10-22-2009

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PCR-Free Quantification of Multiple Splice Variants in a Cancer Gene by Surface-Enhanced Raman Spectroscopy

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Received: May 16, 2009

We demonstrate a surface-enhanced Raman spectroscopy (SERS) based array platform to monitor gene expression in cancer cells in a multiplex and quantitative format without amplification steps. A strategy comprising DNA/RNA hybridization, S1 nuclease digestion, and alkaline hydrolysis was adopted to obtain DNA targets specific to two splice junction variants, \( \Delta (9,10) \) and \( \Delta (5) \), of the breast cancer susceptibility gene 1 from MCF-7 and MDA-MB-231 breast cancer cell lines. These two targets were identified simultaneously, and their absolute quantities were estimated by a SERS strategy utilizing the inherent plasmon–phonon Raman mode of gold nanoparticle probes as a self-referencing standard to correct for the variability in surface enhancement. The results were then validated by reverse-transcription polymerase chain reaction. Our proposed methodology could be expanded to a higher level of multiplexing for quantitative gene expression analysis of any gene without any amplification steps.

Introduction

Studying gene expression at the transcript level is a vital and commonly adopted approach in biological and biomedical sciences. Effective and efficient screening of gene expression requires analytical techniques with the ability to detect multiple transcripts and provide quantitative estimates of especially the low abundantly expressed transcripts, which are most commonly analyzed qualitatively via polymerase chain reaction (PCR) amplification. Conventional PCR for gene expression analysis is semiquantitative in nature. In addition, if more than one pair of primers is used in PCR to amplify the corresponding segments of target DNA simultaneously, extensive optimization is required. Furthermore, different amplicons are often amplified at different efficiencies to result in competition among multiple primer pairs in the reaction. Although real-time PCR is known for its ability to quantify and multiplex, it is not always effective in situations where a high amplification efficiency is not available, and challenges do exist in a few technical aspects, such as the template quality and data interpretation, especially when dealing with a very low copy number of targets. In addition, fluorescence-based detection poses a limit on the degree of multiplexing because of the broad and overlapping emission profiles when higher levels of multiplexing are required. The application of other common methods for gene expression studies, such as Northern blots and quantitative nuclease protection assays, either is semiquantitative or lacks the needed sensitivity. Microarray is undoubtedly an established method for genome-wide analysis. However, such an extensive assay may not always be necessary for many specific applications and is more expensive compared to other techniques. In addition, this approach requires a universal genomic data quality control standard for improved reliability.

To meet the requirements of detection sensitivity and multiplexing capability for gene expression studies, surface-enhanced Raman spectroscopy (SERS) could be an extremely promising alternative because of its intrinsically sharp fingerprints, single-molecule sensitivity, and availability of a wide range of photostable labels. Many SERS-based approaches exist for nucleic acid detection. A typical probe design includes the adsorption of fluorophore-labeled DNA onto colloidal silver nanoparticles, the covalent attachment of fluorophore-labeled DNA onto gold nanoparticles, the use of plasmonic nanoprobes with metal nanoparticles functionalized with a stem-loop DNA molecule tagged with a Raman label, a surface-enhanced resonance Raman scattering beacon approach, and the sequential attachment of DNA strands and nonfluorescent Raman tags (RTags) to gold nanoparticles. In addition, different substrates and detection platforms have been developed, and multiplex detection has been demonstrated.

Although successful in qualitative studies, previous SERS applications were limited in providing quantitative information because of the variability of surface enhancement. Ongoing efforts in fabricating tunable and reproducible substrates are aimed at controllable SER activity; however, uniform SERS for reliable quantification has not been fully achieved. Although quantitative behavior has been shown with a dilution series of dye-labeled DNA sequences, no quantitative detection of unknown DNA targets was demonstrated using these systems. An alternative SERS-based quantification strategy using isotope-edited dyes as internal standards was proposed; however, this approach entails the synthesis of new dyes and requires optimization. Hence, a consistent SERS-based quantification method to evaluate unknown nucleic acid targets without amplification will be highly desirable.

In our study, we demonstrate a SERS approach to identify and quantify multiple gene segments extracted from cells based on the concept of using the plasmon–phonon bands as a self-referencing standard. We used the breast cancer susceptibility gene 1 (BRCA1) as a model gene and its two alternative splice variants, \( \Delta (9,10) \) (splice junction skipping exons 9 and 10) and \( \Delta (5) \) (splice junction skipping exon 5), as targets. Alternative splicing is a crucial mechanism for generating protein diversity, and BRCA1 is a good model because it exhibits cell- and tissue-
specific alternative splicing patterns that could serve as a potential cancer marker. We compare the differential expression of these two BRCA1 splice variants (Δ(9,10) and Δ(5)) in two breast cancer cell lines, MCF-7 and MDA-MB-231, simultaneously using nonfluorescent Raman probes by isolating RNA from cells. DNA/RNA hybridization, S1 nuclease digestion, alkaline hydrolysis, and an array-format SERS sandwich assay. Quantification results were validated by reverse-transcription polymerase chain reaction (RT-PCR) and compared with the literature, where available. Because the demonstrated SERS method can monitor multiple fragments at a high level of sensitivity without any amplification steps, lowly expressed splice variants could be detected and quantified on an absolute scale. Thus, this method can serve as a complementary detection tool to the existing methods and yet fill the niche in detecting very low quantities accurately and efficiently.

Experimental Methods

Preparation of DNA Targets from Cancer Cells. MCF-7 and MDA-MB-231 cells were grown in Eagle’s Minimum Essential Medium (ATCC) supplemented with 0.01 mg/mL bovine insulin and 10% FetalPlex animal serum complex (Gemini Bio-Products) and an ATCC-formulated RPMI-1640 medium supplemented with 10% FetalPlex animal serum, respectively. Total RNA was first isolated from approximately 10⁷ cells using the QIAGEN RNEasy Mini Kit. A total of 30 µL of hybridization cocktail (20 µL of total RNA, 9 µL of a 3X aqueous hybridization solution, and 1 µL of DNA oligonucleotides with sequences complementary to the RNA segments of interest) was then added and heated to 75 °C for 10 min, and the mixture was incubated overnight at 55 °C. Next, 270 µL of S1 nuclease mix was added, and the mixture was incubated at 37 °C for 45 min. A hydrolysis solution (1.6 M NaOH and 135 mM EDTA) was then added, and the mixture was incubated at 95 °C for 15 min. Finally, a neutralizing solution (1 M HEPES, 1.6 M HCl, and 6X SSC) was added to suspend the DNA targets of interest in a neutral medium.

SERS Sandwich Assay. Sequences of BRCA1 splice junctions Δ(9,10) and Δ(5) were obtained from an NH-maintained GenBank (accession number U14680) as listed in Table 1. A SERS sandwich assay was performed on a gold-coated glass slide covered with a silicone mask containing an array of wells (3 mm in diameter). In a typical assay, 10 µL of 1 µM capturing strand (CS) with the sequence spanning the downstream half of the exon/exon junction with 3'-thiol modification was spotted onto the slide and incubated in a humidity chamber at room temperature to reduce nonspecific binding. Next, 10 µL of the target strand (TS) with the sequence complementary to the exon/exon junction was added to hybridize with the immobilized CS by incubating in a humidity chamber at 37 °C for 4 h. Finally, 10 µL of a DNA-AuP-RTag probe (synthesis procedure detailed in work by Sun et al.) with probing strands (PSs) containing the same sequence as that of the upstream half of the exon/exon junction with 5'-thiol modification and nonfluorescent RTags covalently attached to 40 nm gold nanoparticles was hybridized to the overhanging part of the TS by incubating in a humidity chamber at 37 °C for 4 h. In this study, 4-mercaptopyridine (RTag-1) and 2-thiazoline-2-thiol (RTag-2) were used as nonfluorescent Raman labels. For multiplex detection, sandwich structures formed by different sets of sequences were immobilized on the same detection spots. All tests can be performed on the same slide.

A SENTERRA confocal Raman system (Bruker Optics) was used in this study. A 785 nm diode laser with 10 mW power at the laser source and a 50× air objective (N.A. 0.7, infinity- and flat-field-corrected) were used to acquire all of the Raman spectra. A 20 s integration time was chosen. OPUS software was used to analyze all of the spectra.

RT-PCR. After RNA was treated with DNase I (New England BioLabs), the iScript cDNA Synthesis Kit (Bio-Rad Laboratories) was used to synthesize cDNA. Then PCR was performed in a DNA Engine (PTC-220) Peltier Thermal Cycler (MJ Research) using Taq DNA polymerase (Fisher Scientific) and the corresponding primers. Finally, PCR products were analyzed in 3% agarose gel prestained with ethidium bromide.

Results and Discussion

To elucidate expression patterns of specific genes in different cell lines, three major tasks were defined: (1) isolation of the nucleotide targets of interest from the isolated RNA pool; (2) qualitative assessment of these targets by SERS; (3) quantification by SERS and validation by conventional methods. In this study, BRCA1 was used as the model gene, and absolute expression levels of the two splice junction variants, Δ(9,10) and Δ(5), in the two breast cancer cells lines, MCF-7 and MDA-MB-231, were determined using our SERS platform.

Detailed detection procedures are described in Figure 1. First, a mixture of single-stranded DNA oligonucleotides (40 bp) complementary to exon/exon junctions Δ(9,10) and Δ(5) was added to the isolated total RNA from the corresponding cancer cell line and allowed to hybridize. Then S1 nuclease was added to digest the single-stranded nucleic acids, including unhybridized parts of the specific mRNA, all of the nonspecific RNA,
controls were first performed using fabricated DNA targets to demonstrate the experimental procedure is effective, two positive probes; detection of splice junctions in MDA-MB-231 using 

of splice junctions in MCF-7 using 

detection of the splice variant expression, four cases were used for the duplex detection spot. To demonstrate duplex complementary to the downstream half of the interested targets and a mixture of DNA-AuP-RTag probes bearing sequences complementary to the upstream half of the interested targets targets simultaneously, a mixture of CSs comprising sequences 

remove heteroduplexes containing mismatched regions. Al- single-stranded RNA or DNA into 5'-mononucleotides, leaving only the specific DNA–RNA duplexes intact because S1 nuclease can hydrolyze single-stranded RNA or DNA into 5’-mononucleotides and remove heteroduplexes containing mismatched regions. Alkaline hydrolysis was subsequently applied to inactivate S1 nuclease and hydrolyze the RNA components of the DNA–RNA duplexes. The resulting DNA TSs contained sequences specific to Δ(9,10) and Δ(5) in proportion to their mRNA expression. Quantification of these DNA targets provides an estimate of the expression of splice junctions Δ(9,10) and Δ(5) in the corresponding cell line.

Figure 1 shows the sandwich structure formed by linking CS and PS with the corresponding TS via hybridization. When the sandwich structure is formed, Raman signals from the corresponding nonfluorescent RTag would be detected. It should be noted that T10 was used as a linker for both CS and PS to minimize nonspecific adsorption of DNA molecules onto the gold surface because thymine has the lowest affinity to gold compared to other nucleotides. A layer of 6-mercaptop-1-hexanol molecules was applied after CS immobilization to prevent nonspecific adsorption of DNA from solution and displace nonspecifically adsorbed CS molecules to increase the detection specificity. To monitor the Δ(9,10) and Δ(5) DNA targets simultaneously, a mixture of CSs comprising sequences complementary to the upstream half of the interested targets and a mixture of DNA-AuP-RTag probes bearing sequences complementary to the downstream half of the interested targets were used for the duplex detection spot. To demonstrate duplex detection of the splice variant expression, four cases were investigated: detection of splice junctions in MCF-7 using Δ(9,10)-AuP-RTag-1 and Δ(5)-AuP-RTag-2 probes; detection of splice junctions in MCF-7 using Δ(9,10)-AuP-RTag-2 and Δ(5)-AuP-RTag-1 probes; detection of splice junctions in MDA-MB-231 using Δ(9,10)-AuP-RTag-1 and Δ(5)-AuP-RTag-2 probes; detection of splice junctions in MDA-MB-231 using Δ(9,10)-AuP-RTag-2 and Δ(5)-AuP-RTag-1 probes. To ensure that the experimental procedure is effective, two positive controls were first performed using fabricated DNA targets Δ(9,10) and Δ(5). In positive control 1, Δ(9,10)-AuP-RTag-1 and Δ(5)-AuP-RTag-2 probes were used, while in positive control 2, Δ(9,10)-AuP-RTag-2 and Δ(5)-AuP-RTag-1 probes were used. As shown in Figure 2, the controls show distinct characteristic peaks attributable to the corresponding RTags (1001, 1091, and 1200 cm\(^{-1}\) for DNA-AuP-RTag-1; 1295 and 1349 cm\(^{-1}\) for DNA-AuP-RTag-2). These characteristic peaks were also observed for all of the detection cases, indicating that the two splice variants are expressed in the chosen breast cancer cell lines. To ensure detection reliability, the same targets were monitored by different RTags for cross-validation. Detection specificity was tested for a control sample using a target solution containing all of the components except the DNA targets. As expected, a very weak nonspecific signal was obtained (Figure 2). It should be noted that a higher level of multiplexing (8-plex) has been achieved using DNA-AuP-RTag probes. Therefore, it is possible to monitor more gene segments from the same cell line or different cell lines simultaneously using the proposed strategy to improve the detection efficiency.

For quantification, we use the normalizing standard developed in our previous studies\(^\text{29,30}\) based on the plasmon–phonon band to quantify two different DNA targets identified in the duplex detection scheme discussed in Figure 1. Individual sets of CSs and DNA-AuP-RTag probes were used to quantify the corresponding DNA targets in the target mixture obtained from different cell lines through DNA/RNA hybridization, S1 nu-
The normalized control signal or nonspecific signal was then equation (Intensitynorm signal, and the resulting intensity was used in the calibration expressed in MCF-7 and MDA-MB-231 cells.

$Δ$
clease digestion, and alkaline hydrolysis. Here we quantify $Δ$(9,10) from the characteristic band of the RTag used) from the characteristic band of the tag (1091 cm$^{-1}$ for RTag-1) and the respective plasmon–phonon band (around 220 cm$^{-1}$) of each spectrum. The corrected characteristic peak ($I_0$) was then normalized against the corrected plasmon–phonon band ($l_0$). The normalized control signal or nonspecific signal was then subtracted from the plasmon–phonon band-normalized sample signal, and the resulting intensity was used in the calibration equation ($\text{Intensity}_{\text{norm}} = 6.842\times10^{0.2169} + 0.1138$) developed in our previous study to evaluate the target concentration. A similar procedure was used to quantify $Δ$(5) expressed in both MCF-7 and MDA-MB-231 cells. The estimated DNA targets for splice junctions $Δ$(9,10) and $Δ$(5) from both MCF-7 and MDA-MB-231 cells are summarized in Table 2. To validate these results, a conventional RT-PCR method was used to compare the expression levels of the splice junctions $Δ$(9,10) and $Δ$(5) from both cell lines qualitatively. Primers for these splice junctions were designed to cross specific exon/exon junctions, and a housekeeping gene, $β$-actin, was used as an internal control for PCR analysis. The quantification results presented in Table 2 reveal that the expression of $Δ$(9,10) was higher in MCF-7 than in MDA-MB-231, which agrees well with the results from the study of a comparison of expression profiles of predominant BRCA1 splice variants in tumor and normal cell lines, while the expression of $Δ$(5) was lower in MCF-7 than in MDA-MB-231. The expression of $Δ$(9,10) was found to be higher than that of $Δ$(5) in MCF-7 and lower than that of $Δ$(5) in MDA-MB-231. The RT-PCR validation shown in Figure 4 confirms the SERS results. It should be noted that customization of the slide could be done to integrate array spots for multiplex detection and quantification with controls as well as any other necessary tests to perform all of the corresponding sandwich assays simultaneously by choosing appropriate sets of CSs and PSs.

**Conclusions**

This study demonstrated the use of SERS as an efficient and quantitative analytical tool for breast cancer cell gene expression analysis. The proposed methodology capitalizes on the multiplex sensing capability coupled with highly sensitive and unambiguous detection by SERS using inexpensive nonfluorescent RTags to quantify gene fragments without amplification. The strategy also bypasses the need for a highly uniform SERS substrate to achieve reliable quantification. Duplex detection and quantification of the BRCA1 expression at the resolution of alternative splicing were successfully achieved. This method could be extended to study any segment of the genome in a multiplex and quantitative manner.

**Acknowledgment.** Funding from NIH-NCI SRO3CA121347-02, a CFSE grant, a Bilsland Fellowship, and the Oncological Science Center from Purdue University are acknowledged. The work was conducted at the Physiological Sensing Facility of the Bindley Bioscience Center at Purdue University.

**References and Notes**

(20) Bell, S. E. J.; Srimuthu, N. M. S. Chem. Soc. Rev. 2008, 37, 1012.


JP908225F