Deciphering the Finger Prints of Brain Cancer Glioblastoma Multiforme from Four Different Patients by Using Near Infrared Raman Spectroscopy

Hirendra Nath Banerjee1,*, Arnold Banerji1, Arunendra Nath Banerjee1, Eilena Riddick1, Jenae Petis1, Shivonda Evans1, Megha Patel1, Carl Parson1, Valerie Smith1, G. Gwebo1 and Sarah Voisin1

1Department of Natural Sciences and Pharmaceutical Sciences, ElizabethCity State University, University of North Carolina, ElizabethCity, NC-27909, USA
2University De Lyon, France

Abstract
To explore the effectiveness of Raman spectra to diagnose brain cancer glioblastoma multiforme (GBM), we investigated the Raman spectra of single cell from four different GBM cell lines developed from four different patients and analyzed the spectra. The Raman spectra of brain cancer (GBM) cells were similar in all these cell lines. The results indicate that Raman spectra can offer the experimental basis for the cancer diagnosis and treatment.

Keywords: Raman spectroscopy; Glioblastoma multiforme; Cancer; Optical tweezers

Introduction
Cancer is one of the most serious diseases threatening human health and life, and the influence of this disease is steadily increasing. Due to the fact that pathogeny of cancer and some correlative illnesses have not been found, and the effective diagnosis and complete therapy cannot be carried out at present, it is impossible to control the progress of the state of the illness for cancer patients in terminal stages. Therefore, early diagnosis and timely therapy is the most effective approach in improving the surviving chance of the patient. It is very difficult to diagnose cancer in modality because the early symptom of cancer is not evident and it has no distinct difference from that of some other illness. Therefore, it is a research emphasis to find new, effective diagnosis technology and treatment method [1]. At present a diagnosis of cancer is made on histological evaluation with possible prior cytological evidence, but early diagnosis of cancer affords early intervention and greatest chance of cure. Because the ability to detect early biochemical changes associated with carcinogenesis prior to the changes a pathologist identifies would revolutionize cancer diagnostics, in recent years there has been much interest in the use of optical diagnostics in cancer detection [2]. Raman spectroscopy is a non-destructive optical technique based on the inelastic scattering of monochromatic light. When a sample is irradiated, an exchange of energy takes place between the excitation light and the molecules of the sample, which results in a measurable shift in the wavelength of the incident laser light [3]. And because the energy-levels involved are unique for every molecule, the resulting Raman spectrum is essentially a ‘biochemical fingerprint’, containing bands representing molecular normal modes of vibration of all molecules within the interrogated region of the sample. This technique has several advantages on the other major vibrational spectroscopy technique (IR spectroscopy), which are the absence of interference due to water, sharp bands and a spatial resolution down to ~1 µm [4]. The oncological applications of Raman spectroscopy are already diverse and have shown to be successful in classification, discrimination, and bimolecular investigation of cancer cells [4–7]. Both Raman spectroscopy and surface enhanced Raman spectroscopy (SERS) are proving to be invaluable tools in the field of biomedical research and clinical diagnostics. The robust, compact, fit-for-purpose Raman spectrometer designs are appropriate for use in surgical procedures to help surgeons assess tumors and allow rapid decisions to be made. Raman systems are also being developed for molecular diagnostic testing to detect and measure human cancer biomarkers. Based on the SERS technique, this approach potentially could change the way bioassays are performed to improve both the sensitivity and reliability of testing. Some of the many applications of Raman and SERS in cancer biology include:

• Examination of biopsy samples
• In vitro diagnostics
• Cytology investigations at the cellular level
• Bioassay measurements
• Histopathology using microscopy
• Direct investigation of cancerous tissues
• Surgical targets and treatment monitoring
• Deep tissue studies
• Drug efficacy studies

Astrocytes are cells found in the brain. They are closely related to the neurons of the brain and form the blood brain barrier. Astrocytoma is Astrocyte cells that have become cancerous. These cells can form two types of tumors: The first type is benign or slow growing, and the second type is malignant or fast growing. A specific molecular diagnostic marker still does not exist for this cancer which has a very high mortality rate. We earlier reported the difference in Raman spectroscopy in normal human Astrocytes and Astrocytoma [1]. In this study we report the results of a comparison study of Raman spectroscopic analysis of Astrocytoma cell lines developed from four different male patients of different age but of same stage cancer.
Materials and Methods

Cell culture

GBM cell lines were obtained from three white male patients of different age groups HTB12, HTB14, HTB159 (obtained from ATCC, USA) and one African American male patient GBM A172 (obtained from Duke University Tisch Brain Tumor Center). HTB12, HTB14, HTB159 were cultured in L 15 medium supplemented with 10% calf serum and antibiotics in a carbon dioxide incubator at 37°C. GBM A 172 was cultured in DMEM medium under similar conditions mentioned above. Cell viability and count was determined by the Trypan Blue method using a standard hemocytometer.

Analysis of individual cells by the use of optical tweezers, and near-infrared Raman spectroscopy

The experimental setup of the optical tweezers and Raman spectroscopy was discussed in detail previously [8]. Briefly, a laser beam at 785 nm from a wavelength-stabilize, beam shaped-circularized semiconductor diode laser is introduced into an inverted microscope (Nikon 200005) through a high numerical aperture objective (100x, N. A. =1. 30) to form an optical trap. The wavelength of the diode laser was temperature stabilized to avoid drifts. A cell in a liquid medium was selectively trapped with the radiation force yielded by the focused laser beam. The same laser excited Raman scattering from the trapped cell. The back scattered light was collimated with the same objective lens and passed through a 100 um confocal pinhole aperture to reject most of the off focusing Rayleigh scattering light. Two interfering notch filters were used to remove most of the onfocusing Rayleigh scattering light. The Raman scattering light was then focused onto the entrance slit of an imaging spectrograph equipped with liquid nitrogen cooled charged coupled detector (CCD). The image of the trapped cell was observed with an illumination lamp and a video camera system, the spectral resolution of the system was estimated to be ~6 cm^{-1}. In order to sample and average most of the trapped cell, the laser beam was steered rapidly (~200 Hz) by a pair of computer controlled Galvo mirrors across the major area of the cell. Because the steering speed was very fast, the cell was found not to follow the steering beam so that the major portion of the cell was excited within the acquisition time. The acquisition time was typically 120s for an individual cell with an excitation power of 15 mW at 785 nm. Each spectrum was averaged from more than 15 separate cells and each experiment repeated three times.

Software and data processing

All our data analysis was performed with R, using different packages (hyperSpec, MASS, permax). The raw spectra were first cut in the 500-1800 cm^{-1} range of the Raman shift. Then, the cut spectra were baseline corrected thanks to automatically fit polynomials: this step allows getting spectra with band edges of up to the theoretical baseline. The back scattered light was collimated with the same objective lens and passed through a 100 um confocal pinhole aperture to reject most of the off focusing Rayleigh scattering light. Two interfering notch filters were used to remove most of the onfocusing Rayleigh scattering light. The Raman scattering light was then focused onto the entrance slit of an imaging spectrograph equipped with liquid nitrogen cooled charged coupled detector (CCD). The image of the trapped cell was observed with an illumination lamp and a video camera system, the spectral resolution of the system was estimated to be ~6 cm^{-1}. In order to sample and average most of the trapped cell, the laser beam was steered rapidly (~200 Hz) by a pair of computer controlled Galvo mirrors across the major area of the cell. Because the steering speed was very fast, the cell was found not to follow the steering beam so that the major portion of the cell was excited within the acquisition time. The acquisition time was typically 120s for an individual cell with an excitation power of 15 mW at 785 nm. Each spectrum was averaged from more than 15 separate cells and each experiment repeated three times.

The use of PC scores instead of the raw data to remove the outliers was repeated from the beginning until outliers were no longer detected. The use of PC scores instead of the raw data to remove the outliers eliminates collinearity problems and avoids inverting a singular matrix [10]. After the removal of outliers, the remaining spectra recorded in every given cell were averaged and all the spectra belonging to a given cell line were grouped together.

Details on the parameters set for the statistical analysis

The PMAV was applied directly to the spectral data and the Euclidean method was used to calculate pairwise distances for every permutation. After centering the data, a PCA was performed with either the 3 cell lines taken into account, using both sets of experiments (n = 8 + 9 = 17 cells/cell line). After centering the data, several PCA were performed on couples of cell lines, using also both sets of experiments. Then, LDA, followed by the LCP, were conducted with several PCs taken into account. The automated TSPWT function in R uses by default the entire permutation distribution, which is only feasible if the sample sizes are fairly small. For each of our TSPWT, the sample sizes were n = 17 cells/cell line, which is too high to use the entire distribution. Therefore, we fixed the number of permutations to N = 15000.

Results

Similarities in band intensities were observed from Astrocytoma cancer cells obtained from four different patients (Figure 1). Table 1 shows the Raman bands and their tentative molecular assignation based on earlier studies on mammalian cells [4,7,11]. Comparison of the band intensities in these cells showed an increase in emission at 878, 1004, 1264, 1302, 1442 and 1660 cm.

Our results, thus, shows distinct similarities in biochemical composition of Astrocytoma grade 4 obtained from four different patients as evidenced by band intensity in Raman spectra.

Discussion

Optical trapping is a powerful tool that uses optical forces to hold a micrometer-sized particle near the focus of a single tightly focused laser beam in three dimensions [2]. This technique has been used to capture and manipulate biological particles such as cells, bacteria, virus, and chromosomes without obvious damage using near-infrared laser beams [3,4]. Biomechanical properties of single cells and single biomolecules have been extensively studied with optical angular momentum of the laser beam to form micromachines [7,11]. The combination of Raman spectroscopy with optical trapping offers a new degree of direct analysis of chemical spectroscopic studies of single optically trapped living cells or single organelles for the purpose of the analysis and identification, without the need of mechanical immobilization or introducing biochemical stains or tags. With the introduction of vibrational spectroscopy [12-16], the constitution and conformation of macromolecules inside the trapped cells can thus be identified and studied. The essence of this paper is to demonstrate that this technique is an effective method to study in real-time changes in biochemical properties of a single cell (or organelle) which can therefore be used to detect cancer. It is known that biological cells contain a very complex mixture of organic, inorganic and biochemical components enclosed in cell membrane. A large number of biological molecules such as proteins, nucleic acids, polycyclicarides, and lipids, etc. and a variety of electrolytes populate the interior of the cell. The
observations and moved away from the coverslip up into solution. A mobile particle to be held at the focus of a laser beam for long-time confocal Raman spectroscopy. In our scheme, optical trapping allows artifacts, we used the LTRS system that combines optical trapping and electrochemical potentials across the cell membrane which is known to affect cellular functions. In order to avoid this problem and consequent drawbacks of the conventional method is that the living cells under study must be immobilize either on a glass cover slip or in a solid culture medium in order to avoid Brownian motion or cell motility when suspended in a liquid medium. Cell immobilization using a cover slip changes the chemical and physical micro environment of this living microorganism and may yield among other changes in the electrochemical potentials across the cell membrane which is known to affect cellular functions. In order to avoid this problem and consequent artifacts, we used the LTRS system that combines optical trapping and confocal Raman spectroscopy. In our scheme, optical trapping allows a mobile particle to be held at the focus of a laser beam for long-time observations and moved away from the coverslip up into solution. Furthermore, levitations of the particle in the bathing solution causes a reduction in fluorescence and stray scattering interference form the coverslip thus undermining the weak Raman signals from the cell. In addition, optical trapping permits optimum excitation and collection for Raman scattering in a confocal configuration because the particle is maintained in the focus of the laser beam and within the focal plane of the objective. Although some degree of photo damage to the living cells may be caused by the trapping beam, this effect can be decreased substantially by using low power and selecting near-infrared laser wavelength [9,18,19]. Thus the vibration spectrum has promising potential as an analytical tool for diagnosing cancer because it can probe the chemical composition and molecular structure of the normal and pathological tissue, so that researchers have paid much attention to the field over the past decade. The Raman method has the relative lower requirement to the preparation of sample, thus ordinary biological sample such as cell, living tissue, DNA and RNA can be measured directly. The measurement has no damage to sample and a large amount of data can easily be obtained. Therefore, many application of Raman spectrum have been gained in biology, medicine, medicament analysis and filtration [6,7,11,20,21]. We studied the Raman spectra of GBM cell lines obtained from three white male patients of different age groups (HTB12, HTB14, HTB15) and one African American male patient (GBM A172). GBM A172 was analyzed in a separate set up using the same technique at Purdue University while the other GBM cell lines were analyzed at University of California-Davis Biophotonics Center. The results showed Raman spectra of brain cancer cells obtained are very similar in all of these cells of same stage cancer as analyzed at the two laboratories. The results indicate that Raman spectra may offer the experimental basis for cancer diagnosis and treatment especially for brain cancer where cancer diagnosis and staging is difficult by invasive procedures. Light interacts with tissue in a number of ways including, elastic and inelastic scattering, reflection and absorption, leading to fluorescence and phosphorescence. These interactions can be used to measure abnormal changes in tissue. Initial optical biopsy systems have potential to be used as an adjunct to current investigative techniques to improve the targeting of blind biopsy. Future prospects with molecular-specific techniques may enable objective optical detection providing a real-time, highly sensitive and specific measurement of the histological state of the tissue. Raman spectroscopy has the potential to identify markers associated with malignant change and could be used as diagnostic tool for the early detection of precancerous and cancerous lesions in vivo. The clinical requirements for an objective, non-invasive, real-time probe for the accurate and repeatable measurement of pathological state of the tissue are overwhelming. Thus Raman spectroscopy will develop as a future tool for early cancer diagnostics and staging.

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### References


