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Lymph node characterization in vivo using endoscopic ultrasound spectrum analysis with electronic array echo endoscopes

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Introduction
Endoscopic ultrasound (EUS) is often employed for locoregional staging of gastrointestinal malignancy. However, differentiating between benign and malignant lymph nodes is still sometimes difficult based on EUS appearance alone [1,2]. Although EUS-guided fine needle aspiration (FNA) can obtain diagnostic cytologic material, targeting of tissue for FNA still depends on imaging, and there is still an important need for better methods for distinguishing benign from neoplastic tissue. Our previous studies with mechanical echo endoscopes have shown that spectral parameters from backscattered signals can be used to quantitatively distinguish between benign and malignant lymph nodes [3,4].

In spectrum analysis, the radiofrequency (RF) data of the ultrasound energy that is backscattered from local tissue inhomogeneities is analyzed [5]. The nature of the backscattered signals depends on the effective size and acoustic concentration of the scattering tissues as well as the local variations in acoustic impedance (product of density and sound speed). Different tissue types may then be distinguished because they have different microstructures that scatter ultrasound differently. Spectral parameters provide a quantitative assessment that is independent of the system being used and of the user, given proper calibration. In the context of gastrointestinal cancer, ex vivo studies of lymph node metastases of colorectal cancer have shown that ultrasound backscatter analysis performed better than B-mode ultrasound [6] even when multiple B-mode sonographic parameters were considered [7]. Recent investigations using high frequency ultrasound (25.6 MHz center frequency) to examine ex vivo lymph node specimens have shown specificity and sensitivity as high as 95% with multiple backscatter parameters [8,9].

Over the past decade, radial and curvilinear electronic array echo endoscopes have widely replaced mechanical echo endoscopes. Our aim in this study was to test the ability of spectral analysis of EUS backscattered signals to distinguish between benign and malignant lymph nodes.
Patients and methods

Diagnostic criteria

Patients in the benign node group met all of the following criteria: (i) referred for EUS for reasons other than cancer staging; (ii) no prior or current diagnosis of malignancy; (iii) no identification of mass lesions at EUS; (iv) each lymph node showing at least one of the following EUS features suggestive of benign nodes: (a) draping configuration (b) hyperechoicity (c) node diameter < 1 cm; (v) no diagnosis of cancer in the year following EUS examination as determined by follow-up telephone questionnaire and/or review of medical records.

Patients in the malignant node group met both the following criteria: (i) lymph node identified at EUS; (ii) FNA cytology positive for carcinoma.

Patients

A total of 19 patients were included in the study. FNA-proven malignant lymphadenopathy was shown in 11 patients, associated with malignancy of the pancreas (n=3), lymphoma (n=2), rectum (n=1), lung (n=2), and esophagus (n=3). Benign-appearing lymph nodes were imaged in eight patients, without a diagnosis of malignancy. Indications for these procedures were incidental mediastinal adenopathy (n=3), benign pancreatic disease (n=4), and gastrointestinal stromal tumor (GIST) (n=1). In some cases, multiple lymph nodes were imaged in a patient.

RF data acquisition and analysis

We used radial (GF-UE160-AL5) and curvilinear (GF-UC140P-AL5) electronic-array ultrasonic gastrovideoscopes operated in the 6 MHz mode with a commercially available clinical ultrasound system (Exera EU-ME1; Olympus America, Center Valley, Pennsylvania, USA). The echo endoscopes contained ultrasound transducer arrays which are electronically excited in sets to generate 360° (radial) or 180° (curvilinear) B-scan cross-sectional images, composed of hundreds of A-scan lines. Each image and its corresponding RF data were simultaneously digitized by the Exera EU-ME1 system and automatically saved.

We then imported the RF data into our custom-designed analysis software (written using MATLAB 2010b; Mathworks, Natick, Massachusetts, USA) for offline image reconstruction and data processing. We independently verified that the gain, contrast, and frame correlation settings, which affect the appearance of the system image, did not change the underlying RF data acquired by the system.

Prior to RF data analysis, clinical endoscopists identified and manually segmented regions of interest (ROIs) on the system image according to the evaluation criteria described below. Data analysts then independently translated the ROIs onto the reconstructed image to select corresponding segments of RF data. The sector-shaped ROIs were maximized in size within the designated areas. The fast Fourier transform (FFT) was used to find the power spectra for each segment of A-scan RF data within the ROI after gating the segment by a series of sliding Hamming windows of 0.8 μs (~0.62 mm), each offset by 0.1 μs (~0.077 mm).

Fig. 1a shows a typical example of an ROI chosen for a lymph node using the curvilinear echoendoscope, while Fig. 1b shows that the FNA used to determine the malignancy of the node was obtained in the same tissue area under EUS guidance.

Spectrum calibration

Previous studies have typically removed the artifacts associated with the composite transfer function of the electronic transmitter/receiver and transducer of the EUS system by dividing the power spectrum from the tissue by the spectrum of an ideal reflector. However, we did not have control over the electronic gain settings for the RF data acquisition (not the image gain setting), and, as such, we could not use a strong ultrasound reflector (e.g., glass) without saturating the received signal from an ideal reflector. Instead, calibration was performed using reflections from the flat wall of an acrylic plastic chamber filled with an attenuating gelatin-based phantom [4]. The phantom material was independently measured using the substitution method to have an attenuation of 1.5 dB/MHz/cm, which was sufficient to prevent signal saturation while minimally affecting overall spectral shape.

To obtain the RF calibration data, an 18-mm hole in the phantom was filled with water. We then inserted the transducer, and inflated the endoscope’s balloon with water until the balloon was in full contact with the phantom. We manually adjusted the orientation of the endoscope tip to make the central axis of the endoscope parallel to the wall. We then acquired the RF data, and computed by spectrum by FFT. The calibration spectrum was then compensated for round-trip attenuation. To obtain the calibrated spectra, we then divided the tissue spectra by the calibration spectrum.

Because the calibrated spectra are typically quasi-linear over the ultrasound frequency band used [3], they can be effectively characterized by linear regression using their slope, intercept, and their midband fit, which is the value of the linear function eval-
by computing the area under the ROC curve (AUC) from ROCKIT. To perform all statistical calculations, we used SPSS (Version 16, Chicago, Illinois, USA). We fitted a binormal receiver operating characteristic (ROC) curve to the resulting discriminant function coefficients, and then assessed classification performance through the leave-one-out approach for cross-validation. The overall accuracy of classification was 89% (See Fig. 2 for the corresponding scatterplot and receiver operating characteristic [ROC] curve.)

Results

Table 1 lists the means and standard deviations of the sets of mean spectral parameters obtained from averaging over the ROIs for each benign and malignant lymph node. The t tests showed that midband fit, intercept, and RMS deviation were significantly different between the benign and malignant nodes with the malignant nodes having lower midband fit and intercept, but greater RMS deviation. Table 2 shows the results of the LDA classification using midband fit and RMS deviation and leave-one-out cross-validation. The canonical discriminant function coefficients were $-0.108$ for midband fit and $-1.704$ for the RMS deviation, with a constant value of $-14.877$. (The use of intercept and RMS deviation or in...
tercept and midband fit gave comparable or worse results.) With the malignant outcome considered a “positive” test result, the classification had sensitivity of 83%, specificity of 94%, positive predictive value of 91%, negative predictive value of 88%, and overall accuracy of 89%.

**Fig. 2a** shows the corresponding scatterplot of the data using midband fit and RMS deviation. The discriminant line shows the approximate dividing line between benign and malignant cases according to the LDA. **Fig. 2b** shows the corresponding binormal estimate of the ROC curve with AUC of 0.95.

**Discussion**

In our previous studies, midband fit and intercept were found to best distinguish benign and malignant lymph nodes, although only with a maximum sensitivity of 67%, specificity of 82%, overall accuracy of 73%, and ROC AUC of 0.90 with all data from both studies [4], and therefore below the level of the current results. In those studies two different echo endoscopes had to be used to obtain the RF data and FNA biopsies; however in the current study the curvilinear echoendoscope could obtain both at once (see **Fig. 1**). It is possible that this factor could account for the improved classification. In the current study we also computed the RMS deviation and found that it provided another relatively uncorrelated parameter for discrimination between the benign and malignant nodes.

The lower midband fit and intercept observed here are consistent with our previous studies with the single-element echo endoscopes. The ability of intercept to distinguish between benign and malignant lymph nodes is consistent with previously reported higher (less negative) intercepts for metastatic nodes and malignant lymph nodes. However, it may be difficult to exactly compare these results with the current study given the different transducer types and more highly controlled conditions of our ex vivo studies.

Additional work will be useful to address the limitations of this study. Because histological results were only available for cases with suspected malignancy (malignant lymph nodes), the benign cases had to be inferred from other more indirect clinical and “classic” EUS diagnosis criteria. As such, the study only included fairly well-defined cases, and additional work is needed to assess the utility of the approach in more ambiguous situations. Also, the modest sample size precluded the use of additional criteria (e.g., texture or morphological parameters) with more sophisticated classification methods (e.g., support vector machines) that could result in an improved ability to discriminate between tissue states.

**Conclusion**

This study shows that spectral analysis of the EUS RF backscatter signals from radial and curvilinear electronic-array echo endoscopes can discriminate between benign and malignant lymph nodes in vivo. With further development, this method may prove useful for providing real-time “digitally stained” images with coloration corresponding to the probability of various normal or disease states, thereby providing endoscopists with more timely and improved accuracy of diagnosis with EUS.

**Competing interests:** None

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