Raman Spectroscopy of Algae: A Review

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Abstract

Algae are eukaryotic microorganisms which contain chlorophyll and are capable of photosynthesis. In various studies, Raman spectra have been used to identify a particular genus in a group of different types of algae. Each biomolecule has its own signature Raman spectrum. This characteristic signal can be used to identify and characterize the biomolecules in algae. Raman spectrum can be used to identify the components, determine the molecular structure and various properties of biomolecules in algae. With this view, this work presents a comprehensive review of current practices and advancements in Raman spectroscopy of Algae as well as in Raman spectroscopy of component biomolecules of different genus of Algae.

Keywords: Raman spectroscopy; Algae; Biomolecules; Bio-analysis; Identification

Introduction

Raman spectroscopy has proven to be a powerful and versatile characterization tool used for determining chemical composition of material systems such as nanoscale semiconductor devices or biological systems. Raman spectroscopy is based on the concept of the Raman effect [1]. The principle behind the Raman Effect is based on the inelastic scattering of incident photons by atoms and molecules in a sample. The incident photons enter a virtual energy state when they interact with sample. The eventual return of photons to ground state results in the inelastic scattering. The wavelength of scattered photons can be determined by calculating the induced dipole moments in molecules due to vibrational displacements. If the final ground state has more energy than initial state, then emitted photon will be shifted to lower frequency. This scattering is called as Stokes scattering. However, if the final state is more energetic than initial, the emitted photon will be shifted to higher frequency, resulting in anti-Stokes scattering. Depending on the amount of scattered photons, Raman spectrum shows various peaks which undergo changes with changes in the characteristics of a sample. These characteristic peaks can be used to identify the structural components or chemical composition of the sample.

One major advantage of Raman spectroscopy in the case of biological molecules is that water gives very weak, uncomplicated Raman signal [2]. Biological systems are essentially wet systems hence Raman spectrum of a biological system can be easily obtained by filtering the water’s Raman signal. Another advantage of Raman spectroscopy in the case of biological molecules is the ability of Raman spectroscopy to analyze in-vivo samples [3]. Raman spectroscopy generally does not require sample preparation for obtaining the response from varied biological samples such as algae cells [3]. This aspect gives this technique an edge over other methods such as Infra-Red (IR) spectroscopy which requires elaborate signal preparation for excitation and complex instrumentation for signal processing after the excitation [3]. Raman spectroscopy is fast emerging as an important characterization tool for biological systems. With this view, the present review focuses on presenting information on advancements made regarding the Raman spectroscopy of Algae.

Algae are eukaryotic microorganisms which contain chlorophyll and are capable of photosynthesis. Algae are a rich source of carbohydrates and other nutrients [5]. Some of the important algal extracts are used in food, cosmetics, and pharmaceutical industry [5]. Algae are known to show changes in composition or the structure depending on the changes in environmental conditions [3]. Based on such attributes, algae have led to new development in applications such as sensing elements in biosensors [6-15]. Algae cells have also been used as an aid in controlling water pollution and heavy metal pollution [10,12,13,15]. One of the most important and visible uses of algae has been in the domain of biofuel development due to superior lipid content in algae cells compared to other plant cells [16,17].

Algae cells contain mainly five types of biomolecules: proteins, carbohydrates, lipids, nucleic acids, and pigments [18]. Each type of biomolecules has been shown to have its own characteristic signature Raman spectrum. Characteristic peaks of each biomolecule can be used to identify them from Raman spectrum of algae cell. Raman spectrum of an algae cell will be the sum of Raman spectra of its constituent biomolecules [16]. In various studies, Raman spectra have been used to identify a particular type of algae species in a group of different types of algae or in different environmental conditions [19-22]. Raman spectra of individual biomolecules can be used to determine the molecular structure and various properties of the biomolecules and hence have also been studied extensively. Figure 1(a) shows the overview of applications of Raman spectroscopy in study of algae. There have been several reviews of applications of Raman spectroscopy in bioanalysis [4,23-24], however none of the available review deals with Raman spectroscopy of algae cells.

This present review is divided in 5 parts. Introduction is provided in section 1. In section 2, the basic instrumentation required for performing Raman spectroscopy is presented along with recent development of advanced methods such as Surface Enhanced Raman Spectroscopy (SERS). In section 3, various studies pertaining to identification of algae species using Raman spectroscopy are reviewed. In section 4, studies of component biomolecules of algae using Raman spectroscopy are presented. Conclusion and future prospects are presented in section 5.

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**Instrumentation**

Schematic of a typical Raman setup is shown in Figure 1(b). In a typical Raman experiment, a laser beam excitation is provided by a laser source such as an Argon ion laser or Nd:YAG laser. Wavelengths of laser lines of Argon ion laser and Nd:YAG lasers are most common incident wavelengths used to study algae Raman response. These wavelengths include 488nm, 514.5nm and 1024nm among other wavelengths. The laser signal is focused on the sample that is to be analyzed using optical systems such as a microscope and lens-mirror assembly and scattered response from the sample is recorded. The scattered response has wavelength higher or lower than that of the original laser depending on whether the scattered response follows Stokes or anti-Stokes scattering. The scattered beam is directed through a series of filters to obtain Raman response in the form of a spectrum of lines with varying intensity as a function of wavelength. This signal is then recorded on a computer for further processing and analyses [25].

Several advances in instrumentation have resulted in the more applications of Raman spectroscopy in biological samples [4]. Better lasers, filters, and fiber optics have improved Raman signals because of improvements in excitation of sample and better acquisition of response signals [4]. Incorporation of diode laser as optical pump in Nd:YAG lasers has provided better pump stability and hence reduced flicker noise, which increases overall signal to noise ratio [4]. Ti:Sapphire and other solid state lasers are also used as excitation sources for Raman

**Figure 1:** (a) A schematic of experimental set-up of a typical micro-Raman spectrometer and (b) a schematic showing applicability of micro-Raman spectroscopy to different aspects of Algae.
Raman spectrum of Na alginate: system hence making hardware gives better spatial resolution and portability to the Raman size and increased the flexibility of the Raman systems [4]. Improved optical excitation and collection of response signal have decreased the biological samples. Incorporation of fiber optics for delivering the rastering [4]. Liquid crystals have high throughput, large spectral range, optical element which produces 2-D images without surface or beam plasmons in metal structures and second, intensity increase due to electromagnetic enhancement correlated with the excitation of surface features [28]. The enhancement in response signal intensity is because of two different mechanisms. First, increase in the intensity due to electromagnetic enhancement correlated with the excitation of surface plasmons in metal structures and second, intensity increase due to chemical enhancement related to the adsorbate-substrate complexes providing different orbitals for excitation of the Raman processes [28]. The enhancement in intensity of the response can also be obtained using the extent of increase in intensity of each Raman mode [27]. The enhancement in intensity of the response is not observed for any other metal colloids or surfaces having large features [28]. The enhancement in response signal intensity is because of two different mechanisms. First, increase in the intensity due to electromagnetic enhancement correlated with the excitation of surface plasmons in metal structures and second, intensity increase due to chemical enhancement related to the adsorbate-substrate complexes providing different orbitals for excitation of the Raman processes [28]. Surface Enhanced Resonance Raman Spectroscopy (SERRS) combines the advantages offered by Resonance Raman spectroscopy and surface enhancement. SERRS can be used for higher specificity in excitation spectroscopy because of their high harmonic conversion efficiencies and ability to tune them to different wavelengths [4]. Along with advances in laser systems, tunable filters such as liquid crystal and acousto-optic filters have introduced electronically controlled, fixed optical element which produces 2-D images without surface or beam rastering [4]. Liquid crystals have high throughput, large spectral range, and fast scan times making them ideal for use in the spectroscopy of biological samples. Incorporation of fiber optics for delivering the optical excitation and collection of response signal have decreased the size and increased the flexibility of the Raman systems [4]. Improved hardware gives better spatial resolution and portability to the Raman system hence making in-vivo analyses easier.

Several new discoveries in the physics of Raman Effect have resulted in the advances in the Raman spectroscopic technique for bioanalysis. Resonance Raman spectroscopy uses the lasers which have their energy adjusted such that the energy of the laser or the that of the scattered signal coincides with electronic transition energy of a particular molecule or crystal [26]. This technique has become more prevalent with the advances in tunable lasers [4]. In biological samples, the advantage of resonance Raman spectroscopy is that the only modes associated with specific chromophoric group of molecules are enhanced due to resonance effect [26]. Resonance Raman spectroscopy makes it possible to study particular molecule in the algae cell thus enabling analysis of individual component from cell. Recently, there have been some major advances in Raman spectroscopy techniques to analyze the biological samples. These include Surface Enhanced Raman Spectroscopy (SERS), Tip Enhanced Raman Spectroscopy (TERS), Coherent Anti-Stokes Raman Spectroscopy (CARS) and Laser Tweezers Raman Spectroscopy (LTERS).

SERS has become important Raman spectroscopic analysis tool because it offers 10^3 to 10^5 enhancement in the intensity of the Raman response signal of an analyte when the analyte is adsorbed on the surface of some noble metals with nanoscale features [23]. Surface enhancement permits a single molecule detection as large intensity enhancement in response signal makes the response signal detectable even when response is obtained from a single molecule [23] and improves the spatial resolution to lateral resolutions better than 10 nm [27]. Information about the surface-interface processes can also be obtained using the extent of increase in intensity of each Raman mode [27]. The enhancement in intensity of the response is not observed for any other metal colloids or surfaces having large features [28]. The enhancement in response signal intensity is because of two different mechanisms. First, increase in the intensity due to electromagnetic enhancement correlated with the excitation of surface plasmons in metal structures and second, intensity increase due to chemical enhancement related to the adsorbate-substrate complexes providing different orbitals for excitation of the Raman processes [28]. Surface Enhanced Resonance Raman Spectroscopy (SERRS) combines the advantages offered by Resonance Raman spectroscopy and surface enhancement. SERRS can be used for higher specificity in excitation...
thus opening interesting opportunities for studying specific parts of large biomolecules [27]. SERS and SERRS make it possible to analyze single cell of the algae. It is also possible to study individual molecules in algae cells using SERS.

The concept of TERS is based on the concept of “hot spots” in surface Raman spectroscopy [28]. This concept states that the surface enhancement is produced by a small Raman active areas called as “hot spots” on the surface and rest of the surface is inactive [28]. In TERS, single such hot spot is created by keeping a sharp protrusion or a tip made of gold or silver at a small distance from the sample [28]. The enhancement in intensity in TERS is because of localized enhanced electromagnetic field near the tip apex [28]. The enhanced electromagnetic field provides the enhancement in the intensity of Raman spectrum like in SERS. Currently, TERS provides two to four orders of magnitude increase in response signal intensity. The response intensity enhancement is expected to increase further on improvements in excitation and radiation efficiency in surface plasmons [28]. TERS provides better spatial resolution compared to traditional Raman spectroscopy due to local enhancement provided by hot spots thus making the single cell analysis possible.

In CARS, two high powered (generally pulsed) laser beams are focused together on the sample. Due to the mixing of two lasers, a coherent beam resembling low intensity laser beam is generated [29]. Due to the coherence of the beam, the spectrum obtained using CARS is orders of magnitude stronger than Raman spectrum obtained using traditional Raman spectroscopy [29]. Two interacting beams give high 3D sectioning capability [23]. CARS signals can also be easily picked in fluorescent background due to blue shift of the response signals [23].

LTRS utilizes laser tweezers to immobilize a single cell of algae or any other organism [17]. LTRS can be used to obtain the Raman spectrum of single cell or even a particular part of an immobilized algae cell [17].

Advances in the spectroscopic techniques have opened a new frontier of very high resolution Raman spectroscopy, even of the specific parts inside the cell. They also enable the monitoring of real time changes in algae cells using Raman spectroscopy thus improving the ability to detect the changes in algae cells in different environments. In the next section, a summary of the studies done on the identification of algae species using Raman spectroscopy is provided.

**Identification of Algae Species using Raman Spectroscopy**

Several studies focusing on the identification of algae using Raman spectroscopy have been reported in literature [19-22]. The technique of identification of algae uses the characteristic peaks in the Raman spectra to identify unique biomolecules in the cells which are correlated to the specific species of algae [19,20].

Identification process often involves analysis of Raman spectra from multiple components of cells. Analysis of multi-component systems is performed using statistical multivariate analysis. Most of the times,
linear analysis is used [30]. Two basic assumptions are made in linear analysis. First, Raman spectrum of mixture of biomolecules is assumed to be linear superposition of component spectra of biomolecules in mixture [30]. It is also assumed that signal intensity and concentration of biomolecule in mixture has linear relationship [30]. For finding major components, either explicit or implicit methods are applied. In explicit methods, such as ordinary least squares or classical least squares, Raman spectra of all component biomolecules are previously known beforehand and hence implicit analysis is performed to analyze the components. Principal component analysis [32] and partial least squares, Raman spectra for components are not known explicitly and hence implicit analysis is performed to analyze the major components, either explicit or implicit methods are applied. Statistical analyses help in extracting the information from Raman spectrum and analyze the component mixtures effectively.

In reference [22], Raman spectroscopy is applied to differentiate between non-toxic and toxic algal strains. Four different species of algae including *Pseudo-nitzschia*, some of which are capable of producing toxin domoic acid, are studied by the means of resonance Raman spectra excited at 457.9 and 488 nm. It was observed that Raman spectra for all algae species contain major peaks near 1000 cm\(^{-1}\), 1153 cm\(^{-1}\), and 1523 cm\(^{-1}\), all of which are strongly enhanced due to carotenoids [22]. Features between 920-980 and 1170-1230 cm\(^{-1}\) are relatively weaker and are more characteristic of algae species [22]. It was observed that all *Pseudo-nitzschia* species produce a Raman response signal which is different from other species [22]. In the study by Brahma et al. [19], various Marine algae species were identified from the medium containing different algae species using resonance Raman spectroscopy. In reference [20], different types of seaweeds

### Table 1: Studies on biomolecules in algae using Raman spectroscopy.

| Biomolecule | Algae species studied | Observations | Conclusions | Ref |
|-------------|-----------------------|--------------|-------------|-----|
| **Proteins** |                       |              |             |     |
| Adhesion proteins | Coccomyxa sp. and Glaphyrya trebouziodes | Amide III band peaks at 1224 cm\(^{-1}\) and 1260 cm\(^{-1}\) | Consistent with generic amyloid structure with strong hydrophobic core | 55 |
| Hemoglobin | C. Eugametos | Fe-CN ferric derivative peak at 440cm\(^{-1}\) | Fe-C-N moiety adopts highly bent structure due to H bonding | 38 |
| **Carbohydrates** |                       |              |             |     |
| Polysaccharides |                       | Peaks in the range 350-600 cm\(^{-1}\) | Skeletal pyranose ring modes | 56,44 |
| *β*-D-glucosides |                       | 377 cm\(^{-1}\) characteristic peak |            | 56,44 |
| *α*-D-glucosides |                       | Strong peaks from 479 to 483 cm\(^{-1}\) | Peaks from Amylopectin and amyrase | 56,44 |
| Alginites |                       | Peaks at <1300 cm\(^{-1}\) | Vibration of polymer backbone | 46 |
| Calcium Alginate |                       | Change of band position from alginites | Symmetric COO- stretching peak | 46 |
| *ε*-carrageenan and *η*-carrageenan | Eucheuma cottonii Eucheuma spinosa | Various peaks from 700 to 1200 cm\(^{-1}\) | Peaks are assigned to respective molecular structure of carrageenans | 48 |
| **Lipids** |                       |              |             |     |
| Hydrocarbons (Figure 3) | Botryococcus braunii | 1650-1670 cm\(^{-1}\) and 2800-3000 cm\(^{-1}\) peaks | Double bond stretching peak in long chain unsaturated hydrocarbons | 57 |
| Chlorella sorokiniana Neochloris oleoabundans | 1650 cm\(^{-1}\) peak 2800-3000 cm\(^{-1}\) peak | C=C stretching peak C=C-H vibration peak | 16 |
| Botryococcus sedetricus Chlamydomonas sp. Trachydiscus minutus | 1656 cm\(^{-1}\) peak 1445 cm\(^{-1}\) peak | cis C=C stretching mode proportional to no. of unsaturated bonds CH\(_1\), scissoring mode proportional to no. of saturated bonds Trachydiscus minutus has significantly higher content of unsaturated fatty acids | 56 |
| **Pigments: Chlorophylls** |                       |              |             |     |
| Chlorophyll a (Figure 4) |                       | Peaks in the range 1100-1600 cm\(^{-1}\) | Stretching motions of C-C and C-N bonds Planar deformations of tetrapyrrol macrorycle Mg related vibrations Mg atoms in hexacoordinated state | 35 |
| Chlorophyll b (Figure 5) |                       | 1361 cm\(^{-1}\) peak | C-N ring breathing mode | 58 |
| Chlorophyll c (Figure 6) | Acaryochloris marina | Very different Raman response as compared to Chlorophyll a and b | Presence of formyl group at C-3 position | 37 |
| **Pigments: Carotenones** |                       |              |             |     |
| *β*-carotene (Figure 7) | Chlorella sorokiniana Neochloris oleoabundans Euglena, Chlamydomonas | Peak at 1520 cm\(^{-1}\) Peak at 1000 cm\(^{-1}\) Strong peak in the eyespot region | C=C stretching mode C-C stretching mode C=CH\(_3\) stretching mode High content of carotenoids in eyespot region | 59, 16, 51-52, 60 |
| **Nucleic acids** |                       |              |             |     |
| DNA and RNA |                       | Peaks between 600 cm\(^{-1}\) and 800 cm\(^{-1}\) Intense peak at 1671 cm\(^{-1}\) Peak at 1100 cm\(^{-1}\) | Ring breathing modes of DNA and RNA bases C=O stretch vibrations Symmetric PO\(_2\) stretching vibration | 53 |
were identified using Fourier Transform based Raman spectroscopy (FT-Raman). It was observed that FT-Raman spectra have higher resolution than FTIR spectra hence it was concluded that FT-Raman spectroscopy is better for identification of different species from a medium containing different species of algae. In reference [21], Algae and bacteria deposits on Ti sheets are detected using Surface enhanced Raman spectroscopy. Spatial distribution of algae on the surface was also identified.

Raman spectroscopy provides a high resolution method with high accuracy for identification and detection of algae species. It can be used to detect the algae species from the biomass coatings or in algal blooms. With smaller and portable equipments, it is possible to detect the algae species in their natural habitat [33]. Recent developments in instrumentation and spectroscopic techniques such as SERS will further help in identification of algae. The identification of algae using Raman spectroscopy is based on the identification of component biomolecules. Hence it is important to study the individual biomolecules using Raman spectroscopy. These studies are summarized in the next section.

**Study of Component Biomolecules**

Algae cell contains mainly five types of biomolecules: proteins, carbohydrates, lipids, nucleic acids and pigments [18]. Raman spectroscopy has been used to find the molecular structure of the biomolecules [34-43] and to analyze other properties such as the location of biomolecules in the cell [16,44-52]. Various methods are used to predict the molecular structure of biomolecules using Raman spectroscopy. In isotope labeling, certain atoms in molecule are replaced by their isotopes and changes in the Raman spectra are observed for the new isotope labeled molecules. The bands which show differences from original spectra are assigned to the labeled parts of the molecule. In site-directed mutagenesis, specific sites of cells are modified using mutation. The changes in Raman spectra can then be assigned to the mutated parts of cells and hence the biomolecules that are present in mutated parts or to the changes in biomolecules due to mutations. In normal coordinate analysis, the vibrational modes are analyzed theoretically using quantum mechanics of coupled harmonic oscillators [54] and each mode is assigned to a peak in Raman spectrum. Now, because of advances in high performance computing, it is possible to find the vibrational modes associated with each peak in Raman spectrum of each biomolecule using molecular simulations such as density functional theory.

The reference Raman spectra for important biomolecules are collected in [53]. The reviews on the use of Raman spectroscopy in the analysis of biomolecules are presented in various references [4,23,24]. The summary of Raman spectroscopic studies of each type of biomolecule in algae is presented in table 1.

**Conclusions**

Raman spectroscopy is an important tool in the analyses of algae cells and component biomolecules. It is an ideal experimental measurement system for the characterization of “wet” biosystems on the account of weak Raman signal of water. It also provides high spatial and temporal resolution required for the characterization of biological samples at cellular level. Raman spectrum also has a lot of information focusing on vibration states of the molecules. Any single Raman spectrum can be used to extract large amount of data about the systems including components of system and temperature of system. Using the information contained in the spectrum, various algae species can be identified and analyzed for various properties such as locations of particular biomolecule in cell.

With advances in spectroscopic techniques in the form of SERS, TERS, CARS and LTRS, it is now possible to obtain Raman spectra of specific parts of a cell. This will enable better understanding of particular parts of cells and hence the constituent biomolecules. Use of pulsed lasers has enabled monitoring of the changes in system with respect to time. In future, Raman spectroscopy can be used to map the real time changes in the biological systems with very high spatial resolution and also very high accuracy.

Raman spectroscopy can expand its application base in future including clinical diagnosis because of various advantages. The most important applications of Raman spectroscopy in the study of algae in future could well be for studying in-vivo samples and for the analysis of algae cell metabolism in real time.

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