscopy can be dramatically enhanced to 10$^6$ to 10$^{15}$ times by surface-enhanced Raman spectroscopy (SERS) technology when the analytes of interest are in close proximity to gold or silver nanoparticles, which could allow for ultrasensitive discrimination of biological molecules between different samples.8 Kneipp et al.9 reported that SERS can be a single molecule tool for bioanalytics.

Our group previously published preliminary findings on DNA variations in the saliva, blood, and tissues from patients with NPC by SERS.10–12 However, the interference of other native constituents and exogenous substances in untreated
saliva, blood, and tissue samples compromised the diagnostic accuracy of NPC using SERS and overwhelmed most DNA signals. To capture more DNA signatures for genetic testing, use of purified DNA for SERS analysis may be an ideal alternative for improving the early diagnosis of NPC.

At present, the application of purified DNA in the diagnosis of NPC has not been reported in the literature. In this work, we sought to use DNA SERS for discrimination between early stage NPC, advanced stage NPC, and normal tissues at the molecular level.

2 Materials and Methods

2.1 Subjects and Protocol

A total of 91 tissue DNA samples (from 26 early stage NPC patients, 35 advanced stage NPC patients, and 30 healthy volunteers) were used in the study (Table 1). All patients were confirmed cases of NPC on histopathology. The disease staging was based on the seventh edition of the International Union Against Cancer/American Joint Committee on Cancer staging system for NPC.16,17 The study was approved by the Fujian Provincial Tumor Hospital Ethics Committee (Reference No. FICH-09911).

Tissue specimens obtained on pharyngorhinoscopy were kept in liquid nitrogen for half an hour. DNA extraction was performed according to the Takara manufacturer’s protocol as previously described by Paterson et al.18 A schematic illustration of the preparation of DNA samples is shown in Fig. 1(a).

Ultimately, DNA was purified in distilled water, and the concentration was detected by biomedical analyzer. Before SERS measurement, the DNA solutions were stored in a −20°C refrigerator to restrain degradation.

2.2 Preparation of Silver (Ag) Colloids and DNA Mixture

Ag colloids were prepared using hydroxylamine hydrochloride and Ag nitrate, in accordance with the method reported by Leopold and Lendl.19 In short, a total of 9 mL of 0.1 M sodium hydroxide solution was mixed with 10 mL of 0.06 M hydroxylamine hydrochloride solution to adjust the pH value of hydroxylamine hydrochloride solution. Then, the mixture was quickly added to 180 mL of 0.0011 M Ag nitrate solution, and the resulting mixture was shaken to obtain a homogeneous milky gray mixture. Absorption spectroscopy was utilized to characterize the prepared Ag colloids. The measurement demonstrated an absorption maximum at 418 nm. The particle sizes were determined by transmission electron microscopy, as described in our previous paper.20 The particle sizes followed a normal distribution with a mean diameter of 35 nm and a standard deviation of 5 nm. Next, the Ag colloids were concentrated by centrifugation at 10,000 rpm for 10 min to get high-concentration Ag NPs as the SERS substrate. Before being subjected to SERS measurement, the DNA solution (about 250 ng/μL) was mixed with silver colloid (about 2000 ng/μL) (at a 1:1 ratio). Finally, a fixed amount (2 μL) of DNA-silver colloid solution was dropped onto an aluminum (Al) plate for SERS measurement.

2.3 SERS Measurement

All SERS measurements were performed on a confocal microspectrometer equipped with a pellicle cooled charge coupled device camera for spectral detection and a 785 nm diode laser for Raman excitation. A Leica 20x objective was used to collect the SERS signals from the sample illuminated by 1 mW of incident laser power. The SERS spectra were measured in the wave number range of 500 to 1700 cm⁻¹. Each specimen was measured several times in different locations. The spectral acquisition and analysis were conducted using the package WIRE 2.0 (Renishaw plc).

<table>
<thead>
<tr>
<th>Specimen characteristics.</th>
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<tbody>
<tr>
<td>Normal group</td>
</tr>
<tr>
<td>Mean age, years</td>
</tr>
<tr>
<td>Mean DNA concentration</td>
</tr>
</tbody>
</table>

Fig. 1 (a) The schematic diagram of the preparation procedure of DNA-Ag NPs mixture. (b) Spectra taken from the DNA-Ag NPs mixture, the DNA sample without Ag NPs, and the Ag NPs substrate.
2.4 Data Preprocessing

In order to deduct background and to acquire the pure DNA SERS signal, the original spectral data were preprocessed using a five-order polynomial fitting algorithm. After fitting, area-under-the-curve normalization is applied to each spectrum to correct for variations in spectral absolute intensity.

2.5 PCA-DFA Analysis

Efficient diagnostic algorithms for differentiating SERS spectra between the early NPC, advanced NPC, and normal nasopharyngeal tissue DNA were applied using principal component analysis (PCA) combined with discriminant function analysis (DFA) by SPSS 15.0 software package (SPSS Inc., Chicago). PCA was used to reduce the dimensionality of spectral data to several principal components (PCs). A dependent-sample t-test was conducted to select the most diagnostically significant PCs ($P < 0.05$). DFA was performed based on these diagnostically significant PCs to maximize between-group variance and minimize within-group variance. In this way, one sample (i.e., one spectrum) was excluded from the dataset, and the algorithm based on PCA-DFA was redeveloped using the remaining DNA spectra. To further evaluate the performance of the PCA-DFA algorithm, receiver operating characteristic (ROC) curves were generated.

3 Results and Discussion

Figure 1(b) shows the DNA SERS spectra after silver colloid enhancement. The DNA SERS peaks were abundant while the normal Raman spectral peaks were few due to the interference of large autofluorescence background. Meanwhile, the interference from silver colloid substrate was scarcely seen. The observations indicated that there were strong interactions between the silver nanoparticles and the DNA molecule. Because of this interaction, a DNA sample can closely adsorb on the silver colloid particles’ surfaces, leading to quenching of biological autofluorescence and simultaneously increasing the intensity of the DNA Raman scattering. This is in agreement with the principle by Dulkeith et al.

Raman bands, also referred to as “molecular fingerprints,” can identify a particular chemical bond or a single functional group in the molecule. Therefore, the more abundant spectral peak signals emerged, the more the material component was reflected. In this study, prominent SERS peaks were observed in all 91 samples at 678, 729, 788, 1093, 1337, 1421, 1460, 1506, and 1573 cm$^{-1}$ [Fig. 2(a)]. All SERS peaks were assigned to DNA according to the literature listed in Table 2.

Figures 2(b) and 2(c) show that the major SERS spectral differences between the DNA of normal, early, and advanced cancer tissue had different intensities of the bands. For example, the modes at 678, 729, 788, 1093, 1337, 1421, 1460, 1506, and 1573 cm$^{-1}$ were due to DNA bases (adenine, thymine, cytosine, and guanine); 1093 cm$^{-1}$ was attributed to the symmetric PO$_2^-$ stretching vibration of the DNA; and 1460 cm$^{-1}$ was assigned to CH$_2$/CH$_3$ deformation of deoxyribose. It is common knowledge that carcinogenesis is prominently associated with DNA damage, especially via nucleobase lesions formed by reactive oxygen species. Meanwhile, a slight modification in the DNA can affect the intensity of Raman vibrations in the DNA.
molecules. Malins et al. and Kourkoumelis et al. had reported that the intensity of spectral peak in 1093 and 1460 cm$^{-1}$, respectively, were different between the normal cells and the cancer cells. These changes probably reflect the variations in relative DNA content and conformations that are involved in the occurrence and development of NPC.

Genomic instability is the hallmark of human cancer, being responsible for abnormal cancer cell proliferation, invasion, and metastasis. Many genetic mutations have been linked to pathogenesis of NPC, such as those linked to Epstein–Barr virus (EBV) infections. There exists some potential mechanisms that may explain the spectral differences among normal tissues, early stage, and advanced stage NPC tissues. For early stage NPC, lower intensities of most SERS peaks may be due to high frequency of loss of heterozygosity in the allelic status. With the progression of NPC, there was still a lack of heterozygosity on chromosome in advanced stage NPC. For advanced stage NPC, it was also found to be infected by EBV because of its high-grade dysplastic lesions, while EBV was hardly found in low-grade dysplastic lesions and the histologically normal nasopharyngeal epithelia. These variants may cause the differences among the three types of tissue DNAs. However, the other mechanisms for these spectra intensity differences of SERS peaks between normal, early stage, and advanced stage NPC warrant further investigation.

To quantitatively analyze the ability of DNA SERS technique for NPC detection, PCA-DFA was performed on the obtained SERS spectra. Forty-nine PCs (99% of the total explained variance) were extracted by PCA. Then, the most significantly significant PCs (i.e., PC2, PC7, PC9, PC15, PC17, and PC18) were determined by the dependent-sample t test. Figure 2(d) shows the plot of these six PC loadings. The plot enables one to know which spectral variables are dominating or influencing the PCA-DFA model. As the most diagnostically significant PCs, the prominent Raman peaks that appeared in these PC loadings, such as 678, 729, 788, 1093, 1337, 1421, and 1573 cm$^{-1}$, are considered as the diagnostically relevant spectral bands. These bands are also easily identifiable in the difference spectrum [Fig. 2(b)]. Next, DFA was used to discriminate among the three groups based on these six PCs and a priori knowledge of which spectra were replicates, a process that does not bias the analysis.

Figure 3(a) shows the scatter plot of the coefficient for each sample according to the two discriminant functions after PCA for normal, early stage, and advanced stage NPC samples. The three scatter plots were distributed in three relatively separate areas in spite of the few overlaps, which indicates that the SERS spectra of the three different types of DNA samples could be well discriminated. Ultimately, use of the PCA-DFA algorithm for differentiation of DNA of normal, early stage, and advanced stage NPC achieved higher diagnostic sensitivities of 83.3%, 80.8%, and 85.7%; specificities of 83.6%, 84.6%, and 82.1%; and accuracies of 87.9%, 86.8%, and 92.3%, respectively (Table 3).

We have done direct Raman spectral detection of tissue, which provided only a few characteristic peaks related to DNA, because most DNA signals were overwhelmed by other native constituents and exogenous substances in untreated tissue samples. DNA extraction and detection by SERS can capture more DNA fingerprints for improving the early diagnosis of NPC. In our study, the extracted DNA SERS spectra obtained by SERS techniques achieved...
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Table 3: Classification results of SERS spectra prediction of three DNA types using PCA-DFA.

<table>
<thead>
<tr>
<th>DNA type</th>
<th>Normal group</th>
<th>Early NPC</th>
<th>Advanced NPC</th>
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<tbody>
<tr>
<td>Normal</td>
<td>25</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>Early NPC</td>
<td>3</td>
<td>21</td>
<td>2</td>
</tr>
<tr>
<td>Advanced NPC</td>
<td>3</td>
<td>2</td>
<td>30</td>
</tr>
<tr>
<td>Sensitivity</td>
<td>83.3(25/30)</td>
<td>80.8(21/26)</td>
<td>85.7(30/35)</td>
</tr>
<tr>
<td>Specificity</td>
<td>83.6(55/61)</td>
<td>84.6(58/65)</td>
<td>82.1(54/56)</td>
</tr>
<tr>
<td>Accuracy</td>
<td>87.9(80/91)</td>
<td>86.8(79/91)</td>
<td>92.3(84/91)</td>
</tr>
</tbody>
</table>

86.8% of accuracy, superior to the results of 75% accuracy provided by direct tissue Raman spectra for early NPC detection.

To further evaluate the performance of the PCA-DFA algorithm, ROC curves were generated [Fig. 3(b)]. The integration of the ROC curves was 0.981, 0.940, and 0.978 for stage NPC versus normal tissues, for early stage NPC versus normal tissues, and for both early and advanced stage NPC versus normal tissues, respectively. These high ROC areas demonstrate the good performance of the PCA-DFA method based on DNA SERS spectra for promoting early diagnosis of NPC. Furthermore, the complete process of SERS detection for DNA of NPC tissue, including the additional step of DNA extraction, takes only 1 day, which is shorter than that of the direct pathological detection for NPC tumor tissue (usually needs 3 to 5 days). Consequently, it is hoped that early detection of NPC may be achieved by specific SERS DNA, such as a single EBV DNA molecule, to provide a specific, rapid, and accurate diagnosis of NPC at an early stage.

In conclusion, the different intensities of significant SERS peaks could be recognized between the early stage, advanced stage, and normal tissue samples, which could reflect gene variations in NPC. PCA-DFA multivariate analysis was performed to classify three types of DNA SERS data, with a diagnostic accuracy of 87.9%, 86.8%, and 92.3% for the three tissue groups, respectively. This exploratory study first underlines the potential of the SERS technique for early molecular diagnosis of NPC.

Meanwhile, we also recognize that it is still a challenge to apply this DNA-based SERS to clinical applications due to some limitations. First, our study has not identified the specific spectra of the DNA of EBV and cannot provide the detailed construction information of DNA. Second, the tissue sample size is limited, too small to comprehensively evaluate the reproducibility and accuracy of this detection strategy. Therefore, further research needs to be conducted to sensitively detect the specific base sequence of EBV-DNA and to systematically verify the utility of the proposed method in real clinical application.

Disclosures

There is no competing interest in this work.

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