Differentiation of peripheral nerve functions and properties with spectral analysis and Karnovsky-Roots staining: a preliminary study

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Abstract: Objective: The purpose of this study was to explore the possibility for analyzing and differentiating between motor and sensory functions of peripheral nerve axons using spectral technology. Methods: 10 μm slide section of S1 anterior and posterior rabbit spinal nerve roots were made and then stained with Karnovsky-Roots method for molecular hyperspectral imaging microscopy analysis. In addition, Raman spectra data of nerve axons on each slide was collected after Karnovsky-Roots staining for 30 minutes. Results: Motor axons were differentiated from sensory axons in a nerve axon section hyperspectral image via Spectral angle mapper algorithm. Raman scatterings could be detected near 2110 cm⁻¹ and 2155 cm⁻¹ in motor axons after Karnovsky-Roots staining. The value of I₂₁₀₀/I₁₄₄₀ in motor axons are significantly different (P<0.001) than in sensory axons after staining for 30 minutes. Conclusions: Motor and sensory nerve axons can be differentiated from their counterparts in 30 minutes by using Raman microspectroscopy analysis assisted with Karnovsky-Roots staining.

Keywords: Peripheral nerve, molecular hyperspectral imaging, Raman spectroscopy

Introduction

Peripheral nerve injury is one of the most common causes for deficiencies in motor and sensory functions leading to loss of labor. The repair and reconstruction of injured peripheral nerves has always been an important issue. One of the reasons for lacking efficacy of nerve repair is attributed to the mismatch of nerve stump orientation and its motor and sensory components.

Cholinesterase staining is currently one of the most frequently used intra-operative methods in clinical practice. The Karnovsky-Roots staining protocol requires at least 8-12 hours for most AChE positive areas to properly manifest, which is time consuming and unsuitable for operation. Many surgeons have attempted to modify the formula to speed up the staining procedure. However, without enhancement in the basic principles, the sensitivity and specificity of the differentiation method remains unchanged.

The application of spectral analysis in medical science has been growing over recent years. Spectral technologies used in this study were molecular hyperspectral imaging (MHI) and Raman microspectroscopy. MHI integrates conventional imaging with spectroscopy and is capable of obtaining both spatial and spectral information from a specimen, a technique which enables investigators to analyze the chemical composition of traces while simultaneously visualize their spatial distributions [1]. MHI has already been applied in monitoring cancer [2], hemodynamics [3] and diabetic foot ulcers [4] as spectra of diseased cells deviated from their normal exhibitions. Preliminary research has already been successful regarding to the ability to describe nerve morphometry
Spectral differentiation of nerve functions

with MHI [5]. Raman microspectroscopy is a label-free method for rapid and sensitive detection of the status and changes in cells biomolecular microenvironments and is capable of both in vivo and ex vivo study. It has been used in detecting cancer cells [6], stem cells [7], inflammation [8], etc., based on recognizing specific Raman signatures expressed by certain radical groups.

We hypothesized that Karnovsky-Roots stained motor axons would exhibit different hyperspectral plots and Raman spectra from their sensory counterparts since the staining process would alter the chemical composition of the samples and generate its own unique spectral signals. The purpose of this study aimed to take advantage of the qualitative analyzing attributes of spectral technologies to identify and differentiate between motor and sensory functions and histological properties of peripheral nerve axons.

Methods

Twenty New Zealand rabbits (ordinary grade, 2.5 kg, random sex) were sacrificed through venous air injections for the bilateral harvest of their S1 anterior and posterior spinal nerve roots which strictly contained either motor or sensory peripheral nerve axons.

Three random samples from both anterior and posterior groups were separately cryosectioned into 10 μm thick slides with CM1850 cryostat (Leica, Germany), Karnovsky-Roots stained for four hours and viewed under a custom assembled molecular hyperspectral imaging system (Li, et al. [9], magnification 40x) with ENVI 4.0 software (Exelis, USA). Fifteen regions of interest (ROI) of motor and sensory axons and myelin sheaths were manually drawn over each gathered image to acquire their respective hyperspectral curves for computer analysis. Spectral angle mapper (SAM) algorithm was used to differentiate between the two kinds of nerve fibers and their sheaths.

Twenty anterior and twenty posterior root samples were randomly chosen and made into 30 μm frozen sections. Sections of every sample were scanned with Raman micro-spectroscopy (HORIBA Jobin Yvon LabRam-1b, 633 nm laser, 4.3 mW, 400 × magnification, France) after 30 minutes of conventional Karnovsky-Roots staining. For each slide, a blank background was also scanned. Raman spectra acquisition range was set for 400 cm\(^{-1}\) to 2400 cm\(^{-1}\) and time for 50 s. Raman spectra data were manually adjusted to eliminate background noise and fluorescence peaks in Labspec 3.0 (ASD, USA) by a single experienced Raman technician. Independent samples t-test was performed on \(I_{1210}/I_{1440}\) values of each slide with SPSS 19.0 (IBM, USA). A classification algorithm was formulated from the statistic results and was put to use in a diagnostic test performed with the remaining twenty anterior and twenty posterior spinal nerve roots, comparing our method against anatomical proof which was considered as “the golden standard”. Raman spectra were drawn with OriginPro 7.5 (Microcal, USA).

The staining protocol adopted in our study was the conventional Karnovsky-Roots acetylcholinesterase staining method. The incubation solutions were created and mixed 30 minutes before staining in the following order: 0.1 M Acetylthiocholine iodide 20 ml, 0.1 M pH = 6.5 phosphate buffer 160 ml, 0.1 M sodium citrate 10 ml, 30 mM copper sulfate 25 ml, 5 mM potassium ferricyanide. Staining was completed under room temperature.

Results

Karnovsky-Roots stained motor and sensory hyperspectral plots had respective characteristics (Figure 1). They were somewhat different and were able to be distinguished with ENVI’s Spectral Angle Mapper and shown on synthesized false-color images over the original section images (Figure 2).
Raman spectra of motor axons showed Raman peaks in 2100 cm$^{-1}$ and 2155 cm$^{-1}$ whereas spectra of sensory axons remained near baseline value (Figure 3). We defined $I_{2100}$ and $I_{1440}$ as the maximum value of Raman scatterings in respectively 2100 cm$^{-1}$ to 2160 cm$^{-1}$ and 1425 cm$^{-1}$ to 1455 cm$^{-1}$. The value of quotient $I_{2100}/I_{1440}$ was analyzed with independent sample t-test. The average $I_{2100}/I_{1440}$ was higher ($t = -0.606$, p<0.05) in motor spectra (1.54 ± 1.11) than in sensory spectra (0.03 ± 0.03). ROC curve was calculated and cut-off point was between 0.17 and 0.33, where sensitivity and 1-specificity were respectively 1 and 0 at 0.17, 0.95 and 0 at 0.33. Therefore we chose an intermediate one-digit decimal of 0.2 as the cut-off.

Based on our criterion of classification, the Raman spectrum of any nerve axon would have to have a value of $I_{2100}/I_{1440}$ higher than 0.2 to be classified as motor. Twenty anterior and twenty posterior nerve root samples were committed in a diagnostic test comparing quick staining and spectral analysis method against
the golden standard of anatomical/histological characteristics which suggest anterior roots be comprised of motor nerves and posterior roots of sensory nerves. All twenty anterior roots and nineteen posterior roots were correctly identified. One posterior root was false-positive. The sensitivity, specificity and accuracy of our method were 100%, 95%, and 97.5%, respectively.

Discussion

In hyperspectral imaging, myelin sheaths and nerve axons had very different hyperspectral plots and were easily identified with Spectral Angle Mapper (SAM). SAM is an algorithm that compares the cosine values between two spectral plots and determines whether they were similar or not. Our finding of myelin sheaths’ distinct plots would be useful for histological and pathological studies as MHI may negate the need for myelin staining (i.e. Sudan black, Luxol blue, silver or potassium dichromate) which are extremely time-consuming and return with low specificity and contrast.

The stain intensities are not always equal in motor nerve fibers. Therefore, the hyperspectral plots of stained motor axons may vary in a certain pattern. As shown in Figure 5, unstained motor axons exhibit an “S” shaped hyperspectral plot and through the increase of stain intensity the plot gradually became “U” shaped while the intensity values decreased.

One single false-positive result can be explained by non-specific staining of butyrocholinesterase (BuChE). Posterior spinal nerve roots contain a small amount of unmyelinated C fibers which appear scattered between the myelinated fibers in a spot or fleck form. Unmyelinated C fibers exhibit BuChE activity and are more intensely stained than myelinated fibers’ AChE under the same circumstance [12]. It has been reported that iso-OMP can suppress BuChE staining and can be added to staining process if higher specificity is desired [13].

There were limitations in our study. First, we were unable to avoid sample collection and sectioning, although this was merely an attempt to test whether spectral analysis was capable of differentiating similar cells with different functions. Secondly, only rabbit spinal nerve roots were used as specimen as they consisted mostly of either motor or sensory axons. Under such unification, a nerve sample would represent any single one of its axons. Human periph-
eral nerve samples are either mixed or sensory nerves. Therefore, any effort made to recognize and match a single specifically targeted axon from an entire nerve cross section would have been overwhelmed. In further studies, we plan to test our algorithms in mixed nerves under Raman spectral imaging systems.

In conclusion, we have made preliminary efforts in differentiation of motor and sensory nerve axons via Raman spectroscopy and hyperspectral imaging. Staining was conducted for four hours before MHI could properly detect motor spectra, and thirty minutes for Raman spectroscopy. These findings provide evidence for further investigation of utilizing spectral analysis to identify nerve fascicle properties and hopefully could achieve in situ spectral identification and differentiation.

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Disclosure of conflict of interest

None.

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