Correlation mapping: rapid method for identification of histological features and pathological classification in mid infrared spectroscopic images of lymph nodes

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Abstract. In this work, a novel technique for rapid image analysis of Fourier transform infrared (FTIR) data obtained from human lymph nodes is explored. It uses the mathematical principle of orthogonality as a method to quickly and efficiently obtain tissue and pathology information from a spectral image cube. It requires less computational power and time compared to most forms of cluster analysis. The values obtained from different tissue and pathology types allows for discrimination of noncancerous from cancerous lymph nodes. It involves the calculation of the dot product between reference spectra and individual spectra from across the tissue image. These provide a measure of the correlation between individual spectra and the reference spectra, and each spectrum or pixel in the image is given a color representing the reference most closely correlating with it. The correlation maps are validated with the tissue and pathology features identified by an expert pathologist from corresponding hematoxylin and eosin stained tissue sections. Although this novel technique requires further study to properly test and validate this tool, with inclusion of more lymph node hyperspectral datasets (containing a greater variety of tissue states), it demonstrates significant clinical potential for pathology diagnosis.

Keywords: correlation mapping; FTIR spectroscopy; lymph node; spectral imaging; orthogonality; colinearity; cancer.

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1 Introduction

Cancer is a disease where by normal cell division has become uncontrolled and unregulated through mutations in the cell’s DNA, which leads to the formation of a growth or lump of tissue made up of these abnormal tumor cells. These tumor growths can become malignant (cancers), gaining the ability to spread to nearby tissues and organs by invasion and metastasis. Cancer cells from this growth may also break off from the primary tumor and invade the bloodstream or lymphatic vessels to be carried to other parts of the body to multiply and form secondary tumors (metastases). The progression of a primary tumor to formation of a secondary tumor or metastasis carries a worse prognosis and can be associated with poor survival in most cancers. Therefore, lymph node sampling and pathological examination is important in assessing whether a primary tumor has spread or has the potential to spread to distant sites to develop secondary tumors.

The current gold standard of conventional lymph node histopathology staging and grading relies on morphological analysis (i.e., tissue architecture, cell shape, and staining features) by a pathologist. This is subject to significant inter- and intraobserver variation, and is also time consuming and difficult to automate. The introduction of immunophenotyping and molecular biology techniques has increased the number of objective parameters that help in distinguishing a normal sample versus a pathologic one. However, there is a need for a more sensitive, fully automated, cost-effective and objective tool for screening of tissue slides in lymph node diagnostics. Fourier transform infrared (FTIR) imaging coupled to spectral orthogonality/correlation image analysis may make this a feasible option in the future. This is explored here.

FTIR microscopy has the potential to become such a screening technique. It is an optical technique that utilizes absorption spectroscopy, which can be employed to interrogate biological tissues to understand the changes in molecular constituents that are associated with cancer progression from normal to malignant during carcinogenesis. FTIR spectroscopy detects molecular bond vibrations within a sample via interrogation of the sample using the mid-infrared region of light, where the loss of light at each wavelength is measured as it is absorbed by the molecules in the tissue. Using the position and intensity of infrared absorption bands, particularly in the mid-infrared region, it has the ability to provide detailed biochemical information, since the spectrum obtained...
from cellular and tissue measurements is an intrinsic molecular fingerprint of the sample that reveals information about the nucleic acids, carbohydrate, protein and lipid content. Imaging methodologies employing FTIR spectroscopy are of particular interest because of their ability to provide qualitative and quantitative biochemical information from large areas of biological tissue. FTIR spectral imaging has already been applied in a number of studies using human tissue, including lymph nodes, prostate, lung, brain gliomas, bone, lymphoma, cervix, and breast.

Imaging studies involving cluster analysis have successively shown that it is possible to identify the different tissue layers in a tissue section as clusters. However, as observed by Romeo and Diem’s group, the acquisition of large IR datasets for each image exceeds 300 MB of memory at times, meaning that the correlation matrix required to be created in cluster analysis programs, such as hierarchical cluster analysis, can exceed 4 GB. The complex computations needed for generation of the different cluster matrices means that calculations require 64-bit processors with large memory access. In addition, the separation of tissue types in a cluster map for a particular tissue section and translation to other tissue sections cannot take place without the use of learning algorithms such as artificial neural networks, as demonstrated by Lasch et al.

The orthogonality/colinearity or cosine of the angle between two vectors (spectra) can be calculated using an operation called the inner or dot product of two entities, \( \mathbf{a} \) and \( \mathbf{b} \). To be orthogonal, the inner product of \( \mathbf{a} \) and \( \mathbf{b} \) must be 0. The inner product is one of the most fundamental operations in useful vector spaces. One can apply an orthogonality/colinearity measure in spectroscopy to determine how close one spectrum resembles another spectrum. The colinearity of a spectrum against another spectrum can be calculated using the equation:

\[
\mathbf{a} \cdot \mathbf{b} = |a||b|\cos \theta,
\]

where \( |a| \) and \( |b| \) denote the length (magnitude) of \( \mathbf{a} \) and \( \mathbf{b} \), and \( \theta \) is the angle between them.

In the equation, a resulting dot product value of 1 represents perfect colinearity, whereas a dot product value of 0 represents perfect orthogonality. This method effectively provides a value for the correlation of one spectrum with the next. To date, we have been unable to identify any previous tissue studies that used this mathematical correlation procedure with vibrational spectroscopy.

In this study, the dot product was used to compare how similar a set of reference or component spectra are to tissue spectra from a hyperspectral cube, and thus can be used as a tool to visualize regions in a sample having similar spectral properties. This helped to discriminate individual tissue types in a section compared to the hematoxylin and eosin (HE) stained slide. By extracting reference spectra to represent the main tissue types in a noncancerous lymph node, as well as tissue spectra from a cancerous lymph node, we were able to illustrate a novel technique to discriminate noncancerous from cancerous nodes using correlation maps.

2 Materials and Methods

2.1 Sample Preparation

During oesophagectomies to remove the primary tumors from patients (who also consented to donation of lymph nodes for spectroscopic studies), lymph node samples were selected and resected. Each lymph node was cut in half, where one half was placed in formalin and sent for routine histopathology (paraffin-embedded sectioning and standard tissue processing), while the other half was snap-frozen in liquid nitrogen and stored in −80 °C freezers for spectroscopic analysis.

Frozen unstained sections 15 \( \mu \)m thick were prepared using a cold cryotome and were mounted on a calcium fluoride (CaF\(_2\)) slide. Prior to spectroscopic measurements, these frozen sections were allowed to defrost and air-dry for at least 45 min.

Infrared spectra were collected using a Perkin Elmer Spotlight 300 Fourier transform infrared (FTIR) spectroscopy system in transmission mode. The system has a liquid-nitrogen-cooled linear mercury-cadmium-telluride (MCT) array detector, a video camera attached to a microscope to view optical images, and a programmable computerized \( x-y-z \) stage. Image maps (64 sample-background rationed coscans with aperture size of 100 \( \times \) 100 \( \mu \)m) within the wave number range 720 to 4000 cm\(^{-1}\) with a spectral resolution of 4 cm\(^{-1}\) were performed on the tissue sections.

2.2 Spectral Processing and Analysis

All data analysis was performed using Matlab\(^{7}\) software (MathWorks, Natick, Massachusetts). FTIR map spectra were converted and loaded into Matlab to generate a 3-D hyperspectral matrix for each section. These were then converted to absorbance using:

\[
\{\log(\text{max(transmittance)}) - \log(\text{transmittance})\},
\]

where transmittance = \( I_{out}/I_{in} \).

Infrared spectra were smoothed using a second-order Savitzky–Golay filter over 19 wavenumber data point spacing to remove short term variations or noise in the spectral dataset.

Two representative lymph nodes were selected for a negative noncancerous lymph node and a positive cancerous lymph node. After consultation with a histopathologist using the stained HE sections, areas representative of noncancerous tissue and of cancerous tissue were selected from the total intensity image of the hyperspectral matrix. An in-house Matlab script was developed that allowed the spectra from the areas of interest to be selected and extracted from the hyperspectral matrix of the two representative lymph nodes. From the noncancerous lymph node, areas representing normal cortex, follicle, medulla, capsule, and fat were selected and extracted; whereas areas of cancerous and remaining normal tissue were selected from the cancerous lymph node. A representative morphological-derived tissue spectral dataset was compiled for these tissue spectra. Each pathology-type dataset contained, on average, 3000 spectra. Each dataset was checked for the quality and colinearity of the spectra (the average colinearity within each dataset was \( \sim 0.92 \) with a standard deviation of \( \sim 0.03 \)). Using these tissue-derived datasets, a mean spectrum was calculated to represent each of...
Fig. 1 Extraction of representative tissue spectra from hyperspectral matrix of noncancerous (Co=cortex; M=medulla; CP=capsule; F=fat; Fo=follicle) and cancerous lymph node (Ca=cancerous tissue; Nm=noncancerous or remaining normal tissue).

Fig. 2 Performing the correlation calculation on a lymph node (cancerous) FTIR hyperspectral dataset.
the tissue types (Fig. 1), which could then be used in the
colinearity/correlation calculation with the lymph node hyper-
spectral datasets. A calcium fluoride spectra dataset was also
collected from the hyperspectral images, and another in-house
script was developed to filter out the calcium fluoride or non-
tissue pixels within the hyperspectral images so as not to com-
plicate the colinearity calculation and to provide better visu-
alization of the resulting correlation maps.

An in-house Matlab script was developed to perform the
correlation procedure. The script compares each morphologi-
cal representative tissue spectra to each spectrum in the hy-
perspectral map, and generates an array consisting of a value
0 to 1 (0 is no correlation and 1 is perfect correlation) for each
pixel of each representative tissue type. The program then
determines the highest (most correlated) value for each pixel
to determine which representative tissue type spectrum each
pixel most represents.

This orthogonality-correlation script, in combination with
the tissue type representative dataset, was then tested on six
noncancerous lymph nodes and seven cancerous lymph nodes.
The average time to run this correlation program on a hyper-
spectral image consisting of ~30,000 spectra was around
20 to 30 min, and required a standard 32-bit PC with 4-GB
RAM. Acquisition of the original 30,000 FTIR spectral im-
ages of the tissue section takes an additional 20 min approxi-
mately. Therefore, image acquisition and running of the or-
thogonality correlation should take under 60 min in total.

A correlation map was generated, displaying an image
where each pixel has a color assigned to it, indicating the
tissue or pathology type most representative for that particular
spectrum. Figure 2 illustrates this procedure.

3 Results and Discussion

Figure 3 illustrates the correlation maps generated from per-
forming the correlation algorithm on each of the 13 lymph
node hyperspectral cubes (six noncancerous and seven cancer-
ous). The cancerous lymph node correlation maps mostly
consisted of bright red (cancerous infiltrate representative spec-
tra) and some orange (medulla representative spectra), while
the noncancerous lymph node correlation maps mostly con-
sisted of blue (normal parenchymal tissue representative spec-
tra) and green (fat representative spectra) pixels.

In summary, this work demonstrates that correlation maps can be used to quickly (~25 min) and efficiently visualize regions of the sample, which have similar spectral properties to key reference spectra. These features correlate favorably with the features identified from the HE-stained slide by a pathologist. By using an in-house-developed orthogonality-correlation script, it was possible to take representative pathology/tissue type spectra and apply a similarity test to spectra of different lymph node samples for both noncancerous and cancerous pathology types. These were reconstructed into correlation maps with specific color coding for cancerous and noncancerous lymph nodes, demonstrating a novel pathological tool in lymph node diagnostics. This technique could be implemented in other tissue types for pathology classification.

While histopathology, which is based on tissue architecture, cell shape, and staining features, is the gold standard in lymph node sampling, it can also be time and labor intensive for pathologists and suffers from significant inter- and intraobserver variation. There is also the possibility of missing micrometastases, which can be difficult to identify in tissue architecture when using conventional HE staining. Immuno-histochemistry and other molecular biology techniques, such as reverse transcriptase polymerase chain reaction (RT-PCR), have helped to improve micrometastasis detection and are also semiautomated; however, these methods can also suffer from high false positive results as well as be expensive to perform routinely for lymph node sampling. Infrared spectroscopy is a nondestructive diagnostic tool requiring very little or no tissue processing (and therefore requires less labor and time costs). It also does not require the use of contrast agents and is based on the chemical characteristics of the tissue without the subjective analysis of a pathologist.

To validate this procedure, a more robust representative tissue dataset needs to be created from a larger number of lymph nodes from a variety of lymph node disease states. Further inclusion of more lymph node hyperspectral datasets are also needed to properly test and validate this tool; however, this initial study demonstrates real clinical potential for correlation maps.

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