Characterization of Microorganisms Using Raman Microscopy

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Abstract

Raman spectroscopy has recently gained popularity as an attractive approach for the biochemical characterization, rapid identification, and an accurate classification of a wide range of prokaryotes and eukaryotes organisms. In the case of eucariotes it is necessary to obtain higher number of Raman spectra in order to perform statistical analysis and to draw conclusions.

“Raman spectroscopy (RS) is a powerful molecular fingerprinting technique which analyzes materials through the interaction of the material’s molecules with an incident laser beam” (Hanlon et al., 2000).

Vibrational spectroscopic technique, Raman spectroscopy (RS), has been used extensively to identify samples of different microorganisms by a careful investigation of the vibrating modes of the molecules in the microorganisms (Rösch et al. 2005). Raman spectroscopy has recently gained popularity as an attractive approach for the biochemical characterization, rapid identification, and an accurate classification of a wide range of prokaryotes and eukaryotes organisms (Hamasha 2011).

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The Raman spectra of the microorganisms are superposition of spectra of the biochemical components inside the cells like e.g. protein, DNA, RNA, lipids, carbohydrates, water, as well as a few components with minor concentrations (Rösch et al. 2011).

Accordingly, the Raman spectra of two different species or strains show minor variations which originate from different chemical compositions due to variations in e.g. the cell wall (Rösch et al. 2011).

For the Raman spectroscopic characterization of eukaryotes like yeasts or fungi, different approaches are mandatory. It is not recommendable to use only one Raman spectra, in case of eukaryotes, because of the variations due to various organelles. An average of fifty spectra is necessary in order to perform statistical analysis and to draw conclusions. On this basis, it can be concluded, that the Raman spectroscopy can be used to identify yeasts or fungi (Stöckel et al. 2015).

For example, distribution of the width of the bands mirrors the different compounds and parts of the yeast cell (Figure 1a). Characteristic C = O stretch vibrations ($1731–1765 \text{ cm}^{-1}$) represent the lipid fraction; mapping over the amide I region ($1624–1687 \text{ cm}^{-1}$) produces bands arousing from the C= C lipid bonds. The phenylenic C = C Raman band ($1567–1607 \text{ cm}^{-1}$) can be only seen in the periphery of the cells (Rösch et al., 2005).

During recording of a spectrum problems can occur. Fluorescence often appears when the examined material is complex and in color (Figure 1b), (Jang and Akkus, 2013). Another problem that might appear is burning of a sample (Figure 1c).

1 Preparation of the sample

1.1 Yeasts

The yeast cells were incubated at 28°C in a nutrient-rich YPD medium. Aliquots of the cell suspension were centrifuged (3000 rpm, 2min), washed three times with sterile water, and final suspended in the new aliquot of water (original suspension: water = 1:9). Spectra of the yeast (Fig. 1a).

Acquisition parameters

Recommendations for the spectra recording:

- Laser wavelength: 532 nm, Greeting: 1200 gr/mm, Slit: 50 µm, Hole: 500 µm,
- Acquisition time: 20s, Range: 400–3200 cm$^{-1}$. Use a quartz plate!

1.2 Bacteria

The bacteria cells were incubated at 30°C–37°C in an appropriate nutrient-rich medium.

Sample preparationis the same as for the yeasts!
Figure 1a: Spectrum of the yeast. Different pronounced bands originated from lipids and proteins can be observed.

Figure 1b: Spectrum of fluorescence is marked by red label.
Acquisition parameters

Recommendations for shooting:

- Laser wavelength: 532 nm, Greeting: 600 gr/mm, Slit: 50 µm, Hole: 500 µm,
- Acquisition time: 30s, Range: 400–3200 cm$^{-1}$. Use a quartz plate!

In addition: All obtained spectra have to be processed using R program, which includes: spike removal, calibration, background removal, cutting, vector normalization, removing of the silent region.

2 References


Figure 1c: Spectrum of burned cells is presented by red label.


