Hyperspectral imaging in detecting dermal invasion in lentigo maligna melanoma

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N. Neittaanmäki1, M. Salmivuori2, I. Pölönen3, L. Jeskanen4, A. Ranki4, O. Saksela4, E. Snellman5, M. Grönroos2

(1) Department of Clinical Pathology, Sahlgrenska University hospital, Göteborg, Sweden
(2) Department of Dermatology and Allergology, Päijät-Häme Social and Health Care Group, Lahti, Finland
(3) Department of Mathematical Information Technology, University of Jyväskylä, Finland
(4) Department of Dermatology and Allergology, Helsinki University and Helsinki University Hospital, Finland
(5) Department of Dermatology, Tampere University and Tampere University Hospital, Finland

Corresponding author: Noora Neittaanmäki, MD, PhD,
Department of Clinical Pathology, Sahlgrenska University Hospital, Gula Stråket 8, 41345 Göteborg, Sweden
E-mail: noora.neittaanmaki@fimnet.fi

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**What’s already known about this topic?** Recently, various non-invasive imaging tools have been introduced for detection of skin cancers. Hyperspectral imaging is a novel promising technology, which combines spectroscopy and skin imaging and enables analysis of large skin areas at once.

**What does this study add?** This pilot study shows that hyperspectral imaging can be used in preoperative staging of lentigo maligna melanomas

**TO THE EDITOR**

Invasive lentigo maligna melanoma (LMM) and its *in situ* precursor lentigo maligna (LM) are the most common melanoma types on the head and neck areas. Clinically unsuspected invasion is revealed histologically in 5-52% of LMs. Thus, correct early diagnoses are crucial for determining accurate resection margins and the discussion of sentinel lymph node examination.

Hyperspectral imaging is based on the detection of spectral differences reflecting a tissue’s biological properties and can be used for detecting skin malignancies. The hyperspectral imaging is rapid and spectral data can be obtained from images with a large imaging field. We determined how accurately a hyperspectral imaging system (HIS) detected LMM invasion.

A total of 31 patients, with 32 lesions, with histologically confirmed LM or LMM, participated in the study. The patients’ mean age was 77 years (range 56-97), 16 were female and 15 were male. Four patients had skin photo-type I, 17 had type II and 10 had type III. Fifteen patients had earlier had keratinocyte (pre)malignancies. None of the patients had received immunosuppressant, cytostatics, biological agents or radiotherapy to the studied areas. Two patients had previously received phototherapy.

Of the 32 LM/LMM lesions, 10 were histologically defined as LMMs (eight on the face or scalp, two on the trunk) with Breslow thicknesses between 0.4–1.6 mm (mean 0.85 mm). The remaining 22 lesions were histologically LMs (20 on the face or scalp, two on the trunk).

All lesions were evaluated with dermatoscopy (Dermlite® 3 GenCA, USA) and photographed (Canon Ixus 115 HS, 12.1 megapixel). Hyperspectral images were taken *in vivo* using a HIS camera prototype (VTT FPI VIS-VNIR Spectral Camera), based on a Fabry-Perot interferometer. Within seconds HIS captures the diffuse reflectance of visible and near-infrared light (500–900 nm), with a 12 cm² field of view (spatial resolution 6400 pixels/cm², pixel=125 μm) and imaging depth of approx. 2 mm. In acquired hyperspectral images every pixel...
represents a diffuse reflectance spectrum formed from the mixing of spectra from the different materials in the image. With mathematical modelling this spectrum can be separated into the constituent spectra from these different materials. These "pure" spectra that contribute to the mixed spectrum are called 'endmember' spectra, Fig 1. End-members can further be used to create abundance images showing localization of these spectra in the imaged area, Fig 2-3. The detailed description of the technique has been reported elsewhere 4-5.

For the first four patients, the lesions were excised with 5mm-margins immediately after imaging. In the consequent cases, punch-biopsies were taken from the suspected invasion sites, and thereafter excised entirely using a 5 mm-margin for LM and a 10 mm-margin for LMM. The specimens were fixed in 4% formalin, embedded in paraffin, sectioned in 3-5 mm intervals and stained with hematoxylin-eosin and if needed with MART1 immunohistochemistry. The samples were evaluated by an experienced dermatopathologist (L.J.) blinded to the HIS outcome.

In abundance images, in situ LMs were seen as homogenous white areas while in the case of LMMs the abundance maps showed a dark hole on the site of invasion, and the rest of lesion was seen as a homogenous white area as in LM. These areas, which represented the LMM invasion, were seen as clear white areas in separate abundance maps, Fig 2. This finding was seen in 9/10 LMMs (true positives). HIS did not detect the invasion in one of the 10 LMMs, where the Breslow thickness was 0.5 mm (false negative). In 19/22 LMs HIS did not indicate any sites of invasion (true negatives), Fig 3, whereas in 3/22 cases invasions were suggested, but not detected histologically (false positives). Thus, HIS achieved a positive predictive value of 75%, a negative predictive value of 95%, a sensitivity of 90% and a specificity of 86.3%.

Diffuse reflectance records both light absorption and scattering, and thus provides information on skin morphology and chromophore content. Abundance images represent the localization of each characteristic diffuse reflectance spectra (end-member) in the imaged area. In this study the abundance images showed clear differences between LM and LMM. Most likely, absorption of the skin chromophores (like melanin) at different skin levels explains most of the differences between the spectra.

Many techniques have been developed for melanoma diagnostics. Of these, dermatoscopy and reflectance confocal microscopy (RCM) are widely used 6. Both techniques are excellent in giving the diagnosis, but imprecise in determining the thickness and early invasion. Multispectral imaging devices have also been introduced 7-10. Multispectral devices use multiple (4-15) wide non-continuous bands of wavelengths resulting in discrete spectral images 8,9, while HIS uses continuous narrow wavebands delivered from a tunable filter.

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Different tumor types can be identified by their different spectra\textsuperscript{11-13}. Scanning and microscopic hyperspectral imagers have been assessed for detecting skin cancers\textsuperscript{11-13}. The HIS method used in this study to differentiate the in situ and invasive melanoma from each other, showed much promise in guiding practitioners to obtain more targeted diagnostic biopsies. HIS also proved to be useful for improving the detection and staging of LMM. HIS could also be developed further to measure the thickness of LMM non-invasively prior to surgery, which would save time and resources.

A limitation in our study was its small sample size. A larger multicenter study is warranted to confirm our results. Although the analyzing method is objective, clinicians still need to interpret the HIS images, which, however, needs less expertise than the interpretation of near-histological RCM images. The quality of the HIS images may be affected by artefacts caused by crusts, lesion thickness and differences in the skin color. The HIS is not yet in commercial production. A future goal is to image a large set of various skin tumors for the development of a classification algorithm for non-invasive tumor diagnostics based on spectral data.

A future prospective could be to combine different imaging techniques (dermatoscopy/RCM) and HIS for identification of invasive tumors.

To conclude, HIS is a promising tool to detect basal membrane invasion in LMM, and thus to separate in situ and invasive LMs for more accurate pre-surgical diagnostics.

**CONFLICT OF INTEREST**

The authors declare no conflicts of interest.

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**FIGURE LEGENDS**

Fig 1. Unsupervised mathematical modeling (linear signal un-mixing) was used to separate end-members i.e. characteristic spectra of LM, invasive LMM and healthy skin. These spectra were further used to create abundance images i.e. localization maps of these characteristic spectra in the imaged skin area (Figures 2-3).

Fig 2. Clinical images, histological images and hyperspectral abundance images representing LMM (true positive) a) Photograph of LMM b) Dermatoscopic image of LMM c) Hyperspectral abundance map representing the healthy skin around the lesion (white) d) Hyperspectral abundance map representing the non-invasive part of the LMM-lesion (white) e) Hyperspectral abundance map representing the dermal invasion in

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Fig 3. Clinical images, histopathological images and hyperspectral abundance images representing in situ LM (true negative) a) Photograph of LM b) Dermatoscopic image of LM c) Hyperspectral abundance map representing the healthy skin around a lesion (white) d) Hyperspectral abundance map representing the LM-lesion (white), no suspected invasion was seen e) Haematoxylin and eosin staining of LM f) Haematoxylin and eosin staining of LMM (Breslow 1.6 mm) g) Immunohistochemistry for MART1 in LMM (Breslow 1.6 mm)